MECHANISTIC INVESTIGATION ON SOLAR LIGHT INITIATED DESTRUCTION OF INDICATOR BACTERIA

by

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Dedicated to Lale İpek Lale and Nihat Lale,

to the memory of Ahmet Kara...

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ABSTRACT

MECHANISTIC INVESTIGATION ON SOLAR LIGHT INITIATED DESTRUCTION OF INDICATOR BACTERIA

Due to the increasing shortage of clean drinking water around the world, some point of use disinfection techniques has been improved to obtain purified drinking water from pathogen microorganisms. Water disinfection using natural or artificial sunlight has been widely studied for the inactivation of microorganisms. Pathogens can be destructed with the synergistic effects of the solar light and thermal heat. Recently, application of solar photocatalysis has gained attraction for the effective inactivation of microorganisms. In this study, solar light initiated destruction of bacteria typically Escherichia coli as the model indicator organism and characterization of the released organic matter were evaluated in the presence/absence of photocatalysts, namely TiO₂ and its second generation type Fe-doped TiO₂. Via destruction of cell membrane, contents of released proteins, carbohydrates and potassium were determined. Solar photolytic/photocatalytic inactivation of E. coli was conducted in the presence/absence of water matrix that provides the ionic characteristics of natural waters and in the presence/absence of humic acid that is a dissolved organic matter analogue in natural waters, hence the degradation of humic acid by solar light initiated processes in the presence of E. coli was also investigated. Characterization of organic matter in respect of E. coli inactivation was evaluated by specified and specific ultraviolet-visible and fluorescence parameters, dissolved organic carbon contents, and excitation emission matrix fluorescence contour plots. E. coli inactivation was followed by bacteria reduction kinetic parameters, and detection of organic (protein and carbohydrate) and inorganic (K⁺ ion leakage) contents under specified conditions in relation with the spectroscopic parameters.

ÖZET

İNDİKATÖR BAKTERİNİN GÜNEŞ IŞIĞI İLE BAŞLATILAN TAHRİBİNİN MEKANİK İNCELENMESİ

Dünya genelinde artan temiz içme suyu kıtlığı nedeniyle patojen mikroorganizmalardan arındırılmış içme suyu elde etmek için bazı dezenfeksiyon teknikleri geliştirilmiştir. Mikroorganizmaların inaktivasyonu için, doğal veya yapay güneş ışığı kullanılarak suyun dezenfekte edilmesi konusu detaylı olarak çalışılmıştır. Patojenler güneş ışığı ve termal ısının sinerjistik etkisiyle parçalanabilmektedir. Son zamanlarda, mikroorganizmaların etkin bir şekilde inaktivasyonu için solar fotokatalizin uygulanması önem kazanmıştır. Bu çalışmada, model indikatör organizma olarak kullanılan Escheriscia coli'nin güneş ışığı ile başlatılan tahribi ve ortama salınan organik maddenin karakterizasyonu fotokatalist olarak titanyum dioksit ve onun ikinci nesil çeşidi Fe-yüklü titanyum dioksitin varlığında ve yokluğunda değerlendirilmiştir. Hücre duvarının tahrip edilmesi sırasında ortama salınan protein, karbonhidrat, ve potasyum miktarları tayin edilmiştir. E. coli'nin solar fotolitik/fotokatalitik inaktivasyonu doğal su kaynaklarının iyon karakteristiklerini sağlayan su ortamının varlığında ve yokluğunda, ve doğal su kaynaklarında çözünmüş organik madde analoğu olan hümik asidin varlığında ve yokluğunda yürütülmüş, böylece E. coli varlığında hümik asidin güneş ışığı ile başlatılan degredasyon prosesi de incelenmiştir. E. coli inaktivasyonuyla bağlantılı olarak organik madde karakterizasyonu spesifik ultraviyole-görünür bölge ve floresans parametreleri, çözünmüş organik karbon içerikleri ve eksitasyon emisyon matrisi floresans kontür çizimleri ile değerlendirilmiştir. E. coli inaktivasyonu, bakteri azalım kinetik parametreleri, ve spektroskopik parametrelerle bağlantılı olarak organik (protein ve karbonhidrat) ve inorganik (K+ iyonu sızıntısı) içerik tayinleri ile açıklanmıştır.

TABLE OF CONTENTS

ACKNOWLEDGEMENTS	iii
ABSTRACT	iv
ÖZET	v
TABLE OF CONTENTS	vi
LIST OF FIGURES	xiv
LIST OF TABLES	.xxxvi
LIST OF SYMBOLS/ABBREVIATIONS	xliii
1. INTRODUCTION	1
2. LITERATURE REVIEW	3
2.1. Solar Disinfection of Water	3
2.2. Mechanism of Solar Disinfection	3
2.3. Heterogenous Photocatalysis for the Enhancement of Solar Efficiency	5
2.3.1. Photocatalytic Oxidation Processes	6
2.3.2. Bare and Visible Light Activated Doped TiO ₂ Species	6
2.4. Indicator Organism: Escherichia coli	8
2.4.1. E. coli Motility	10
2.4.2. Surface Properties and Point of Zero Charge of <i>E. coli</i>	11
2.4.3. E. coli Inactivation Mechanism During Solar Light Initiated Photocatalytic	
Processes	12
2.4.3.1. Effect of reactive oxygen species on solar light initiated photocatalytic	2
bacteria destruction	12
2.4.3.2. Cellular targets of reactive oxygen species	14
Bacterial capsule as extracellular polymeric substances	14
Cell wall and cell membrane	14
Enzymes	17
Proteins and DNA	17
2.4.3.3. Detection of bacteria inactivation and mineralization	18
2.5. Water Matrix	18
2.5.1. Effects of Water Matrix Components on Solar Light Initiated Photocatalytic	
Processes	18
2.5.1.1. Effect of chloride	19
2.5.1.2. Effect of nitrate	19

	2.5.1.3.	Effect of sulfate	20
	2.5.1.4.	Effect of phosphate	21
	2.5.1.5.	Effect of bicarbonate	21
	2.5.1.6.	Effect of monovalent cations	21
	2.5.1.7.	Effect of divalent cations	22
2.6. Natu	ral Organi	c Matter	23
2.6.1	. Humic S	Substances and Humic Acid	24
2.6.2	. Photoch	nemistry of Humic Substances in Natural Water Systems	28
MATERI	ALS AND	D METHODS	34
3.1. Mate	rials		34
3.1.1	. Preparat	tion of <i>E. coli</i> Suspension	34
3.1.2	. Luria Be	ertani Broth	34
3.1.3	. CHRON	Magar TM ECC	34
3.1.4	. Isotonic	e Solution	34
3.1.5	. Water N	Matrix Solution	35
3.1.6	. Humic A	Acid Solution	35
3.1.7	. Photoca	atalyst Specimens	35
3.2. Meth	ods		36
3.2.1	. Experin	nental Set-Up	36
3.2.2	. Experin	nental Procedure	36
3.2.3	. Analytic	cal Methods	37
	3.2.3.1.	<i>E. coli</i> enumeration	37
	3.2.3.2.	Total protein determination	39
		Procedure	39
		The Lowry Reagent	39
		Solution A (alkaline solution)	40
		Solution B	40
		Solution C	40
		The Folin-Ciocalteu reagent, 1N	40
	3.2.3.3.	Total carbohydrate determination	40
		Procedure	41
		5 % Phenol solution	41
	3.2.3.4.	K ⁺ leakage determination	41
	3.2.3.5.	Endotoxin determination	41
	3.2.3.6.	UV-vis spectroscopic measurements	42
	 2.6. Natur 2.6.1 2.6.2 MATERI 3.1.1 3.1.2 3.1.3 3.1.4 3.1.5 3.1.6 3.1.7 3.2. Meth 3.2.1 3.2.2 3.2.3 	2.5.1.3. 2.5.1.4. 2.5.1.5. 2.5.1.6. 2.5.1.7. 2.6. Natural Organi 2.6.1. Humic S 2.6.2. Photock MATERIALS ANI 3.1. Materials 3.1.1. Prepara 3.1.2. Luria B 3.1.3. CHRON 3.1.4. Isotonic 3.1.5. Water M 3.1.6. Humic J 3.1.7. Photoca 3.2. Methods 3.2.1. Experir 3.2.2. Experir 3.2.3. Analyti 3.2.3.1. 3.2.3.2. 3.2.3.2.	2.5.1.3. Effect of sulfate. 2.5.1.4. Effect of phosphate. 2.5.1.5. Effect of bicarbonate 2.5.1.6. Effect of monovalent cations. 2.5.1.7. Effect of divalent cations. 2.6. Natural Organic Matter. 2.6.1. Humic Substances and Humic Acid. 2.6.2. Photochemistry of Humic Substances in Natural Water Systems. MATERIALS AND METHODS. 3.1. Materials. 3.1.1. Preparation of <i>E. coli</i> Suspension. 3.1.2. Luria Bertani Broth. 3.1.3. CHROMagar TM ECC 3.1.4. Isotonic Solution. 3.1.5. Water Matrix Solution. 3.1.6. Humic Acid Solution. 3.1.7. Photocatalyst Specimens. 3.2.1. Experimental Set-Up. 3.2.2. Experimental Set-Up. 3.2.3.1. <i>E. coli</i> enumeration. 3.2.3.1. <i>E. coli</i> enumeration. 3.2.3.2. Total protein determination. Procedure. The Lowry Reagent. Solution A. (alkaline solution). Solution C. The Folin-Ciocalteu reagent, IN. 3.2.3.3. Total carbohydrate determination. Procedure. 5% Phenol solution. 3.2.3.4. K ⁺ leakage determination. 3.2.3.5. Endotoxin determinati

3.2.3.7. Fluorescence spectroscopic measurements	43
3.2.3.8. Non-purgeable organic carbon measurements	43
I. RESULTS AND DISCUSSION	44
4.1. Solar Photolytic Inactivation of <i>E. coli</i>	45
4.1.1. Solar Photolytic E. coli Inactivation Under Specified Reaction	Conditions,
Reductions and Kinetics	45
4.1.2. Organic Matter Formation and Removal Upon Solar Photolytic	Treatment
of <i>E. coli</i> in IsoT Solution	48
4.1.3. Mechanistic Evaluation of E. coli Inactivation Upon Solar Phot	olytic
Treatment in IsoT Solution	53
4.1.4. Organic Matter Formation and Removal Upon Solar Photolytic	Treatment
of <i>E. coli</i> in WM solution	55
4.1.5. Mechanistic Evaluation of E. coli Inactivation Upon Solar Phot	olytic
Treatment in WM Solution	60
4.1.6. Organic Matter Formation and Removal Upon Solar Photolytic	Treatment
of <i>E. coli</i> in HA solution	61
4.1.7. Mechanistic Evaluation of E. coli Inactivation Upon Solar Phot	olytic
Treatment in HA Solution	
4.1.8. Organic Matter Formation and Removal Upon Solar Photolytic	Treatment
of <i>E. coli</i> in "HA and WM" solution	67
4.1.9. Mechanistic Evaluation of E. coli Inactivation Upon Solar Phot	olytic
Treatment in "HA and WM" Solution	72
4.1.10. Comparative Evaluation of Different Experimental Matrixes of	n Solar
Photolytic Inactivation Mechanism of <i>E. coli</i>	73
4.2. Solar Photocatalytic Inactivation of <i>E. coli</i>	77
4.2.1. TiO ₂ Solar Photocatalytic Inactivation of <i>E. coli</i>	77
4.2.1.1. TiO ₂ solar photocatalytic <i>E. coli</i> inactivation under spe	cified reaction
conditions, reductions, and kinetics	77
4.2.1.2. Organic matter formation and removal upon TiO_2 solar	r photocatalytic
treatment of <i>E. coli</i> in IsoT solution	79
4.2.1.3. Mechanistic evaluation of <i>E. coli</i> inactivation upon Ti	D ₂ solar
photocatalytic treatment in IsoT solution	84
4.2.1.4. Organic matter formation and removal upon TiO_2 solar	r photocatalytic
treatment of <i>E. coli</i> in WM solution	85
4.2.1.5. Mechanistic evaluation of <i>E. coli</i> inactivation upon Ti	O_2 solar

viii

		photocatalytic treatment in WM solution
	4.2.1.6.	Organic matter formation and removal upon TiO ₂ solar photocatalytic
		treatment of <i>E. coli</i> in HA solution91
	4.2.1.7.	Mechanistic evaluation of E. coli inactivation upon TiO2 solar
		photocatalytic treatment in HA solution97
	4.2.1.8.	Organic matter formation and removal upon TiO ₂ solar photocatalytic
		treatment of <i>E. coli</i> in "HA and WM" solution98
	4.2.1.9.	Mechanistic evaluation of E. coli inactivation upon TiO ₂ solar
		photocatalytic treatment in "HA and WM" solution103
	4.2.1.10	. Comparative evaluation of different experimental matrixes on TiO ₂
		solar photocatalytic inactivation mechanism of <i>E. coli</i> 104
4.2.2.	SynTiO ₂	Solar Photocatalytic Inactivation of <i>E. coli</i> 108
	4.2.2.1.	SynTiO ₂ solar photocatalytic <i>E. coli</i> inactivation under specified
		reaction conditions, reductions, and kinetics108
	4.2.2.2.	Organic matter formation and removal upon SynTiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in IsoT solution110
	4.2.2.3.	Mechanistic evaluation of E. coli inactivation upon SynTiO ₂ solar
		photocatalytic treatment in IsoT solution114
	4.2.2.4.	Organic matter formation and removal upon SynTiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in WM solution115
	4.2.2.5.	Mechanistic evaluation of E. coli inactivation upon SynTiO ₂ solar
		photocatalytic treatment in WM solution120
	4.2.2.6.	Organic matter formation and removal upon SynTiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in HA solution121
	4.2.2.7.	Mechanistic evaluation of <i>E. coli</i> inactivation upon SynTiO ₂ solar
		photocatalytic treatment in HA solution126
	4.2.2.8.	Organic matter formation and removal upon SynTiO ₂ solar
		photocatalytic treatment of E. coli in "HA and WM" solution127
	4.2.2.9.	Mechanistic evaluation of E. coli inactivation upon SynTiO ₂ solar
		photocatalytic treatment in "HA and WM" solution132
	4.2.2.10	. Comparative evaluation of different experimental matrixes on
		SynTiO ₂ solar photocatalytic inactivation mechanism of <i>E. coli</i> 133
4.2.3.	0.25% F	e-TiO ₂ Solar Photocatalytic Inactivation of <i>E. coli</i> 136
	4.2.3.1.	0.25% Fe-TiO ₂ solar photocatalytic <i>E. coli</i> inactivation under specified
		reaction conditions, reductions, and kinetics

	4.2.3.2.	Organic matter formation and removal upon 0.25% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in IsoT solution138
	4.2.3.3.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂
		solar photocatalytic treatment in IsoT solution143
	4.2.3.4.	Organic matter formation and removal upon 0.25% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in WM solution144
	4.2.3.5.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂
		solar photocatalytic treatment in WM solution149
	4.2.3.6.	Organic matter formation and removal upon 0.25% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in HA solution150
	4.2.3.7.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂
		solar photocatalytic treatment in HA solution155
	4.2.3.8.	Organic matter formation and removal upon 0.25% Fe-TiO ₂ solar
		photocatalytic treatment of E. coli in "HA and WM" solution156
	4.2.3.9.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂
		solar photocatalytic treatment in "HA and WM" solution161
	4.2.3.10	. Comparative evaluation of different experimental matrixes on
		0.25% Fe-TiO ₂ solar photocatalytic inactivation mechanism of <i>E. coli</i> 162
4.2.4.	0.50% F	e-TiO ₂ Solar Photocatalytic Inactivation of <i>E. coli</i> 165
	4.2.4.1.	0.50% Fe-TiO ₂ solar photocatalytic <i>E. coli</i> inactivation under specified
		reaction conditions, reductions, and kinetics165
	4.2.4.2.	Organic matter formation and removal upon 0.50% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in IsoT solution167
	4.2.4.3.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-TiO ₂
		solar photocatalytic treatment in IsoT solution172
	4.2.4.4.	Organic matter formation and removal upon 0.50% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in WM solution173
	4.2.4.5.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-TiO ₂
		solar photocatalytic treatment in WM solution177
	4.2.4.6.	Organic matter formation and removal upon 0.50% Fe-TiO ₂ solar
		photocatalytic treatment of <i>E. coli</i> in HA solution178
	4.2.4.7.	Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-TiO ₂
		solar photocatalytic treatment in HA solution183
	4.2.4.8.	Organic matter formation and removal upon 0.50% Fe-TiO ₂ solar
		photocatalytic treatment of E. coli in "HA and WM" solution184

4.2.4.9. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-TiO ₂	
solar photocatalytic treatment in "HA and WM" solution	.189
4.2.4.10. Comparative evaluation of different experimental matrixes on	
0.50% Fe-TiO ₂ solar photocatalytic inactivation mechanism of <i>E. coli</i>	190
4.2.5. 0.25% Fe-SynTiO ₂ Solar Photocatalytic Inactivation of <i>E. coli</i>	193
4.2.5.1. 0.25% Fe-SynTiO ₂ solar photocatalytic <i>E. coli</i> inactivation under	
specified reaction conditions, reductions, and kinetics	.193
4.2.5.2. Organic matter formation and removal upon 0.25% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in IsoT solution	195
4.2.5.3. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-SynTiO ₂	
solar photocatalytic treatment in IsoT solution	200
4.2.5.4. Organic matter formation and removal upon 0.25% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in WM solution	201
4.2.5.5. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-SynTiO ₂	
solar photocatalytic treatment in WM solution	206
4.2.5.6. Organic matter formation and removal upon 0.25% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in HA solution	.207
4.2.5.7. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-SynTiO ₂	
solar photocatalytic treatment in HA solution	212
4.2.5.8. Organic matter formation and removal upon 0.25% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in "HA and WM" solution	213
4.2.5.9. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.25% Fe-SynTiO ₂	
solar photocatalytic treatment in "HA and WM" solution	.218
4.2.5.10. Comparative evaluation of different experimental matrixes on	
0.25% Fe-SynTiO ₂ solar photocatalytic inactivation mechanism of	
E. coli	.219
4.2.6. 0.50% Fe-SynTiO ₂ Solar Photocatalytic Inactivation of <i>E. coli</i>	222
4.2.6.1. 0.50% Fe-SynTiO ₂ solar photocatalytic <i>E. coli</i> inactivation under	
specified reaction conditions, reductions, and kinetics	.222
4.2.6.2. Organic matter formation and removal upon 0.50% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in IsoT solution	224
4.2.6.3. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-SynTiO ₂	
solar photocatalytic treatment in IsoT solution	228
4.2.6.4. Organic matter formation and removal upon 0.50% Fe-SynTiO ₂ solar	
photocatalytic treatment of <i>E. coli</i> in WM solution	229

2	4.2.6.5. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment in WM solution	3
2	4.2.6.6. Organic matter formation and removal upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment of <i>E. coli</i> in HA solution234	4
Ζ	4.2.6.7. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment in HA solution	3
Ζ	4.2.6.8. Organic matter formation and removal upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment of <i>E. coli</i> in "HA and WM" solution239)
Ζ	4.2.6.9. Mechanistic evaluation of <i>E. coli</i> inactivation upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment in "HA and WM" solution244	1
Z	4.2.6.10. Comparative evaluation of different experimental matrixes on	
	0.50% Fe-SynTiO ₂ solar photocatalytic inactivation mechanism of	
	<i>E. coli</i>	5
4.3. Compa	rative Evaluation of E. coli Inactivation Upon Use of Photocatalyst	
Specim	nens)
4.3.1.	Comparative Efficiency Evaluation of Different Photocatalyst Specimens on	
	<i>E. coli</i> Inactivation in IsoT Solution249)
	4.3.1.1. Comparative mechanistic evaluation of <i>E. coli</i> inactivation in IsoT	
	solution upon solar photocatalytic treatment with undoped and	
	Fe-doped TiO ₂ specimens25	l
4.3.2.	Comparative Efficiency Evaluation of Different Photocatalyst Specimens on	
	<i>E. coli</i> Inactivation in WM Solution252	2
	4.3.2.1. Comparative mechanistic evaluation of <i>E. coli</i> inactivation in WM	
	solution upon solar photocatalytic treatment with undoped and	
	Fe-doped TiO ₂ specimens254	1
4.3.3.	Comparative Efficiency Evaluation of Different Photocatalyst Specimens on	
	<i>E. coli</i> Inactivation in HA Solution250	5
	4.3.3.1. Comparative mechanistic evaluation of <i>E. coli</i> inactivation in HA	
	solution upon solar photocatalytic treatment with undoped and	
	Fe-doped TiO ₂ specimens258	3
4.3.4.	Comparative Efficiency Evaluation of Different Photocatalyst Specimens on	
	<i>E. coli</i> Inactivation in "HA and WM" Solution260)
	4.3.4.1. Comparative mechanistic evaluation of <i>E. coli</i> inactivation in	
	"HA and WM" solution upon solar photocatalytic treatment with	
	undoped and Fe-doped TiO ₂ specimens262	2

4.4. Endotoxin Release Upon Solar Photolytic and Photocatalytic Inactivation of E. coli	
in Various Aqueous Solutions	264
5. CONCLUSION	266
REFERENCES	268
APPENDIX A: COMPOSITION OF WATER MATRIX SOLUTION	298
APPENDIX B: SPECTROSCOPIC CHARACTERIZATION OF 100 kDa HUMIC	
ACID SOLUTION	299
APPENDIX C: CHARACTERIZATION OF PHOTOCATALYST SPECIMENS	301
APPENDIX D: CALIBRATION CURVE OF BOVINE SERUM ALBUMIN STANDARD	303
APPENDIX E: CALIBRATION CURVE OF GLUCOSE STANDARD	304

LIST OF FIGURES

Figure 2.1.	Proposed schematic process of cell destruction4
Figure 2.2.	a. Formation of surface hydroxylated species of TiO ₂ according to pH;
	b. pH dependent electrostatic interactions on TiO ₂ surface7
Figure 2.3.	The general structure of a prokaryotic organism. Red-labeled structures
	are found in all bacteria species, while black-labeled ones are not9
Figure 2.4.	a. CCW, CW flagella rotations and movement of <i>E. coli</i> ; b. Run and tumble
	movements of a bacterium. Blue arrows indicate the flagellar rotation direction and
	gray arrows indicate the movement direction; c. Random walk pattern of bacteria10
Figure 2.5.	Transmission electron microscopy (TEM) image of E. coli with TiO ₂ particles
	attached on it12
Figure 2.6.	Possible mechanism of photocatalytic disinfection of bacteria13
Figure 2.7.	a. Structural membrane composition and possible photocatalytic disinfection
	mechanism; b. Lipid peroxidation and membrane leakage process in
	Gram-negative bacteria e.g. <i>E. coli</i> 13
Figure 2.8.	Cell wall and cell membrane structure of Gram-negative bacteria14
Figure 2.9.	Schematic diagram of ROS utilized lipid peroxidation reaction15
Figure 2.10	. Venn diagram of various NOM forms found in natural waters.
	TOM or NOM, TOC, DOM, DOC, POC, DON, DOP
	and humic and non-humic materials are represented23
Figure 2.11	. Organic matter mass (Da) and size (μm) distribution diagram of NOM species
	and organisms in natural waters. Molecular weight (MW), humic acid (HAc),
	fulvic acid (FAc), hydrocarbons (HC), carbohydrates (CHO), fatty acids (FA),

xiv

	amino acid (AA)24
Figure 2.12.	Conceptual model of polydisperse HSs25
Figure 2.13.	The postulated relation of HSs25
Figure 2.14.	a. Conformational changes of HA molecules through alkaline to acidic pH;A. Charge repulsion at alkaline pH, B. Intramolecular aggregation at decreasing pH,C. Intermolecular aggregation at decreasing pH, D. Precipitation at acidic pH;b. Tail, loop, and train formations of a HA polymer chain adsorbed on the surface26
Figure 2.15.	a. Hypothetical structure of HA; b. Functional groups and their range of pK _a values present in HA; c. Theoretical structure of HA; d. Proposed three-dimensional structure of HA
Figure 2.16.	a. UV-vis absorption spectra; b. emission scan; and c. synchronous scan fluorescence spectra of IHSS soil HA (SHA), IHSS HA, IHSS FA, Aldrich HA (AHA), Roth HA (RHA), and Nordic HA (NHA)28
Figure 2.17.	Representative functional groups and building blocks of HSs
Figure 2.18.	Diagram of EEM fluorescence contour plot regions
Figure 3.1. S	Solar simulator Atlas-Suntest CPS+ and reaction vessel
Figure 3.2.	<i>E. coli</i> cultured CHROMagar TM ECC plates. Top Left: 10^{-2} dilution from initial <i>E. coli</i> solution, inoculation volume: $10 \ \mu$ L. Top Right: 10^{-5} dilution from <i>E. coli</i> suspension, inoculation volume: $5 \ \mu$ L. Middle Left: 10^{-3} dilution from initial <i>E. coli</i> solution, inoculation volume: $20 \ \mu$ L. Middle Right: 10^{-3} dilution from t=0 <i>E. coli</i> solution, inoculation volume: $10 \ \mu$ L. Bottom Left: 10^{-2} dilution from t=0 <i>E. coli</i> solution, inoculation volume: $10 \ \mu$ L. Bottom Right: 10^{-2} dilution from t _{irr} =30 min <i>E. coli</i> solution, inoculation volume: $50 \ \mu$ L

Figure 3.3. Schematic reaction of phenol-sulfuric acid method......40

Figure 4.1.	Irradiation time dependent <i>E. coli</i> inactivation upon solar photolytic treatment in different solution matrixes46
Figure 4.2.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon solar photolytic treatment
Figure 4.3.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon solar photolytic treatment49
Figure 4.4.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon solar photolytic treatment50
Figure 4.5.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT upon solar photolytic treatment
Figure 4.6.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon solar photolytic treatment
Figure 4.7.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment
Figure 4.8.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment
Figure 4.9.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment
Figure 4.10	 Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment57
Figure 4.11	. EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment
Figure 4.12	 UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment61

Figure 4.13.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment62
Figure 4.14.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment62
Figure 4.15.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment63
Figure 4.16.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment65
Figure 4.17.	UV-vis absorption spectra of organic matter. Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment67
Figure 4.18.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment
Figure 4.19.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment
Figure 4.20.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment
Figure 4.21.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment71
Figure 4.22.	Initial and t _{irr} =60 min total K, protein, and carbohydrate contents, and bacteria enumeration results of <i>E. coli</i> inactivation upon solar photolytic treatment in various aqueous matrixes73

Figure 4.23. Released protein and carbohydrate contents, and bacteria LRV of E. coli

	inactivation upon solar photolytic treatment in various aqueous matrixes74
Figure 4.24.	Initial and t_{irr} =60 min protein and carbohydrate contents, FI _{sync280} , UV ₂₈₀ , NPOC,
	and bacteria enumeration results upon solar photolytic treatment of E. coli in
	various aqueous matrixes76
Figure 4.25.	Irradiation time dependent E. coli inactivation upon TiO ₂ solar photocatalytic
	treatment in various solution matrixes77
Figure 4.26.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in IsoT solution upon TiO ₂ solar photocatalytic treatment80
Figure 4.27.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent <i>E. coli</i> inactivation in IsoT solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.28.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time
	dependent E. coli inactivation in IsoT solution upon TiO2 solar photocatalytic
	treatment
Figure 4.29.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time
	dependent E. coli inactivation in IsoT solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.30.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in IsoT solution upon TiO ₂ solar photocatalytic treatment83
Figure 4.31.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in WM solution upon TiO ₂ solar photocatalytic treatment85
Figure 4.32.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in WM solution upon TiO ₂ solar photocatalytic
	treatment

Figure 4.33. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time

	dependent E. coli inactivation in WM solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.34.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time
	dependent E. coli inactivation in WM solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.35.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in WM solution upon TiO ₂ solar photocatalytic treatment89
Figure 4.36.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in HA solution upon TiO ₂ solar photocatalytic treatment91
Figure 4.37.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in HA solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.38.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time
	dependent E. coli inactivation in HA solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.39.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time
	dependent E. coli inactivation in HA solution upon TiO ₂ solar photocatalytic
	treatment
Figure 4.40.	EEM fluorescence contour plots of organic matter. Irradiation time dependent
	<i>E. coli</i> inactivation in HA solution upon TiO ₂ solar photocatalytic treatment96
Figure 4.41.	UV-vis absorption spectra of organic matter: Irradiation time dependent E. coli
	inactivation in "HA and WM" solution upon TiO ₂ solar photocatalytic treatment98
Figure 4.42.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in "HA and WM" solution upon TiO ₂ solar
	photocatalytic treatment

Figure 4.43.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon TiO ₂ solar photocatalytic treatment
Figure 4.44.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in "HA and WM" solution upon TiO ₂ solar
	photocatalytic treatment101
Figure 4.45.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in "HA and WM" solution upon TiO ₂ solar photocatalytic
	treatment102
Figure 4.46.	Initial, t=0, and tirr=60 min total K, protein, carbohydrate content, and bacteria
	enumeration results of <i>E. coli</i> inactivation upon TiO ₂ solar photocatalytic
	treatment in various aqueous matrixes105
Figure 4.47.	Released (Left) and degraded (Right) protein and carbohydrate contents, and
	bacteria LRV of E. coli inactivation upon TiO2 solar photocatalytic treatment
	in various aqueous matrixes106
Figure 4.48.	Initial, t=0, and t _{irr} =60 min protein and carbohydrate contents, FI _{sync280} , UV ₂₈₀ ,
	NPOC, and bacteria enumeration results of TiO ₂ solar photocatalytic treatment
	of <i>E. coli</i> in various aqueous matrixes107
Figure 4.49.	Irradiation time dependent E. coli inactivation upon SynTiO ₂ solar
	photocatalytic treatment in various solution matrixes108
Figure 4.50.	UV-vis absorption spectra of organic matter: Irradiation time dependent E. coli
	inactivation in IsoT solution upon SynTiO ₂ solar photocatalytic treatment110
Figure 4.51.	Synchronous scan fluorescence spectra of organic matter: Irradiation
	time dependent E. coli inactivation in IsoT solution upon SynTiO ₂ solar
	photocatalytic treatment111

Figure 4.52. Specified UV-vis and fluroscence parameters of organic matter: Irradiation

	time dependent <i>E. coli</i> inactivation in IsoT solution upon SynTiO ₂ solar photocatalytic treatment111
Figure 4.53.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.54.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.55.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon SynTiO ₂ solar photocatalytic treatment115
Figure 4.56.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.57.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.58.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.59.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon SynTiO ₂ solar photocatalytic treatment
Figure 4.60.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon SynTiO ₂ solar photocatalytic treatment121
Figure 4.61.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon SynTiO ₂ solar

	photocatalytic treatment	121
Figure 4.62.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon SynTiO ₂ solar photocatalytic treatment.	122
Figure 4.63.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon SynTiO ₂ solar photocatalytic treatment	123
Figure 4.64.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon SynTiO ₂ solar photocatalytic treatment	125
Figure 4.65.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment	127
Figure 4.66.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment.	128
Figure 4.67.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment	129
Figure 4.68.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment	.130
Figure 4.69.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment	131

xxii

Figure 4.70. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents, and

	bacteria enumeration results of <i>E. coli</i> inactivation upon SynTiO ₂ solar photocatalytic treatment in various aqueous matrixes133
Figure 4.71.	Released protein and carbohydrate contents, and bacteria LRV of <i>E. coli</i> inactivation upon SynTiO ₂ solar photocatalytic treatment in various aqueous matrixes
Figure 4.72.	Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI _{sync280} , UV ₂₈₀ , NPOC, and bacteria enumeration results of SynTiO ₂ solar photocatalytic treatment of <i>E. coli</i> in various aqueous matrixes135
Figure 4.73.	Irradiation time dependent <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂ solar photocatalytic treatment in various solution matrixes
Figure 4.74.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.75.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.76.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.77.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.78.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment

xxiii

Figure 4.79. UV-vis absorption spectra of organic matter: Irradiation time dependent

	<i>E. coli</i> inactivation in WM solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.80.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.81.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.82.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.83.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in WM solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.84.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.85.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.86.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment
Figure 4.87.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment

xxiv

Figure 4.88.	EEM fluorescence contour plots of organic matter: Irradiation time dependent	
	<i>E. coli</i> inactivation in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment	154
Figure 4.89.	UV-vis absorption spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment	156
Figure 4.90.	Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment	157
Figure 4.91.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment.	158
Figure 4.92.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment	158
Figure 4.93.	EEM fluorescence contour plots of organic matter: Irradiation time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment	160
Figure 4.94.	Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents, and bacteria enumeration results of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂ solar photocatalytic treatment in various aqueous matrixes	162
Figure 4.95.	Released protein and carbohydrate contents, and bacteria LRV of <i>E. coli</i> inactivation upon 0.25% Fe-TiO ₂ solar photocatalytic treatment in various aqueous matrixes.	163
Figure 4.96.	Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI _{sync280} , UV ₂₈₀ , NPOC, and bacteria enumeration results of 0.25% Fe-TiO ₂ solar photocatalytic treatment of <i>E. coli</i> in various aqueous matrixes	164

Figure 4.97.	Irradiation time dependent E. coli inactivation upon 0.50% Fe-TiO2
	solar photocatalytic treatment in various solution matrixes165
Figure 4.98.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	<i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-TiO ₂ solar photocatalytic
	treatment167
Figure 4.99.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in IsoT solution upon 0.50% Fe-TiO ₂ solar
	photocatalytic treatment
Figure 4.100	. Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in IsoT solution upon 0.50% Fe-TiO ₂
	solar photocatalytic treatment
Figure 4.101	. Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in IsoT solution upon 0.50% Fe-TiO2
	solar photocatalytic treatment
Figure 4.102	. EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in IsoT solution upon 0.50% Fe-TiO2 solar photocatalytic
	treatment171
Figure 4.103	. UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in WM solution upon 0.50% Fe-TiO ₂ solar photocatalytic
	treatment173
Figure 4.104	. Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in WM solution upon 0.50% Fe-TiO ₂ solar
	photocatalytic treatment174
Figure 4.105	. Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in WM solution upon 0.50% Fe-TiO ₂
	solar photocatalytic treatment174

xxvi

Figure 4.106.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent <i>E. coli</i> inactivation in WM solution upon 0.50% Fe-TiO ₂
	solar photocatalytic treatment175
Figure 4.107.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in WM solution upon 0.50% Fe-TiO ₂ solar photocatalytic
	treatment
Figure 4.108.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in HA solution upon 0.50% Fe-TiO ₂ solar photocatalytic
	treatment
Figure 4.109.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in HA solution upon 0.50% Fe-TiO2 solar
	photocatalytic treatment179
Figure 4.110.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in HA solution upon 0.50% Fe-TiO2
	solar photocatalytic treatment
Figure 4.111.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent <i>E. coli</i> inactivation in HA solution upon 0.50% Fe-TiO ₂
	solar photocatalytic treatment
Figure 4.112.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in HA solution upon 0.50% Fe-TiO ₂ solar photocatalytic
	treatment
Figure 4.113.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂ solar
	photocatalytic treatment
Figure 4.114.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂
	solar photocatalytic treatment

Figure 4.115.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation time	
	dependent E. coli inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂	
	solar photocatalytic treatment	6
Figure 4.116.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation time	
	dependent E. coli inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂	
	solar photocatalytic treatment	6
Figure 4.117.	EEM fluorescence contour plots of organic matter: Irradiation time dependent	
	E. coli inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂ solar	
	photocatalytic treatment	8
F ' 4 1 1 0		
Figure 4.118.	Initial, t=0, and t _{irr} =60 min total K, protein, and carbonydrate contents, and	
	bacteria enumeration results of <i>E. coli</i> inactivation upon 0.50% Fe-11O ₂ solar	0
	photocatalytic treatment in various aqueous matrixes	'U
Figure 4.119.	Released protein and carbohydrate contents, and bacteria LRV of <i>E. coli</i>	
0	inactivation upon 0.50% Fe-TiO ₂ solar photocatalytic treatment in various	
	aqueous matrixes	1
Figure 4.120.	Initial, t=0, and t _{irr} =60 min protein and carbohydrate contents, FI _{sync280} , UV ₂₈₀ ,	
	NPOC and bacteria enumeration results of 0.50% Fe-TiO ₂ solar photocatalytic	
	treatment of <i>E. coli</i> in various aqueous matrixes19	2
E	Les disting time dans tot E line disting and 0.250/ E. Cautio	
rigure 4.121.	alangh at actualitie tractine and in containing achieve achieve achieve actualities	12
	solar photocatalytic treatment in various solution matrixes	'3
Figure 4.122.	UV-vis absorption spectra of organic matter: Irradiation time dependent	
-	<i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-SynTiO ₂ solar	
	photocatalytic treatment	15
Figure 4.123.	Synchronous scan fluorescence spectra of organic matter: Irradiation time	
	dependent E. coli inactivation in IsoT solution upon 0.25% Fe-SynTiO ₂	
	solar photocatalytic treatment19	6

Figure 4.124.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in IsoT solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment196
Figure 4.125.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.126.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in IsoT solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic
	treatment
Figure 4.127.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in WM solution upon 0.25% Fe-SynTiO ₂ solar
	photocatalytic treatment
Figure 4.128.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in WM solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.129.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent <i>E. coli</i> inactivation in WM solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.130.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in WM solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment203
Figure 4.131.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in WM solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic
	treatment
Figure 4.132.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in HA solution upon 0.25% Fe-SynTiO ₂ solar
	photocatalytic treatment207

Figure 4.133.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
	dependent E. coli inactivation in HA solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.134.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in HA solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.135.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in HA solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.136.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in HA solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic
	treatment
Figure 4.137.	UV-vis absorption spectra of organic matter: Irradiation time dependent
	E. coli inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.138.	Synchronous scan fluorescence spectra of organic matter: Irradiation
	time dependent E. coli inactivation in "HA and WM" solution upon
	0.25% Fe-SynTiO ₂ solar photocatalytic treatment214
Figure 4.139.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in "HA and WM" solution upon
	0.25% Fe-SynTiO ₂ solar photocatalytic treatment215
Figure 4.140.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
	time dependent E. coli inactivation in "HA and WM" solution upon
	0.25% Fe-SynTiO ₂ solar photocatalytic treatment
Figure 4.141.	EEM fluorescence contour plots of organic matter: Irradiation time dependent
	E. coli inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO ₂ solar
	photocatalytic treatment

XXX

Figure 4.142.	Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of <i>E. coli</i> inactivation upon 0.25% Fe-SynTiO ₂
	solar photocatalytic treatment in various aqueous matrixes
Figure 4.143.	Released protein and carbohydrate contents, and bacteria LRV of E. coli
	inactivation upon 0.25% Fe-SynTiO ₂ solar photocatalytic treatment in
	various aqueous matrixes
Figure 4.144.	Initial, t=0, and t _{irr} =60 min protein and carbohydrate contents, FI _{sync280} ,
	UV_{280} , NPOC, and bacteria enumeration results of 0.25% Fe-SynTiO ₂ solar
	photocatalytic treatment of <i>E. coli</i> in various aqueous matrixes221
Figure 4.145.	Irradiation time dependent E. coli inactivation upon 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment in various solution matrixes
Figure 4 146	UV-vis absorption spectra of organic matter: Irradiation time dependent
C	<i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-SynTiO ₂ solar
	photocatalytic treatment
Figure 4.147.	Synchronous scan fluorescence spectra of organic matter: Irradiation time
-	dependent <i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.148.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation
C	time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.149.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation
C	time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment
Figure 4.150.	EEM fluorescence contour plots of organic matter: Irradiation time
	dependent <i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment

xxxi

Figure 4.151.	UV-vis absorption spectra of organic matter: Irradiation time dependent	
	E. coli inactivation in WM solution upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment	229
Figure 4.152.	Synchronous scan fluorescence spectra of organic matter: Irradiation time	
	dependent E. coli inactivation in WM solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.230
Figure 4.153.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent <i>E. coli</i> inactivation in WM solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.230
Figure 4.154.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent <i>E. coli</i> inactivation in WM solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.231
Figure 4.155.	EEM fluorescence contour plots of organic matter: Irradiation time	
	dependent E. coli inactivation in WM solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.232
Figure 4.156.	UV-vis absorption spectra of organic matter: Irradiation time dependent	
	E. coli inactivation in HA solution upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment	234
Figure 4.157.	Synchronous scan fluorescence spectra of organic matter: Irradiation time	
	dependent E. coli inactivation in HA solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.235
Figure 4.158.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent <i>E. coli</i> inactivation in HA solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.236
Figure 4.159.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent E. coli inactivation in HA solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	.236

Figure 4.160.	EEM fluorescence contour plots of organic matter: Irradiation time	
	dependent E. coli inactivation in HA solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	238
Figure 4.161.	UV-vis absorption spectra of organic matter: Irradiation time dependent	
	E. coli inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment	240
Figure 4.162.	Synchronous scan fluorescence spectra of organic matter: Irradiation	
	time dependent E. coli inactivation in "HA and WM" solution upon	
	0.50% Fe-SynTiO ₂ solar photocatalytic treatment	240
Figure 4.163.	Specified UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent E. coli inactivation in "HA and WM" solution upon	
	0.50% Fe-SynTiO ₂ solar photocatalytic treatment	241
Figure 4.164.	Specific UV-vis and fluroscence parameters of organic matter: Irradiation	
	time dependent E. coli inactivation in "HA and WM" solution upon	
	0.50% Fe-SynTiO ₂ solar photocatalytic treatment	242
Figure 4.165.	EEM fluorescence contour plots of organic matter: Irradiation time dependent	
	E. coli inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment	244
Figure 4.166.	Initial, t=0, and t _{irr} =60 min total K, protein, and carbohydrate contents and	
	bacteria enumeration results of <i>E. coli</i> inactivation upon 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment in various aqueous matrixes	246
Figure 4.167.	Released protein and carbohydrate contents, and bacteria LRV of E. coli	
	inactivation upon 0.50% Fe-SynTiO2 solar photocatalytic treatment in	
	various aqueous matrixes	247
Figure 4.168.	Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI _{sync280} ,	
	UV_{280} , NPOC, and bacteria enumeration results of 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment of <i>E. coli</i> in various aqueous matrixes	248

Figure 4.169.	Irradiation time dependent E. coli inactivation in IsoT solution upon solar
	photocatalytic treatments performed in the presence of undoped and Fe-doped TiO ₂ specimens
Figure 4.170.	Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents of
	<i>E. coli</i> inactivation in IsoT solution upon solar photocatalytic treatments
	performed in the presence of bare and Fe-doped TiO ₂ specimens251
Figure 4.171.	Irradiation time dependent E. coli inactivation in WM solution upon solar
	photocatalytic treatments performed in the presence of undoped and Fe-doped
	TiO ₂ specimens
Figure 4.172.	Initial, t=0, and t _{irr} =60 min total K, protein, and carbohydrate contents of
	E. coli inactivation in WM solution upon solar photocatalytic treatments
	performed in the presence of bare and Fe-doped TiO ₂ specimens255
Figure 4.173.	Irradiation time dependent E. coli inactivation in HA solution upon solar
	photocatalytic treatments performed in the presence of undoped and Fe-doped
	TiO ₂ specimens
Figure 4.174.	Initial, t=0, and t _{irr} =60 min total K, protein, and carbohydrate contents of
	E. coli inactivation in HA solution upon solar photocatalytic treatments
	performed in the presence of bare and Fe-doped TiO ₂ specimens259
Figure 4.175.	Irradiation time dependent E. coli inactivation in "HA and WM" solution
	upon solar photocatalytic treatments performed in the presence of undoped
	and Fe-doped TiO ₂ specimens
Figure 4.176.	Initial, t=0, and t _{irr} =60 min total K, protein, and carbohydrate contents of
	E. coli inactivation in "HA and WM" solution upon solar photocatalytic
	treatments performed in the presence of bare and Fe-doped TiO ₂ specimens263
Figure 4.177.	Schematic diagram of LPS structure and positioning on Gram-negative
	bacteria cell wall

Figure B.1.	UV-vis absorption spectra of 100 kDa HA solution	299
Figure B.2.	Synchronous scan fluorescence spectra of 100 kDa HA solution	299
Figure B.3.	EEM fluorescence contour plot of 100 kDa HA solution	300
Figure C.1.	X-Ray Diffraction (XRD) diffractograms of undoped and Fe-doped photocatalyst specimens: a. TiO ₂ , b. 0.25% Fe-TiO ₂ , c. 0.50% Fe-TiO ₂ , d. SynTiO ₂ , e. 0.25% Fe-SynTiO ₂ , f. 0.50% Fe-SynTiO ₂	301
Figure C.2.	SEM micrographs and Energy-Dispersive X-Ray (EDX) spectra of undoped and Fe-doped photocatalyst specimens: a. TiO ₂ , b. 0.25% Fe-TiO ₂ , c. SynTiO ₂ , d. 0.50% Fe-SynTiO ₂	302
Figure D.1.	Calibration curve of bovine serum albumin (BSA) standard	303
Figure E.1.	Calibration curve of glucose standard	304

XXXV

LIST OF TABLES

Table 4.1.	Bacteria LRV in terms of percentage reduction45
Table 4.2.	Solar photolytic treatment of <i>E. coli</i> : Inactivation parameters47
Table 4.3.	Characteristic properties of organic matter present in solar photolytic treatment of <i>E. coli</i> in IsoT solution
Table 4.4.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in IsoT solution upon solar photolytic treatment
Table 4.5.	Characteristic properties of organic matter present in solar photolytic treatment of <i>E. coli</i> in WM solution
Table 4.6.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in WM solution upon solar photolytic treatment
Table 4.7.	Characteristic properties of organic matter present in solar photolytic treatment of <i>E. coli</i> in HA solution
Table 4.8.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in HA solution upon solar photolytic treatment
Table 4.9.	Characteristic properties of organic matter present in solar photolytic treatment of <i>E. coli</i> in "HA and WM" solution70
Table 4.10	D. Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon solar photolytic treatment
Table 4.11	. TiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation parameters78
Table 4.12	 Characteristic properties of organic matter present in TiO₂ solar photocatalytic treatment of <i>E. coli</i> in IsoT solution
Table 4.13.	Total K, protein, and carbohydrate contents of time dependent E. coli
-------------	---
	inactivation in IsoT solution upon TiO ₂ solar photocatalytic treatment84
Table 4.14.	Characteristic properties of organic matter present in TiO ₂ solar
	photocatalytic treatment of <i>E. coli</i> in WM solution
Table 4.15.	Total K, protein, and carbohydrate contents of time dependent E. coli
	inactivation in WM solution upon TiO ₂ solar photocatalytic treatment90
Table 4.16.	Characteristic properties of organic matter present in TiO ₂ solar
	photocatalytic treatment of <i>E. coli</i> in HA solution94
Table 4.17.	Total K, protein, and carbohydrate content of time dependent E. coli
	inactivation in HA solution upon TiO ₂ solar photocatalytic treatment97
Table 4.18.	Characteristic properties of organic matter present in TiO ₂ solar
	photocatalytic treatment of <i>E. coli</i> in "HA and WM" solution101
Table 4.19.	Total K, protein, and carbohydrate contents of time dependent E. coli
	inactivation in "HA and WM" solution upon TiO ₂ solar photocatalytic treatment
Table 4.20.	SynTiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation parameters109
Table 4.21.	Characteristic properties of organic matter present in SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in IsoT solution112
Table 4.22.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i>
	inactivation in IsoT solution upon SynTiO ₂ solar photocatalytic treatment114
Table 4.23.	Characteristic properties of organic matter present in SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in WM solution118
Table 4.24.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i>
	inactivation in wive solution upon Syn 11 O_2 solar photocatalytic treatment120

	٠	٠	٠
XXXV	1	1	1

Table 4.25.	Characteristic properties of organic matter present in SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in HA solution124
Table 4.26.	Total K, protein, and carbohydrate content of time dependent E. coli
	inactivation in HA solution upon SynTiO ₂ solar photocatalytic treatment126
Table 4.27.	Characteristic properties of organic matter present in SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in "HA and WM" solution130
Table 4.28.	Total K, protein, and carbohydrate content of time dependent E. coli inactivation
	in "HA and WM" solution upon SynTiO ₂ solar photocatalytic treatment132
Table 4.29.	0.25% Fe-TiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation parameters137
Table 4.30.	Characteristic properties of organic matter present in 0.25% Fe-TiO ₂
	solar photocatalytic treatment of <i>E. coli</i> in IsoT solution141
Table 4.31.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivation
	in IsoT solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment143
Table 4.32.	Characteristic properties of organic matter present in 0.25% Fe-TiO ₂
	solar photolytic treatment of <i>E. coli</i> in WM solution147
Table 4.33.	Total K, protein, and carbohydrate content of time dependent E. coli inactivation
	in WM upon 0.25% Fe-TiO ₂ solar photocatalytic treatment149
Table 4.34.	Characteristic properties of organic matter present in 0.25% Fe-TiO ₂
	solar photolytic treatment of <i>E. coli</i> in HA solution153
Table 4.35.	Total K, protein, and carbohydrate content of time dependent E. coli inactivation
	in HA solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment155
Table 4.36.	Characteristic properties of organic matter present in 0.25% Fe-TiO ₂
	solar photolytic treatment of <i>E. coli</i> in "HA and WM" solution159

Table 4.37.	Total K, protein, and carbohydrate content of time dependent E. coli inactivation
	in "HA and WM" solution upon 0.25% Fe-TiO ₂ solar photocatalytic treatment161
Table 4.38.	0.50% Fe-TiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation parameters166
Table 4.39.	Characteristic properties of organic matter present in 0.50% Fe-TiO ₂ solar photolytic treatment of <i>E. coli</i> in IsoT solution170
Table 4.40.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in IsoT solution upon 0.50% Fe-TiO ₂ solar photocatalytic treatment172
Table 4.41.	Characteristic properties of organic matter present in 0.50% Fe-TiO ₂ solar photolytic treatment of <i>E. coli</i> in WM solution175
Table 4.42.	Total K, protein, and carbohydrate content of time dependent <i>E. coli</i> inactivation in WM solution upon 0.50% Fe-TiO ₂ solar photocatalytic treatment177
Table 4.43.	Characteristic properties of organic matter present in 0.50% Fe-TiO ₂ solar photolytic treatment of <i>E. coli</i> in HA solution181
Table 4.44.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in HA solution upon 0.50% Fe-TiO ₂ solar photocatalytic treatment
Table 4.45.	Characteristic properties of organic matter present in 0.50% Fe-TiO ₂ solar photolytic treatment of <i>E. coli</i> in "HA and WM" solution
Table 4.46.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i> inactivation in "HA and WM" solution upon 0.50% Fe-TiO ₂ solar photocatalytic treatment189
Table 4.47.	0.25% Fe-SynTiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation parameters
Table 4.48.	Characteristic properties of organic matter present in 0.25% Fe-SynTiO ₂ solar photolytic treatment of <i>E. coli</i> in IsoT solution198

Table 4.49.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivation
	in IsoT solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic treatment200
Table 4.50.	Characteristic properties of organic matter present in 0.25% Fe-SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in WM solution204
Table 4.51.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivation
	in WM solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic treatment206
Table 4.52.	Characteristic properties of organic matter present in 0.25% Fe-SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in HA solution210
Table 4.53.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivation
	in HA solution upon 0.25% Fe-SynTiO ₂ solar photocatalytic treatment212
Table 4.54.	Characteristic properties of organic matter present in 0.25% Fe-SynTiO ₂
	solar photolytic treatment of <i>E. coli</i> in "HA and WM" solution216
Table 4.55.	Total K, protein, and carbohydrate contents of time dependent <i>E. coli</i>
	inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO ₂ solar
	photocatalytic treatment
Table 4.56.	0.50% Fe-SynTiO ₂ solar photocatalytic treatment of <i>E. coli</i> : Inactivation
	parameters223
Table 4.57.	Characteristic properties of organic matter present in 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment of <i>E. coli</i> in IsoT solution226
Table 4.58.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivation
	in IsoT solution upon 0.50% Fe-SynTiO ₂ solar photocatalytic treatment228
Table 4.59.	Characteristic properties of organic matter present in 0.50% Fe-SynTiO ₂
	solar photocatalytic treatment of <i>E. coli</i> in WM solution231

Table 4.60. Total K, protein, and carbohydrate contents of time dependent E. coli inactivation

	in WM solution upon 0.50% Fe-SynTiO ₂ solar photocatalytic treatment	233
Table 4.61.	Characteristic properties of organic matter present in 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment of <i>E. coli</i> in HA solution	237
Table 4.62.	Total K, protein, and carbohydrate contents of time dependent E. coli inactivati	on
	in HA solution upon 0.50% Fe-SynTiO ₂ solar photocatalytic treatment	239
Table 4.63.	Characteristic properties of organic matter present in 0.50% Fe-SynTiO ₂	
	solar photocatalytic treatment of <i>E. coli</i> in "HA and WM" solution	243
Table 4.64.	Total K, protein, and carbohydrate content of time dependent E. coli	
	inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO ₂ solar	
	photocatalytic treatment	245
Table 4.65.	Solar photocatalytic treatment of E. coli in IsoT solution performed in	
	the presence of undoped and Fe-doped TiO ₂ specimens: Inactivation	
	parameters	249
Table 4.66.	Solar photocatalytic treatment of <i>E. coli</i> in WM solution performed in	
	the presence of undoped and Fe-doped TiO ₂ specimens: Inactivation	
	parameters	253
Table 4.67.	Solar photocatalytic treatment of E. coli in HA solution performed in	
	the presence of undoped and Fe-doped TiO ₂ specimens: Inactivation	
	parameters	257
Table 4.68.	Solar photocatalytic treatment of <i>E. coli</i> in "HA and WM" solution	
	performed in the presence of undoped and Fe-doped TiO ₂ specimens:	
	Inactivation parameters	261
Table A.1.	Composition of WM solution	298
Table B.1.	Specified and specific UV-vis and fluorescence spectroscopic properties and	
	NPOC of 100 kDa HA solution	299

xli

Table C.1.	Crystallite particle sizes, absorption wavelengths and band-gap energies (E_{bg})	
	of undoped and Fe-doped TiO ₂ specimens	.301
Table C.2.	BET surface areas of photocatalyst specimens	.301

LIST OF SYMBOLS / ABBREVIATIONS

Symbol	Explanation	Unit
°C	Celsius	
Ca ²⁺	Calcium ion	
Cl [_]	Chloride ion	
Cl	Chlorine radical	
Cl ₂	Molecular chlorine	
Cl ₂ •-	Dichloride radical anion	
cm	Centimeter	
CO ₃ ^{2–}	Carbonate ion	
CO ₃ •-	Carbonate radical	
СООН	Carboxyl group	
¹ DOM	Singlet DOM	
³ DOM	Triplet DOM	
Da	Daltons	
E _{bg}	Band-gap energy	eV
e ⁻	Electron	
e ⁻ _{CB}	Photogenerated electron	
Fe ²⁺	Iron (II) ion	
Fe ³⁺	Iron (III) ion	
FI _{sync}	Fluorescence intensity	
	in synchronous scan mode	A.U.
FI _{sync280}	Fluorescence intensity	
	in synchronous scan	
	mode at λ_{emis} =280 nm	A.U.
FI _{sync470}	Fluorescence intensity	
	in synchronous scan	
	mode at λ_{emis} =470 nm	A.U.
g	Gram	
h	Planck's constant, $6.63 \times 10^{-34} \text{ m}^2 \text{ kg/s}$	
h ⁺	Positive holes	
$h^+ v_B$	Photogenerated hole	
H^+	Hydrogen ion	

H_2O	Water	
H_2O_2	Hydrogen peroxide	
$H_2PO_4^-$	Dihydrogen phosphate ion	
H_2PO_4	Dihydrogen phosphate radical	
H_2SO_4	Sulfuric acid	
HCO ₃ ⁻	Bicarbonate ion	
HCO ₃ •	Bicarbonate radical	
HSO_4^-	Hydrogen sulfate ion	
HO ʻ	Hydroxyl radical	
HO ₂ •	Hydroperoxyl radical	
Io	Incident photon flux	W/m^2
K^+	Potassium ion	
kDa	Kilodalton	
λ	Wavelength	nm
Δλ	Bandwidth	nm
λ_{emis}	Emission wavelength	nm
λ_{exc}	Excitation wavelength	nm
λ_{max}	Maximum wavelength	nm
L	Liter	
Mg^{2+}	Magnesium ion	
m	Meter	
μL	Microliter	
μm	Micrometer	
μΜ	Micromolar	
mg	Milligram	
mL	Milliliter	
v	Frequency of the light	s^{-1}
Ν	Normality	
Na ⁺	Sodium ion	
NaCl	Sodium chloride	
nm	Nanometer	
NO	Nitric oxide radical	
NO_2^-	Nitrite ion	
NO ₂ •	Nitrogen dioxide radical	
NO ₂ •-	Nitrite radical anion	

NO_3^-	Nitrate ion	
NO ₃ •-	Nitrate radical ion	
No	Initial E. coli cell count	CFU/mL
N ₆₀	E. coli cell count at irradiation	
	time of 60 min	CFU/mL
•ox	Oxidizing radical species	
¹ O ₂	Singlet oxygen	
³ O ₂	Triplet oxygen	
O - -	Oxide radical anion	
O ₂	Oxygen	
O ₃	Ozone	
O (³ P)	Atomic oxygen	
O2 ^{•–}	Superoxide radical anion	
OH⁻	Hydroxyl ion	
pKa	p function of acid dissociation constant	
%	Percentage	
рН	Power of hydrogen	
pHzpc	Point of zero charge	
PO ₄ ³⁻	Phosphate ion	
R	First order rate	CFU/mL min
SFI _{sync}	Specific Fluorescence Intensity	
	in synchronous scan mode	
SFI _{sync280}	Specific fluorescence intensity	
	in synchronous scan mode	
	at λ_{emis} =280 nm	
SFI _{sync470}	Specific fluorescence intensity	
	in synchronous scan mode	
	at λ_{emis} =470 nm	
SO4 ²⁻	Sulfate ion	
SO4 ^{•-}	Sulfate radical	
t	Irradiation time	min
t _{1/2}	Half-life	min
TiO ₂	Titanium dioxide	
t _{irr}	Irradiation period	min
W	Watt	

Abbreviation	Explanation	
AOPs	Advanced Oxidation Processes	
ATCC	American Type Culture Collection	
ATP	Adenosine triphosphate	
ATPase	Adenosine triphosphatase	
BET	Brunauer-Emmett-Teller	
CAT	Catalase	
СВ	Conduction Band	
CW	Clockwise	
CCW	Counterclockwise	
CFU	Colony Forming Unit	
Color ₄₃₆	Absorbance at 436 nm	
DBPs	Disinfection by-products	
DNA	Deoxyribonucleic acid	
DOM	Dissolved Organic Matter	
DOC	Dissolved Organic Carbon	mg/L
DON	Dissolved Organic Nitrogen	
DOP	Dissolved Organic Phosphorus	
E. coli	Escherichia coli	CFU/mL
EDX	Energy-Dispersive X-Ray spectroscopy	
EEM	Excitation Emission Matrix	
EPS	Extracellular Polymeric Substances	
EU	Endotoxin Unit	
FA	Fulvic Acid	
FAAS	Flame Atomic Absorption Spectroscopy	
FI	Fluorescence Index	
HA	Humic Acid	
HSs	Humic Substances	
IC	Ion Chromatography	
IEP	Isoelectric point	
IHSS	International Humic Substance Society	
IsoT	Isotonic solution	
L. acidophilus	Lactobacillus acidophilus	
LAL	Limulus Amebocyte Lysate	
LB	Luria Bertani	

LPS	Lipopolysaccharide	
LRV	Logarithmic Reduction Value	%
MDA	Malonaldehyde	
min	Minute	
MW	Molecular Weight	
NADH	Nicotinamide adenine dinucleotide	
NADPH	Nicotinamide adenine dinucleotide phosphate	
NP	Nanoparticle	
NPOC	Non-Purgeable Organic Carbon	mg Org C/L
NOM	Natural Organic Matter	
POM	Particulate Organic Matter	
ROS	Reactive Oxygen Species	
S. cerevisiae	Saccharomyces cerevisiae	
SCoA ₄₃₆	Specific Color Absorbance at 436 nm	L/m mg
SEM	Scanning Electron Microscopy	
SFI	Specific Fluorescence Intensity	
SOD	Superoxide dismutase	
SODIS	Solar water disinfection	
S. typhimurium	Salmonella typhimurium	
SUVA ₂₅₄	Specific UV Absorbance at 254 nm	L/m mg
SUVA ₂₈₀	Specific UV Absorbance at 280 nm	L/m mg
SUVA ₃₆₅	Specific UV Absorbance at 365 nm	L/m mg
SynTiO ₂	Synthesized TiO ₂	
TEM	Transmission Electron Microscopy	
TOC	Total Organic Carbon	
ТОМ	Total Organic Matter	
UV	Ultraviolet	
UV-A	Ultraviolet A	
UV-B	Ultraviolet B	
UV-C	Ultraviolet C	
UV ₂₅₄	Absorbance at 254 nm	
UV ₂₈₀	Absorbance at 280 nm	
UV ₃₆₅	Absorbance at 365 nm	
UV-DRS	Ultraviolet Diffuse Reflectance Spectroscopy	
UV-vis	Ultraviolet-visible	

xlvii

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1. INTRODUCTION

With climate change and increased global population, many people will suffer from lack of clean freshwater which is one of the biggest problems throughout the world. The challenges with water are expected to go worse in following years due to globally occurring water scarcity (Montgomery and Elimelech, 2007). According to UNICEF, 38 % of the world's population, suffered from lack of proper sanitation facilities whereas almost one billion people continue to use unsafe drinking water sources. Especially, in Sub-Saharan Africa, 42 % of people still use unimproved water sources (UNICEF, 2009). No access to good quality drinking water causes a high risk of waterborne diseases like diarrhea, cholera, typhoid fever, hepatitis A, amoebic and bacillary dysentery, and others (WHO, 1999).

Due to the absence of clean drinking water around the world, disinfection of water by the application of different treatment methods has been widely investigated (Helali et al., 2014). Water disinfection is traditionally achieved by using chlorination and ozonation processes. However, the usage of chlorine leads to the formation of disinfection byproducts (DBPs) that have adverse effects on human health and environment by reacting with dissolved organic matter (DOM) in water (Nieuwenhuijsen et al., 2000). Ozonation also causes the formation of organic DBPs like aldehydes, carboxylic acids, ketones and bromates (Huang et al., 2005). However, ozonation process requires more complex technologies than chlorination (Sichel et al, 2007). On the other hand, water disinfection using natural and/or artificial sunlight have been widely studied for the inactivation of a wide range of microorganisms (Busse et al., 2019; Carratala et al., 2016; Mattle et al., 2015; Vivar et al., 2017). Solar water disinfection (SODIS), which is a simple, inexpensive method for the household water treatment, relies on the combined effect of ultraviolet (UV) radiation and thermal heating of solar light. Since 1980s, studies have been conducted to investigate the mechanism of solar disinfection and to develop new technologies to make this technique faster and safer (McGuigan et al., 2012). Literature suggests that SODIS in natural water is in reality an in-situ generated photocatalytic process promoted by solar light (Pulgarin, 2015).

The usage of TiO₂-UV bactericidal activity to decontaminate water was first demonstrated by Matsunaga and colleagues (Matsunaga et al., 1985). Since then, photocatalytic disinfection of microorganisms gained an increasing interest and many review papers are available in literature covering background information as well as recent developments in the field (Bekbolet, 2007; Byrne

et al., 2015; Gupta and Modak, 2020; Helali et al., 2014; Reddy et al., 2017; Uyguner-Demirel et al., 2017, 2018; Vale et al., 2016; Wang et al., 2015).

Solar photocatalytic disinfection studies of *E. coli* as the indicator microorganism in the presence/absence of DOM using bare TiO₂ and visible light activated (VLA) doped TiO₂ (*i.e.*, N-doped, Fe-doped, Se-doped, and Se-N co-doped TiO₂) has been carried out so far by Bekbolet and collegues (Bekbolet and Araz, 1996; Bekbolet, 1997, 2007; Birben et al., 2017a, 2017b, 2021b).

Complimentary to previous research, the main objective of the present study is to evaluate and propose heterogeneous photocatalysis using bare and Fe-doped TiO₂ under solar light for destruction of indicator bacteria. Considering its extensive use in literature and for comparison purposes, the pure cultures of the fecal indicator bacterium *E. coli* were used in photolytic and photocatalytic disinfection studies. Inactivation of *E. coli* was considered under identical solar irradiation conditions. Considering the real case in aquatic systems, effect of DOM on solar light initiated *E. coli* inactivation was evaluated with the use of humic acid (HA) as a DOM analogue in natural waters. Besides ultraviolet-visible (UV-vis) and fluorescence spectroscopic analysis to characterize DOM under photo-initiated reaction conditions, destruction of *E. coli* was followed by released cell components like proteins, carbohydrates, and total potassium content. Further evaluation of released cytoplasmic organic content was followed by the occurrence of fluorophores of microbial by-products and protein-like fluorophores as presented in excitation emission matrix (EEM) contour plots. Moreover, absence or presence of endotoxin was also investigated.

2. LITERATURE REVIEW

2.1. Solar Disinfection of Water

The usage of sunlight for water purification is one of the oldest methods, communities from the Indian sub-continent placed their drinking water in open trays to be "blessed" by the sun nearly 2000 years ago. The first systematic study about the bactericidal effect of sunlight was investigated by Downes and Blunt in 1877. They showed that short wavelength solar radiation has the greatest antimicrobial effect (Downes and Blunt, 1877a, 1877b, 1878). After a century, Acra and colleagues suggested the practical application of sunlight for the disinfection of oral rehydration solutions and drinking water and this process was generally named as solar disinfection (Acra et al., 1980, 1984, 1989). Also, they contemplated that the solar disinfection can be used as an alternative, inexpensive, sustainable and simple water treatment method by people who had no access to other water treatment systems. After these findings, bacterial inactivation potential of solar disinfection namely SODIS has been investigated in detail by several groups (Conroy et al., 1996; Joyce et al., 1992, 1996; Kehoe et al., 2001; Lawand et al., 1988; McGuigan et al., 1998, 1999; Reed, 1997; Sommer et al., 1997; Wegelin et al; 1994; Wegelin and Sommer, 1998).

2.2. Mechanism of Solar Disinfection

The solar irradiance incident on the Earth's outer atmosphere is almost around 1360 W/m². However, received irradiance at the ground level on a horizontal surface of the equator in summer is 1120 W/m^2 . This decrease is caused from the absorption of the irradiance by the water vapor, carbon dioxide, oxygen, ozone and pollutants in the atmosphere (McGuigan et al., 2012). Solar light contains infrared radiation, visible light and ultraviolet radiation. Visible light and infrared light make up approximately 95 % and UV light generates 5 % of the solar light that reaches the Earth's atmosphere. The wavelength ranges of UV light are: 400 to 315 nm for UV-A; 315 to 280 nm for UV-B and 280 to 100 nm for UV-C light (IARC, 2012).

Natural waters can contain a variety of microorganisms such as bacteria, viruses, fungi, protozoa, and algae. The removal or inactivation of pathogenic bacteria in drinking water has high importance to prevent the transmission of waterborne diseases. Due to difficulty and cost of individual detection of pathogenic microorganisms, particular indicator organisms are used to test drinking water safety (Bekbolet, 2007; Tortora et al., 2018). Coliform bacteria were first used as indicator organisms by

Phelps (1909). Moreover, *E. coli*, which is a fecal coliform bacterium, has been widely used as an indicator organism to detect drinking water contamination. Bacteria destruction in solar disinfection process could be formed in two main ways as direct and indirect damage. When genetic material of the organism, namely deoxyribonucleic acid (DNA), is irradiated by UV light, especially UV-B, direct damage of DNA occurs (Halliwell and Gutteridge, 1989). Although UV-A irradiation is not as effective as UV-B to directly damage DNA, it plays an important role on the inactivation of bacteria by causing indirect damage. Indirect damage occurs by the absorption of UV wavelengths via photosensitizers. When photosensitizers are excited, they lead to the formation of reactive oxygen species (ROS) in water such as singlet oxygen, superoxide, hydrogen peroxide, and hydroxyl radical (Cooper et al., 1988a; Foyer et al., 1994).

Photosensitizers could be endogenous (internal) ones such as porphyrins, flavins, quinones, nicotinamide adenine dinucleotide/nicotinamide adenine dinucleotide phosphate (NADH/NADPH), cytochromes, photosynthetic pigments etc. (Curtis et al., 1992; Eisenstark, 1987; Foyer et al., 1994; Jagger, 1981; Sammatro and Tuveson, 1987; Tuveson and Sammartano, 1986) and exogenous (external) ones which are present in surface waters like humic compounds and chlorophylls (Blough and Zepp, 1995; Cooper et al., 1988b; Voelker et al., 1997). ROS generated by solar irradiation react with the cell membrane, membrane lipids, proteins and DNA and cause inactivation of the cell (Figure 2.1) (Bose and Chatterjee, 1995; Futsaether et al., 1995; Gourmelon et al., 1994). When ROS are formed, they cause a series of oxidation reactions as of polyunsaturated fatty acids in lipids, amino acids in proteins and oxidative damage to DNA (Reed et al., 2000). This DNA damage can occur via the strand breakage, which is fatal for the microorganism, and base changes, which can cause some mutagenic effects like blockage of replication (McGuigan et al, 2012).



Figure 2.1. Proposed schematic process of cell destruction (Adhikari et al., 2015).

Other than the effect of UV light, increased temperature also plays an important role in SODIS. The temperature of the water increases to a point that bacteria are inactivated by the absorption of solar light, especially the solar infrared radiation and this thermal inactivation process is named as solar pasteurization (Wegelin and Sommer, 1998). In addition, when the bactericidal effect of UV light combines with thermal inactivation process, a synergistic effect has been observed at the water temperature exceeding 45°C (McGuigan et al., 1998). Researchers suggested that increased water temperatures occurring in the solar pasteurization process inhibits DNA repair mechanisms, therefore, synergistic effect can be more effective (McGuigan et al., 1998).

The main disadvantage of photoinactivation is attributed to microbial regrowth and not being effective in case of bacterial strains such as Gram-negative *S. typhimurium*. Therefore, research was directed to the application of photocatalysis as an alternative.

2.3. Heterogeneous Photocatalysis for the Enhancement of Solar Disinfection Efficiency

Advanced oxidation processes (AOPs) were established by Glaze and colleagues and defined as treatment processes which form hydroxyl radicals (HO[•]) in sufficient quantity to oxidize organic pollutants and microorganisms in water (Glaze et al., 1987). In general, AOPs could be classified in two major groups as homogeneous and heterogeneous AOPs. In homogeneous systems such as photo-Fenton, O₃, O₃/UV, O₃/H₂O₂, H₂O₂/UV, reaction takes part in liquid phase however in heterogenous systems like TiO₂/O₂/UV and TiO₂/H₂O₂/UV, catalysts are not dissolved in liquid, and reaction occurs in liquid-solid phase (Buthiyappan et al., 2016; Mansilla et al., 1997; Centi and Perathoner, 2005; Uyguner and Bekbolet, 2007). In the last few decades, main interest has been directed to heterogeneous photocatalysis which is a promising AOP in various photochemical applications consisting of water splitting, degradation of organic compounds, reduction of CO₂ and water disinfection (Carey et al., 1976; Fujishima and Honda, 1972; Frank and Bard, 1977; Matsunaga, 1985; Matsunaga et al., 1985).

According to IUPAC, photocatalysis is defined as the change in the rate of a chemical reaction or its initiation under the action of UV, visible or infrared radiation in the presence of a photocatalyst which absorbs light and also involves in the chemical transformation of the reaction partners (IUPAC, 2004). One of the earliest studies about TiO₂ and UV light's bactericidal activity was demonstrated in 1985 by Matsunaga and colleagues to inactivate *E. coli*, *L. acidophilus* and *S. cerevisiae* cells covering also a mechanistic explanation (Matsunaga et al., 1985). TiO₂ is the most widely used semiconductor for water treatment processes since its excitation by wavelengths shorter than 390 nm (UV-A) generates ROS, especially HO[•] (Harper et al., 2001; Ollis et al., 1991).

2.3.1. Photocatalytic Oxidation Processes

When a semiconductor like TiO_2 is irradiated by a suitable wavelength, the photocatalytic oxidation processes occur. Throughout the irradiation of TiO_2 , the electrons (e⁻) are excited into the conduction band (CB) which leaves a hole (h⁺) in the valence band (VB) of TiO_2 . The charge carriers (e⁻/h⁺ pairs) migrate to the surface/interface of TiO_2 and take part in redox reactions to form highly active ROS like HO[•], superoxide radical anion (O2^{•-}), hydroperoxyl radical (HO2[•]) and hydrogen peroxide (H₂O₂) (Equations 2.1-2.8) (Rengifo-Herrera et al., 2013).

$$TiO_2 + h\nu \rightarrow TiO_2 (e^-_{CB} + h^+_{VB})$$
(2.1)

$$TiO_2 (h^+ VB) + H_2O_{ads} \rightarrow TiO_2 + HO^{\bullet}_{ads} + H^+$$
(2.2)

$$TiO_{2}(h^{+}_{VB}) + OH^{-}_{ads} \rightarrow TiO_{2} + HO^{\bullet}$$

$$TiO_{2}(h^{-}_{VB}) + O_{2}(h^{-}_{VB}) + O_{2}(h^{-}_{VB}) + O_{2}(h^{-}_{VB})$$

$$(2.3)$$

$$\Gamma_1 O_2 (e^-_{CB}) + O_2 \rightarrow \Gamma_1 O_2 + O_2^{-2}$$

$$(2.4)$$

$$O_2^{\bullet-} + H^+ \leftrightarrow HO_2^{\bullet} \qquad pK_a = 4.8 \qquad (2.5)$$

$$HO_2^{\bullet} + O_2^{\bullet-} + H^+ \rightarrow H_2O_2 + O_2$$

$$(2.6)$$

$$2HO_2 \rightarrow H_2O_2 + O_2 \tag{2.7}$$

$$TiO_2 (e_{CB}) + H_2O_2 \rightarrow TiO_2 + OH^- + HO^{\bullet}$$

$$(2.8)$$

2.3.2. Bare and Visible Light Activated Doped TiO₂ Species

TiO₂ P-25 (Evonik) which is one of the most widely used photocatalyst, mainly contains free nanoparticle (NP) with single anatase (80 %) and rutile (20 %) crystal structure (Jiang et al., 2017). Although both of them are equally detrimental to bacteria (Simon-Deckers et al., 2009), anatase crystal phase shows stronger photocatalytic properties in comparison to rutile phase (Joost et al., 2015). Isoelectric point (IEP) or point of zero charge (pH_{zpc}) of TiO₂ equals to pH=6.25 (Birben et al., 2015). Its pH dependent positively and negatively charged surface-active sites are shown in Figure 2.2. Surface groups of a metal oxide like semiconductor TiO₂ express amphoteric properties and the surface acid-base equilibria at 25°C can be displayed by the following equations (Equations 2.9 and 2.10) (Kormann et al., 1991).

$$TiOH_{2^{+}} \leftrightarrow TiOH + H^{+} \qquad pK_{a1} = 2.4$$

$$TiOH \leftrightarrow TiO^{-} + H^{+} \qquad pK_{a2} = 8.0$$

$$(2.9)$$

$$(2.10)$$



Figure 2.2. a. Formation of surface hydroxylated species of TiO₂ according to pH (Dalrymple and Goswami, 2017); b. pH dependent electrostatic interactions on TiO₂ surface (Beranek, 2011).

Bare TiO₂ has a wide band-gap energy, as electromagnetic band-gap (E_{bg}) = 3.2 eV, that cause some drawbacks on photocatalytic process such as; restriction of light absorption to only UV region which is approximately 5 % of solar spectrum, rapid charge recombination between electron hole pairs and low surface coverage of photocatalyst particles causing slower degradation rates of organic compounds (K11c and Cinar, 2009). The effectiveness of bare TiO₂ can be increased by doping it with metal elements or metallic salts such as iron, silver, copper, tungsten and ruthenium (Dobosz and Sobczyński, 2003; Fung et al., 2003; Koli et al., 2016; Lettmann et al., 2001; Li et al., 2001; Sangchay and Ubonchonlakat, 2015; Sun et al., 2003; Yu et al., 2003) or non-metals like sulphur (Yu et al., 2005) or by coating the surface of TiO₂ with photosensitizing dyes (Cho et al., 2001; Lobedank et al., 1997). TiO₂ doping process forms VLA TiO₂ by shifting the light absorption range of bare TiO₂ from UV to visible light. Doped photocatalyst specimens could be used to improve solar light harvesting in TiO₂ photocatalytic process and improve the photocatalytic activity. Doping TiO₂ with a transition metal like Fe increases the activity of TiO₂ with two main ways; *i.* shifting light absorption to the visible light by narrowing the band-gap and creating additional electronic states within TiO₂ band-gap which could be induced by lower energy light, and *ii.* increased HO[•] formation by reducing electron hole pair recombination speed (Yalcin et al., 2010). Fe and Se-doped TiO₂ specimens were studied in detail by Bekbolet and colleagues (Birben et al., 2015, 2017a, 2017b). Fe-doped TiO₂ specimens were prepared by using TiO₂ P-25 and TiO₂ Hombikat UV-100 by wet impregnation method. Humic acid sample was selected as a specific molecular size fraction as < 100 kDa. Following treatment, humic molecular size fractions were further characterized by UV-vis and advanced fluorescence techniques such as EEM. Furthermore, VLA TiO₂ specimens were selected as Se-doped and Se-N co-doped TiO₂. Solar photocatalytic inactivation kinetics, organic matter degradation was also followed and characterized by UV-vis and EEM (Bekbolet, 1997, 2020; Bekbolet and Araz, 1996; Bekbolet and Sen-Kavurmaci, 2015; Bekbolet and Turkten, 2020; Birben et al., 2015, 2017a, 2017b; Uyguner-Demirel and Bekbolet, 2011; Sen-Kavurmaci and Bekbolet, 2014).

2.4. Indicator Organism: Escherichia coli

As the main indicator organism, *E. coli* is a prokaryotic, single-celled organism lacking a distinct nucleus and membrane enclosed organelles (Figure 2.3).

Taxonomy of <i>E. coli</i> :	Domain: Bacteria
	Phylum: Proteobacteria
	Class: Gammaproteobacteria
	Order: Enterobacteriales
	Family: Enterobacteriaceae
	Genus: Escherichia
	Species: Escherichia coli

Belonging to the main domain of Bacteria and phylum Proteobacteria *E. coli* is from Enterobacteriaceae or enteric family which includes facultative anaerobe microorganisms. As a member of *Escherichia* genus, rod shaped *E. coli* is recognized as Gram-negative bacteria. Motile forms of Enterobacteriaceae have peritrichous multi-flagella like *E. coli* (Hogg, 2005; Tortora et al., 2018). It is a neutrophilic bacterium maintaining its growth between almost at neutral conditions as pH: 6-8 (Ingraham, 1999).



Figure 2.3. The general structure of a prokaryotic organism. Red-labeled structures are found in all bacteria species, while black-labeled ones are not (Tortora et al., 2018).

Approximately 70 % of an *E. coli* cell is made up of water. Remaining part of the bacterium biomass proportions are protein, 50-60 %; RNA, 13-25 %; polysaccharide, 9-17 %; lipid, 8-9 %; and DNA, 2-4 % per gram dry weight (Sucher et al., 2012). Cells are 1.0-2.0 µm long and 0.5 to 1.0 µm in diameter; and volume and surface area of an *E. coli* is nearly 1.0 μ m³ and 6.0 μ m², respectively (Phillips et al., 2013). C, O, H, N, S, P, K, Mg, Ca, Na, Cl and Fe are the macronutrients that E. coli needs to maintain its growth (Dworkin et al., 2006). Also, E. coli can be represented with a chemical formula of CH_{1.74}O_{0.34}N_{0.22} (Duboc et al., 1999). It is an important pathogen founding in soil, plants, animal respiratory and intestinal tracts. The presence of E. coli in water and food indicates fecal contamination. Due to its high numbered permanent presence in human feces, ability of survival in water and simple detectability, it is known as an indicator organism. The usual indicator organisms are coliform bacteria defined as Gram-negative, aerobic or facultative anaerobic, non-endospore forming, rod-shaped bacteria that ferment lactose to form gas in 48 hours after being placed in lactose broth at 35 °C. E. coli is the dominant fecal coliform bacteria because it generates a large portion of human intestinal population. E. coli is normally not disease causing, although some strains of it can cause gastroenteritis, traveler's diarrhea, in developing countries infant diarrhea and urinary tract infections. The best-known and most infectious type of *E. coli* is serotype O157:H7 (Tortora et al., 2018).

2.4.1. E. coli Motility

Most of the bacteria can actively move in liquid media by itself and this movement is called as motility. Flagellar motility requires energy to rotate flagellum from the basal body and this motion could be provided by one or more flagella. Bacteria can modify the speed of the movement and change the rotation direction of the flagella (Singleton, 1997; Tortora et al., 2018). *E. coli* has about 10 flagella arising from different points of the cell surface and this type of positioning of flagella is called as peritrichous (Hogg, 2005; Wilson et al., 2002). Peritrichous flagella of *E. coli* rotate independently from each other (Macnab and Han, 1983). Most of the time (95 %) each flagellum rotates counterclockwise (CCW) and rest of the time clockwise (CW). When most of the flagella rotate CCW, they pile up in a helical bundle at one end of the cell and bacterium moves toward one direction, the movement is called as "run" or "swim" movement. Run movements are periodically interrupted about one per second by directionless "tumbles". Changed direction of flagellar rotation causes tumbles; it occurs randomly (Figures 2.4a and 2.4b). Flagellar rotation determines the type of the movement (Singleton, 1997; Tortora et al., 2018). Consecutive running and tumbling movements cause the formation of a three dimensional "random walk" similar to Brownian motion of electrons. (Figure 2.4c) (Singleton, 1997; Macnab, 1979).



Figure 2.4. a. CCW, CW flagella rotations and movement of *E. coli* (Butler and Camilli, 2005). b. Run and tumble movements of a bacterium. Blue arrows indicate the flagellar rotation direction and gray arrows indicate the movement direction (Tortora et al., 2018). c. Random walk pattern of bacteria (Phillips et al., 2013).

Growth of flagella occurs from the distal tip by polymerization of protein molecules (Rosenbaum and Child, 1967). Naturally or mechanically displaced flagella can regenerate themselves (Levy, 1974). Furthermore, researchers showed that flagellar assembly of bacteria did not detach with the effect of photocatalytic disinfection process. However, after recurrent photocatalytic treatment, genes taking place in the expression of flagellar assembly are downregulated (Zhang et al., 2019). Therefore, it can be concluded that the protein content of the experiment solution could not increase due to separation of flagella.

2.4.2. Surface Properties and Point of Zero Charge of E. coli

Bacterial cells have a net negative charge under the influence of surface components (Corpe, 1970). Dissociation of the charged functional groups founding in the cell wall causes this electronegative characteristic. Neihof and Echols showed that carboxyl groups on the surface of *E. coli* cells are the source of electronegativity (Neihof and Echols, 1973). These carboxylic groups of peptidoglycans are located on the D-glumatic acid, meso-diaminopimelic acid and terminal D-alanine residues of the peptide groups. Due to the presence of the electronegative groups, the cell surface of the bacteria is negatively charged at neutral pH. Furthermore, the presence of amino groups in the peptidoglycan layer and counter-ions from the environment neutralize the negative charges and prevent shreding of the cell wall due to electrostatic repulsion of the negative charges (Thwaites and Mendelson, 1991).

With regard to basic definition of zeta potential as the electric charge at the shear plane, is generally used as a surrogate parameter to explain surface charge of bacteria cells. Proteins and metabolites which are present in the growth medium can be adsorbed to the bacteria cell surface via various electrostatic interactions as well as by *van der Waals* attractions and London forces. These absorbed structures can form layers on the cell wall. Therefore, the bacteria cell wall contains both the cell wall charge and the effect of the formed layer. It was reported that pH_{zpc} of *E. coli* (1.5-3) grown in different growth mediums like Luria Bertani (LB) Lennox, LB Lennox with 2 mg/L glucose and buffered LB Lennox demonstrated similar values with each other (Ng, 2018). Additionally, research investigating the effect of different wash buffers (such as 0.154M NaCl, 0.001M KCl, and 0.1M NaNO₃) on pH_{zpc} of *E. coli* showed that there was no significant effect of the buffers against the pH_{zpc} of the cell surface (Ng and Ting, 2017). The attained results significantly indicated that neither different growth media nor wash buffers substantially exert an effect on the pH_{zpc} of *E. coli* in aqueous medium. Therefore, under almost neutral conditions (pH \approx 6-7), *E. coli* cell surface would be regarded as negatively charged.

2.4.3. E. coli Inactivation Mechanism During Solar Light Initiated Photocatalytic Processes

In photocatalytic inactivation processes in situ generated ROSs attack bacteria cell components, starting with the cell capsule. When ROSs attack the cell capsule, damage lipid contents of the cell membrane and cause increased membrane permeability resulting in the leakage of K^+ , proteins and DNA of the bacteria. Increased ROSs concentration at the periphery and inside the cell could cause inactivation of enzymes and give oxidative damage to DNA and proteins (An et al., 2017).

During exposure of stress like osmotic, heat, or oxidative, bacteria cells may continue their metabolic activities but stop growing. As a consequence of these type of stress factors, bacteria cells start to produce excessive amount of free radicals that could be lethal for the cell and this process is called as "suicide response" (Overman, 2006). Bacteria in exponential phase are more sensitive to suicide response than stationary phase (Aldsworth et al., 1999).

2.4.3.1. Effect of reactive oxygen species on solar light initiated photocatalytic bacteria destruction. ROS not only destroy the chemical contaminants in water but also cause fatal damage to bacteria via oxidation of cellular components, membrane leakage of the cell wall, etc. (Blake et al., 1999; Cabiscol et al., 2000; Reddy et al., 2016). Hydroxyl radicals are very active and primarily responsible for the disinfection of bacteria (Fujishima et al., 2000; Reddy et al., 2017). The oxidative effects of TiO_2 on bacteria occur via direct contact of bacteria with the catalyst (Figure 2.5), therefore the first damage of bacteria occurs at the outer membrane (Figures 2.6 and 2.7a) (Stage A) (Reddy et al., 2017; Sunada et al., 2003). After the first damage on the outer membrane, the fatal damage can occur by the destruction of DNA (Stage B) (McGuigan et al., 2012; Reddy et al., 2017). It was also demonstrated that cell wall damages take place prior to cytoplasmic membrane damage.



Figure 2.5. Transmission electron microscopy (TEM) image of *E. coli* with TiO₂ particles attached on it (Dalrymple and Goswami, 2017).



Figure 2.6. Possible mechanism of photocatalytic disinfection of bacteria (Reddy et al., 2017).



Figure 2.7. a. Structural membrane composition and possible photocatalytic disinfection mechanism, b. Lipid peroxidation and membrane leakage process in Gram-negative bacteria e.g. *E. coli* (Reddy et al., 2017).

The bactericidal inactivation processes generally occur in following stages; *i*. Cytoplasmic membrane destruction, *ii*. Increase in cell permeability, *iii*. Release of intracellular components, *iv*. Loss of viability, *v*. Distortion of cellular surface, *vi*. Change in overall shape and size. Increase in cell membrane permeability (Figure 2.7b) could be related to the increase in the release of protein, carbohydrate, and K⁺ (Carré et al., 2014; Ren et al., 2009). Loss of viability could be related to the photooxidation of co-enzyme A and it causes the inhibition of respiration (Reddy et al., 2017).

2.4.3.2. Cellular targets of reactive oxygen species.

Bacterial capsule as extracellular polymeric substances. Extracellular polymeric substances (EPS) or glycocalyx are produced and secreted by bacteria which are composed of polysaccharides and polypeptides (Decho, 2011). EPS are located outside of the cell wall and help bacteria to maintain moisture, store nutrients, and attach to surfaces and EPS attached tightly on the cell wall is called capsule (Eboigbodin and Biggs, 2008; Tortora et al., 2018). EPS provide contact between bacteria and photocatalyst and protect the cell from ROS attack (Hessler et al., 2012). Researchers showed that bacteria without capsule are more sensitive to photocatalytic inactivation than the capsulated ones (Liu et al., 2007). Also, bacteria that have direct contact with TiO₂ have high sensitivity to photocatalytic inactivation. When bacteria–TiO₂ direct contact is inhibited, the amount of EPS increases which can result in higher resistance capacity to treatment (Huang et al, 2015).

<u>Cell wall and cell membrane.</u> Bacteria cell envelope consists of plasma/cell membrane and cell wall (Figure 2.8). Cell envelope provides stable inner conditions to the cell and prevent unwanted materials from entering the cell.



Figure 2.8. Cell wall and cytoplasmic membrane structure of Gram-negative bacteria (a. Madigan and Martinko, 2006; b. Tortora et al., 2018).

Unlike Gram-positive bacteria, cell wall of Gram-negative bacteria has a thinner peptidoglycan layer and an outer membrane outside this peptidoglycan layer (An et al., 2017; Dalrymple et al., 2010; Tortora et al., 2018). Peptidoglycan layer composed of peptide cross-linked polysaccharide in Gram-negative bacteria generates 10 % of the cell wall (1-2 nm). It provides the shape and internal pressure of the cell and allows passage of particles of 2 nm size through due to its porous structure (Dalrymple and Goswami, 2017; Demchick and Koch, 2006). Outer membrane of Gram-negative bacteria includes lipopolysaccharides (LPS), lipoproteins and phospholipids in its structure. LPS consists of three components as; O polysaccharide, core polysaccharide, and lipid A which released from cell wall after bacteria death and functions as an endotoxin. Moreover, outer membrane has porin proteins allowing to enter some nutrients into the cell, such as disaccharides, peptides, amino acids, nucleotides, vitamin B₁₂, and iron (Tortora et al., 2018).

Bacterial envelope damage may form as a result of lipid peroxidation process induced by the oxidative stress (Figure 2.9) (Dalrymple et al., 2010, Singh et al., 2013). ROS attack to unsaturated lipids and start a chain reaction. Phospholipids constitute most part of the outer membrane in *E. coli* and due to their amplitude and convenience to oxidation they are the first targets to HO[•] attacks (Dalrymple and Goswami, 2017).



Figure 2.9. Schematic diagram of ROS utilized lipid peroxidation reaction (Dalrymple et al., 2011).

Lipid peroxidation occurs in three steps. The first step is initiation, HO[•] takes a H atom from an unsaturated fatty acid and forms a lipid radical. The second step is propagation, the carbon centered lipid radical reacts with an oxygen and generates a lipid peroxyl radical, and then newly formed radical takes one H atom from another unsaturated fatty acid that forms a lipid peroxide molecule and leaves another lipid radical. Generated lipid peroxide degrades into malonaldehyde (MDA) that is

usually used as an indicator of lipid peroxidation. The third step is termination that occurs by the formation of neutral products from two radicals. Because lipid peroxidation is a chain reaction, termination reaction can end far from the first unsaturated fatty acid in the initiation reaction (Dalrymple and Goswami, 2017; Sun et al., 2017).

As a result of bacterial damage, leakage of intracellular components like K^+ could be detected. The main functions of K^+ in bacteria are to sustain membrane potential, osmotic pressure and to conduct signals. Some enzymes those take part in protein synthesis require potassium, as well. In unharmed cells, high intracellular K^+ concentration is provided with Na⁺- K^+ pumps to maintain turgor pressure of the cell (Wang and O'Doherty, 2012). The leakage of intracellular components indicates that bacterial cell wall has been damaged and cell membrane permeability has increased. Moreover, relative permeability of membranes are 10 times higher to K^+ than Na⁺ (Brock et al., 2000). Other than K^+ leakage, endotoxin (an outer membrane component of *E. coli*, Lipid A) release is another sign of the outer membrane damage in *E. coli*. Lu and colleagues reported that ~5 nm TiO₂ particles penetrate *E. coli* cell only after 20 min of photocatalytic treatment and this finding supports that the destruction of cell wall and leakage of macromolecules from the cell occurs during the treatment (Lu et al., 2003). In addition, it has been reported that in the presence of TiO₂, K^+ leakage starts immediately after illumination and K^+ concentration shows an increase up to a stable point. Researchers' findings show a consistency between K^+ leakage and the loss of cell viability (Saito et al., 1992).

During the photocatalytic inactivation process bacterial envelop is damaged and cell membrane permeability is increased. It could also be associated with the disruption of membrane-associated proteins. Researchers found that, respiratory chain membrane proteins and the adenosine triphosphatase (ATPase), which play a crucial role to transform energy into adenosine triphosphate (ATP), were inactivated rapidly and caused inadequate energy to maintain membrane unity (Bosshard et al., 2010a).

Research showed that respiration rate decrease occurs faster than bacteria inactivation (Sun et al., 2017). Damaged membrane proteins participating in bacterial respiratory chain causes a decreased respiration rate, this generates harm but does not inactivate bacteria directly, it can be due to the fermentation ability of *E. coli* (Clark, 1989). ATP generation capability of bacteria decreases through photocatalysis and results in the complete loss of ATP synthesis. Without ATP synthesis, bacteria cannot repair the damaged cell membrane, cannot sustain Na^+ -K⁺ pump activity and membrane potential. In addition, the absence of ATP causes K⁺ leakage from the cytoplasm (Sun et al., 2014;

Bosshard et al., 2009). According to the findings of researchers, it can be concluded that direct cause of bacteria inactivation is due to the loss of ATP synthesis ability instead of damaged membrane structure and decreased respiration (Sun et al., 2017).

<u>Enzymes.</u> Upon photocatalysis, intracellular ROS like H_2O_2 and O_2^{-} which can be named as secondary oxidants are generated. Inside the cell, H_2O_2 and O_2^{-} can produce reactive HO[•] by a Fenton reaction including free iron (Equations 2.11 and 2.12) (An et al., 2017). *E. coli* contains almost 20 μ M of iron for Fenton reactions (Park and Imlay, 2003). O_2^{-} is continuously created as a by-product upon aerobic respiration (Tortora et al., 2018).

$$Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + HO^{\bullet} + OH^{-}$$
 (2.11)

$$O_2^{\bullet-} + H_2O_2 \longrightarrow HO^{\bullet} + OH^- + O_2$$

$$(2.12)$$

Photocatalytic treatment processes and increased inner ROS generate oxidative stress to the cell and bacteria produce some anti-oxidative enzymes against this stress. Superoxide dismutase (SOD) and catalase (CAT) are the main enzymes. SOD catalyzes the conversion and detoxification of O_2^{-} by converting to H_2O_2 and O_2 , afterwards CAT carries out the decomposition of H_2O_2 to harmless H_2O and O_2 (Sun et al., 2017; Chiang and Schellhorn, 2012). Upon photocatalytic inactivation, SOD and CAT levels decrease and as a result of this, intracellular ROS formation increases. The inactivation of these enzymes causes the accumulation of both intracellular and extracellular ROS and accelerates the inactivation rate of the system (An et al., 2017). Moreover, lesion repairing process of DNA after oxidative damage could be achieved by the action of specific enzymes and proteins (Cabiscol et al., 2000).

<u>Proteins and DNA.</u> Researchers detected leakage of intracellular substances like protein and nucleic acids by using scanning electron microscopy (SEM) images after photoelectrocatalytic treatment (Sun et al., 2014). Carré and colleagues showed that after photocatalytic treatment of *E. coli*, bacteria lost membrane associated proteins (Carré et al., 2014). Furthermore, intracellular ROS can lead to the oxidation of proteins which produces carbonyl groups (Dalle-Donne et al., 2003). After solar disinfection process, bacterial proteins can be aggregated by the attack of ROS and this aggregation occurs due to the oxidation of proteins (Bosshard et al., 2010b).

DNA is a more sensitive macromolecule to the oxidative stress than others; therefore, the damage or loss of bacterial DNA is a fatal factor for inactivation. This DNA damage can occur via base losses, breakage of one strand or double strand of the DNA (Hidaka et al., 1997).

2.4.3.3. Detection of bacteria inactivation and mineralization. During photocatalysis and as a result of oxidative degradation, leakage of cytoplasmic components of bacteria to the solution matrix would be expected to occur and leaked cytoplasmic substances would also be subjected to direct interaction with extracellular ROS. Therefore, leaked cytoplasmic components can be oxidized, segmented, or go for complete mineralization upon exposure to ROS attack (Suryo et al., 2012; Hirakawa et al., 2004).

Leung and colleagues examined dissolved organic carbon (DOC) content of the reaction medium after photocatalytic treatment and detected an increase reaching a peak and then a decreasing trend following the experiment. These findings showed that the released bacterial components can be degraded within extended irradiation periods, even complete mineralization of total DOC content can be expected (Leung et al., 2008). Consequently, from a general perspective, it can be expected that bacterial inactivation could also be followed via DOC variations in the reaction medium along with cell enumeration.

2.5. Water Matrix

Natural aquatic systems contain natural organic matter (NOM), microorganisms, anions, cations and pollutants giving chemical, physical and biological characteristics to water. In this study, organic materials like NOM and microorganisms and a water matrix (WM) comprised of inorganic materials as common anions (Cl⁻, NO₃⁻, SO₄^{2–}, PO₄^{3–}, HCO₃⁻/CO₃^{2–}) and cations (Na⁺, K⁺, Ca²⁺, Mg²⁺) were used to simulate natural water conditions.

2.5.1. Effect of Water Matrix Components on Solar Light Initiated Photocatalytic Processes

In photocatalytic degradation processes, the solution matrix is very important since the adsorptive effect of the components can inhibit or alter the reaction rate of the process. The competition and retardation effects of different types of NOM, photocatalysts, metal ions, common anions and cations and inhibitory effects of alkalinity on TiO₂ photocatalysis process have been investigated to understand the limitations of the photocatalytic system (Bekbolet, 1997; Bekbolet and Balcioglu, 1996; Bekbolet et al., 1998; Gonenc and Bekbolet, 2001; Uyguner and Bekbolet, 2007, 2009; Uyguner-Demirel and Bekbolet, 2011).

Related to pH_{zpc} values of the particles and the pH of the solution, surface of the particles gains negative or positive charges. The presence of common anions and cations could hinder or improve

the generation of electrostatic attractions between organic matter, bacteria and photocatalyst surfaces. Uyguner and Bekbolet reported that the presence of common anions could lead the shrinkage of HA volume and shape however the presence of divalent, trivalent cations could cause enhanced intermolecular attractions and could end up with coagulation or precipitation (Uyguner and Bekbolet, 2009).

<u>2.5.1.1.</u> Effect of chloride. The presence of chloride ions (Cl⁻) causes decreased oxidation rates in the system since they are adsorbed by the catalyst. Chloride ions could block the photocatalyst's active surface sites thus the catalyst could be deactivated against organic molecules (Abdullah et al., 1990; Bekbolet et al., 1998). This can also occur by the scavenging effect of chloride. Chloride ions decrease the oxidation rate by scavenging oxidizing radical species ('ox) (Equation 2.13) (Abdullah et al., 1990; Matthews, 1988). Moreover, Cl⁻ may react with the HO' as given in the following reactions (Equations 2.14-2.16) (Grigor'ev et al., 1987; Jayson et al., 1973).

$ox + Cl^- \rightarrow Cl^{\bullet} + ox^-$	(2.13)
$HO^{\bullet} + Cl^{-} \rightarrow ClOH^{\bullet-}$	(2.14)
$\text{ClOH}^{\bullet-} \rightarrow \text{HO}^{\bullet} + \text{Cl}^{-}$	(2.15)
$ClOH^{\bullet-} + H^+ \longrightarrow Cl^{\bullet} + H_2O$	(2.16)

Also, some competitive reactions could form chloride ion and photogenerated holes, h^+_{VB} / electrons, e^-_{CB} and NOM species (Equations 2.17-2.19) (Uyguner and Bekbolet, 2009). Formed dichloride radical anion (Cl₂⁻⁻) is less reactive than chlorine radical (Cl^{*}), however, reactive for both bacteria and organic matter (Alegre et al., 2000; Rincón and Pulgarin, 2004). Re-combination of both two previously generated Cl^{*} (Equation 2.20) and Cl₂⁻⁻ (Equation 2.21) form molecular chlorine (Cl₂) in water. These reactions can enhance the inactivation rate due to disinfectant effect of chlorine (Rincón and Pulgarin, 2004).

$$Cl^{-} + e^{-}_{CB} \rightarrow Cl^{-}$$

$$(2.18)$$

$$Cl^{+} + Cl^{-} \rightarrow Cl^{*-}$$

$$(2.19)$$

 $Cl' + Cl' \rightarrow Cl_2 \tag{2.20}$

 $Cl_2^{\bullet-} + Cl_2^{\bullet-} \rightarrow Cl_2 + 2Cl^-$ (2.21)

<u>2.5.1.2.</u> Effect of nitrate. Nitrate ions (NO₃⁻) could block the active sites of the photocatalysts partially thus its inhibitory effect is less than Cl⁻. (Abdullah et al., 1990). On the other hand, NO₃⁻

ions have reactivity in natural water. As they are irradiated (λ =320 nm), HO[•] form thus they could enhance the degradation rate of organic compounds. Nitrate and nitrite (NO₂⁻) give the following reaction in water (Equations 2.22-2.30) (Bekbolet et al., 1998; Uyguner and Bekbolet, 2009; Vione et al., 2014; Zafiriou and True, 1979a, 1979b).

$$\begin{split} & \text{NO}_{3}^{-} + \text{hv} \to \text{NO}_{2}^{-} + \text{O} (^{3}\text{P}) & (2.22) \\ & \text{NO}_{3}^{-} + \text{hv} \to \text{NO}_{2}^{+} + \text{O}^{-} & (2.23) \\ & \text{O}^{-} + \text{H}_{2}\text{O} \to \text{HO}^{+} + \text{OH}^{-} & (2.24) \\ & \text{NO}_{3}^{-} + \text{H}_{2}\text{O} + \text{hv} \to \text{NO}_{2} + \text{HO}^{+} + \text{OH}^{-} & (2.25) \\ & \text{NO}_{3}^{-} + \text{H}^{+} + \text{hv} \to \text{NO}_{2}^{+} + \text{HO}^{*} & (2.26) \\ & \text{NO}_{2}^{-} + \text{H}^{+} \to \text{NO}_{2}^{*-} \to \text{NO}^{*} + \text{O}^{-} & (2.27) \\ & \text{O}^{*-} + \text{H}^{+} \to \text{HO}^{*} & (2.28) \\ & \text{NO}_{2}^{-} + \text{H}_{2}\text{O} + \text{hv} \to \text{NO}^{*} + \text{HO}^{*} + \text{OH}^{-} & (2.29) \\ & \text{NO}_{2}^{-} + \text{H}^{+} + \text{hv} \to \text{NO}^{*} + \text{HO}^{*} & (2.30) \\ \end{split}$$

HO[•] formed in natural waters mainly give rections with HAs. This non-selective aqueous HO[•] can seperate H atoms from organic molecules or add to C=C double bond or accept an electron (Hoigné, 1990). NO₃⁻ have ability to photolyze under sunlight conditions to form HO[•] in natural water bodies; therefore, oxidation of humic substances (HSs) in sunlit natural surface waters could be induced in the presence of NO₃⁻ (Kotzias et al., 1982; Zafiriou, 1974).

<u>2.5.1.3. Effect of sulfate.</u> Sulfate ions (SO_4^{2-}) could directly be adsorbed on positively charged TiO₂ surface sites and deactivate some portion of the photocatalyst. This deactivation can cause retardation in the oxidation rates of organic molecules. Moreover, SO_4^{2-} could bind with the positive holes (h⁺) and form sulfate radicals (SO_4^{--}) if all oxidizing sites of the surface are available for adsorption (Equation 2.31) (Abdullah et al., 1990).

$$h^+ + SO_4^{2-} \rightarrow SO_4^{--} \tag{2.31}$$

Also, HO[•] radicals could react with SO_4^{2-} species to give $SO_4^{\bullet-}$ (Equations 2.32 and 2.33) (Matthews at al., 1972; Uyguner and Bekbolet, 2009). With the effect of these reactions, oxidation rate of the organic molecules could be enhanced.

$$\mathrm{HO}^{\bullet} + \mathrm{HSO}_{4^{-}} \longrightarrow \mathrm{H}_{2}\mathrm{O} + \mathrm{SO}_{4^{\bullet}}^{\bullet}$$

$$(2.32)$$

$$\mathrm{HO}^{\bullet} + \mathrm{SO}_4^{2-} \longrightarrow \mathrm{H}_2\mathrm{O} + \mathrm{SO}_4^{\bullet-} \tag{2.33}$$

<u>2.5.1.4.</u> Effect of phosphate. Phosphate ions $(H_2PO_4^-)$ react like SO_4^{2-} anions by adsorbing on the surface active sites of the photocatalyst and decrease the rate of oxidation (Abdullah et al., 1990; Uyguner and Bekbolet, 2009). Phosphate ions could react similarly with sulfate ions by binding with the positive holes and form dihydrogen phosphate radicals $(H_2PO_4^{\bullet})$ (Equation 2.34) (Abdullah et al., 1990). Formed $H_2PO_4^{\bullet}$ is not reactive as HO[•] (Rincón and Pulgarin, 2004).

$$h^+ + H_2 PO_4^- \to H_2 PO_4^{\bullet} \tag{2.34}$$

Moreover, H₂PO₄⁻ anion actions like a screen by absorbing UV radiation and this low amount of photo absorption could reduce the light reaching bacteria in the solution and provide some light protection to bacteria (Rincón and Pulgarin, 2004).

<u>2.5.1.5. Effect of bicarbonate.</u> Bicarbonate ions (HCO₃⁻) behave as scavengers of h^+ formed on irradiatied TiO₂ surface (Equation 2.35) and decrease the inactivation rate (Abdullah et al., 1990).

$$h^+ + HCO_3^- \to HCO_3^{\bullet} \tag{2.35}$$

At neutral pH conditions (pH: 6-7), the surface charge of TiO₂ is neutral as well (pH_{zpc} \approx 6.5). Under these conditions, HCO₃⁻ adsorbs on TiO₂ and forms a negatively charged layer on the surface causing less positive surface charge. Due to this change on the surface charge, TiO₂ adsorption on *E*. *coli* decreases and causes reduced inactivation rates. Moreover, HCO₃⁻ could demonstrate photo absorption and reduce light penetration into the bacteria solution, resulting in some light protection to bacteria like phosphate ions (Rincón and Pulgarin, 2004).

 HCO_3^- could react with HO[•] radical to produce less reactive carbonate radical (CO_3^{-}) (Equations 2.36 and 2.37) (Chen et al., 1997), its reactions are slower than HO[•] (Rincón and Pulgarin, 2004).

$$HO^{\bullet} + HCO_{3}^{-} \rightarrow CO_{3}^{\bullet-} + H_{2}O$$

$$(2.36)$$

$$\mathrm{HO}^{\bullet} + \mathrm{CO}_{3}^{2-} \to \mathrm{CO}_{3}^{\bullet-} + \mathrm{OH}^{-} \tag{2.37}$$

<u>2.5.1.6.</u> Effect of monovalent cations. The presence of both Na⁺ and K⁺ as monovalent cations decrease the inactivation rate of bacteria. In addition, researchers reported that inhibitory effect of K⁺ is higher than Na⁺. Rincón and Pulgarin suggested that the addition of K⁺ into the solution containing *E. coli*, compensates K⁺ leakage from the cell membrane and cause the retardation of cell damage (Rincón and Pulgarin, 2004).

2.5.1.7. Effect of divalent cations. The presence of divalent cations as Ca^{2+} and Mg^{2+} in low concentrations displayed enhanced DOC removal efficiency in high DOC containing samples. Uyguner and Bekbolet reported that Ca^{2+} has more effectiveness than Mg^{2+} (Uyguner and Bekbolet, 2009).

Rincón and Pulgarin performed a study to elucidate the effect of individual common anions and cations on TiO₂ photocatalytic inactivation of E. coli (Rincón and Pulgarin, 2004). It was reported that preliminary experiments conducted in the absence of photocatalyst specimen and sole HCO3⁻ showed an enhancement in the inactivation rate of bacteria. However in the presence of SO₄²⁻, NO₃⁻, Cl⁻ and H₂PO₄²⁻ anions a retardation effect was observed in inactivation rates expressing no significant difference between *E. coli* count in the presence of SO_4^{2-} , NO_3^{-} , Cl^- and $H_2PO_4^{2-}$ anions. In the presence of TiO₂, anions decrease the inactivation rates in an increasing order as: Cl⁻, NO₃⁻, SO_4^{2-} , HCO_3^{-} and HPO_4^{2-} ; the effectiveness of HCO_3^{-} and HPO_4^{2-} was higher than other anions. Besides the effects of individual anions, researchers further studied the removal extends of anions from the system with respect to their dissappearnce from filtered solution via E. coli intake and/or adsorption onto TiO₂ surface occurring mainly throuh electrostatic interactions. Only in the presence of TiO₂ and negatively charged anions, removal % of anions by adsorption onto photocatalys surface showed a decreasing order as: $NO_3^- \ge Cl^- > SO_4^{2-} > HPO_4^{2-} > HCO_3^-$. On the other hand in the presence of *E. coli* and anions, removal % of anions followed a decreasing trend as: $HCO_3^- > SO_4^{2-}$ > Cl⁻ >> HPO₄²⁻ > NO₃⁻. Conclusively, in the presence of all reactive species as anions, *E. coli*, and TiO₂, anion removal % via both bacteria uptake and adsorption onto TiO₂ showed a decreasing trend as: $HCO_3^- > HPO_4^{2-} > SO_4^{2-} > NO_3^- > Cl^-$. According to removal % of anions, HCO_3^- was the most absorbed anion by E. coli, causing enhanced sensitivity to irradiation by changing membrane characteristics of bacteria, and increased bacteria inactivation efficiency (Rincón and Pulgarin, 2004).

Romazzoni and colleagues studied SODIS and photo-Fenton processes in the presence of inorganic ions and suggested that *E. coli* K-12 inactivation in both processes increased with the usage of 10 mg/L HCO₃⁻ (Romazzoni et al., 2020). This enhancement could be due to formation of CO₃⁻⁻ radical which has much longer lifetime in aqueous solutions than HO⁺. Addition of 100 mg/L Cl⁻⁻ provides enhanced disinfection rates in both processes. When researchers used increased concentrations of Cl⁻, SODIS showed enhanced results but photo-Fenton process was disrupted. SO₄²⁻⁻ addition in the range of 0-500 mg/L showed no difference in SODIS but led to increased inactivation rates in photo-Fenton process. NO₂⁻⁻ and NO₃⁻⁻ usage provide increased inactivation to both processes and the effect of NO₂⁻⁻ was higher than NO₃⁻⁻. Moreover, the addition of NH₄⁺⁻ showed an enhancement for photo-Fenton, while had no effect on SODIS (Romazzoni et al., 2020).

2.6. Natural Organic Matter

NOM can be defined as all the organic matter, in both dissolved and particulate form in nature other than living organisms and synthetic materials (Buffle, 1984; Buffle, 1988). Dissolved form of NOM in oceans and in all types of freshwaters is named as dissolved organic matter (DOM) and particulate fraction of NOM is referred to as particulate organic matter (POM) (vanLoon and Duffy, 2000). Various forms of NOM under the concept of total organic matter (TOM) and total organic carbon (TOC) present in natural waters are displayed as a Venn schema in Figure 2.10. Dissolved organic nitrogen (DON) species e.g., proteins, peptides, amino acids and dissolved organic phosphorus (DOP) species e.g., phospholipids and polyphosphates are also considered in general domain of TOM, explicitly in DOM and more specifically as a part of DOC. Non-humic fraction should also be expressed within the dissolved organic matrix.



Figure 2.10. Venn diagram of various NOM forms found in natural waters. TOM or NOM, TOC, DOM, DOC, POC, DON, DOP and humic and non-humic materials are represented (Pagano et al., 2014).

DOM is the main carbon-based component that can range in molecular weight from a few hundred to 100,000 daltons (Da), in the colloidal size range (Aiken et al., 1985; Thurman, 1985). A schematic mass (MW, Da) and size (μ m) distribution diagram of NOM species and organisms in natural waters is shown in Figure 2.11. Expressing comparatively larger size mostly greater than 1 μ m, species like zooplankton, phytoplankton and bacteria could be regarded as POM. The International Humic Substance Society (IHSS) uses natural waters to isolate NOM fractions.



Figure 2.11. Organic matter mass (Da) and size (μ m) distribution diagram of NOM species and organisms in natural waters. Molecular weight (MW), humic acid (HAc), fulvic acid (FAc), hydrocarbons (HC), carbohydrates (CHO), fatty acids (FA), amino acid (AA) (Artifon et al., 2019).

NOM contains HSs like HAs and fulvic acids (FAs) and non-humic substances covering proteins, polysaccharides and other various components in its structure (MacCarthy et al., 1990). NOM can originate from two different types of sources as allochthonous and autochthonous. While allochthonous NOM is generated from terrestrial or watershed sources, it indicates the presence of HS, autochthonous NOM arises from algae, bacteria and macrophytes and it mainly indicates the presence of algal derived organic matter. NOM chemistry, composition and amount can change with the source of the organic matter, climate, geology of the field and other cases like algal blooms and runoffs (Leenheer and Croué, 2003).

2.6.1. Humic Substances and Humic Acid

HSs are polyelectrolytic, dark colored organic acids expressing highly complex and heterogeneous chemical nature. From a general perspective, HSs can also be defined as the heteropolycondensed products of carbohydrates, proteins, fatty acids, lignins, tannins and many other compounds based on their origin (Gjessing, 1976). HSs display polydisperse characteristic with respect to their wide range of molecular sizes (Figure 2.12) (Aiken et al., 1985). HSs constitute 60 to 80 % of DOM in natural waters. The elementary composition of most HSs could be represented in the following order as; C: 45-55 %; O: 30-45 %; H: 3-6 %; N: 1-5 %; and S: 0-1 % (Manahan, 1999).


Figure 2.12. Conceptual model of polydisperse HSs (Haberhauer et al., 2000).

HSs can be examined in three different types according to their solubility characteristics (Figure 2.13). HA is not soluble at pH below 2 but soluble at higher pH. FA is soluble at all pH values. And lastly humin is the non-extractable plant residue that is insoluble at any pH (Gaffney et al., 1996).



Figure 2.13. The postulated relation of HSs (Uyguner et al., 2007).

The humic content of various soils expressing diverse characteristics display variations in the range of 0 to 10 %. On the other hand, humic content of surface waters is expressed as DOC differing from 0.1 to 50 ppm in dark-water swamps. Whereas, in ocean water DOC changes in the range of 0.5 to 1.2 ppm at the surface, and 0.1 to 10 ppm in deep groundwaters (Choppin and Allard, 1985). Moreover, approximately 10 % of DOC in surface waters is found as suspended matter that can be present in the form of organic or organically coated inorganic particulates e.g., clay minerals and oxide surfaces (Gaffney et al., 1996). HSs generate 60-80 % of DOC and POC in surface waters

(Reuter and Perdue, 1977). As the acid insoluble fraction of HS, elemental content of HA could be expressed in the following order as: C: 60.0 %; O: 34.5 %; H: 4.47 %; N: 0.96 % (Mao et al., 1998).

It is reported that pH_{zpc} of HA can vary in the narrow range of pH: 4-5 due to the presence of functional groups (vanLoon and Duffy, 2000). Therefore, under acidic or alkaline pH conditions, structure of HA can exhibit conformational rearrangements (Figure 2.14). While at low pH conditions, HA shows a compact and coiled structure, however, at high pH values HA has almost linear structure resembling extended polymer configuration. At neutral pH conditions (pH: 6-7) resembling natural water, the surface of HA can be recognized as partially-negatively charged due to the deprotonation of carboxylic functional groups (pH: 3-5). Deprotonation of phenolic functional groups can also increase the number of negatively charged sites on HA in aqueous medium conditions expressing alkaline pH > 8 conditions.



Figure 2.14. a. Conformational changes of HA molecules through alkaline to acidic pH; A. Charge repulsion at alkaline pH, B. Intramolecular aggregation at decreasing pH, C. Intermolecular aggregation at decreasing pH, D. Precipitation at acidic pH (de Melo et al., 2016). b. Tail, loop, and train formations of a HA polymer chain adsorbed on the surface (Adachi et al., 2012).

The hypothetical structure of HA has a hydrophobic aromatic core which is substituted with functional groups like free and bonded phenolic groups, quinine structures, nitrogen and oxygen as bridge units and carboxyl groups (COOH) (Figure 2.15). The hypothetic model contains both



hydrophilic and hydrophobic sites and has an immensely polyelectrolytic feature and some available sites to bind with organic compounds, mineral surfaces, and metal ions (Suffet and MacCarthy, 1989).

Figure 2.15. a. Hypothetical structure of HA (Fetsch et al., 1998); b. Functional groups and their range of pK_a values present in HA (Nada et al., 2019); c. Theoretical structure of HA (Stevenson, 1982); d. Proposed three-dimensional structure of HA (Engebretson and von Wandruszka, 1994).

2.6.2. Photochemistry of Humic Substances in Natural Water Systems

From a general perspective aquatic photochemistry is mainly dependent upon solar irradiation. Besides various photo-reactions occurring by inorganic species, as the principal component of natural waters, NOM may undergo either direct or indirect photolysis although both of the mechanisms may occur simultaneously.

UV-vis absorption spectra and fluorescence spectral features display the presence of chromophores and fluorophores present in NOM. Although a continuous decreasing absorbance with respect to increasing wavelength is a general trend of HSs, origin and type dependent variations may also be assessed (Figure 2.16) (Uyguner et al., 2007).



Figure 2.16. a. UV-vis absorption spectra, b. emission scan, and c. synchronous scan fluorescence spectra of IHSS soil HA (SHA), IHSS HA, IHSS FA, Aldrich HA (AHA), Roth HA (RHA), and Nordic HA (NHA) (Uyguner et al., 2007).

The chromophoric centers in numerous compounds that are also considered as building blocks of natural organic matter express quite different absorption and photochemical properties (Figure

2.17) (Vione et al., 2014). Examples of these reactions could simply be photoionization (phenolic groups), photoreduction (quinonoid groups, conjugated systems) and homolysis.



Figure 2.17. Representative functional groups and building blocks of HSs (Vione et al., 2014).

Aquatic NOM can either be originated from terrestrial runoff as allochthonous or directly formed within the system as autochthonous. Therefore, the photosensitizing properties are extensively investigated and generally considered as the same for both allochthonous and/or autochthonous organic matter. Upon light absorption, changes in absorption and fluorescence properties of organic matter may occur via photolysis and irreversible reactions may lead to structural changes in organic matter. During indirect photolysis process, photosensitization process predominates over direct photolysis. HSs are known to be the main photosensitizers in natural waters since HSs are Omni present organic matter expressing ill-defined chemical structure as explained in Section 2.6.1.

In natural waters, HSs are the most important sunlight absorbing substances leading to photosensitized oxygenation reactions (Zepp et al., 1977). Photosensitization can be defined as a process initiating a photochemical reaction using a photosensitizer which is a substance having the ability of absorbing light and transferring the energy to another molecule (Zafiriou et al., 1984). When a photosensitizer is excited by photon energy, fundamental changes in its electron distribution occurs as described by internal homolysis of electron pairs. Electron pair homolysis of a molecule results in formation of excited singlet and triplet states (Schenck, 1963). Photosensitizers may not be consumed during these reactions, they may excite and/or transfer energy to other molecules or simply return to their original states (IUPAC, 2004). The altered molecule by photosensitizer is called as substrate or acceptor. Further reactions could result transformation and even degradation of organic compounds through various pathways such as photochemical, redox and hydrolysis.

Photochemical performances of NOM and its analogs has been widely studied since 1970s (Zepp et al., 1976, 1977; Aguer and Richard, 1993, 1996; Aguer et al., 1999). DOM mainly HSs control photochemical reactions in water systems. The photoinduction of DOM causes the formation of ROS. Probable photochemical reactions of DOM in aqueous media presented below (Equations 2.38-2.49), where the contributors are singlet DOM (¹DOM), triplet DOM (³DOM), $e^{-}(aq)$, HO[•], scavenger (R), $O_2^{\bullet-}$, HO₂[•], H₂O₂, triplet oxygen (³O₂) and singlet oxygen (¹O₂) (Cooper, 1988; Uyguner-Demirel et al., 2017; Zepp et al., 1985). Reactions begin with the absorption of light by the functional groups of DOM which generates the photo-exited ¹DOM and ³DOM, and following reactions form ROS.

$DOM + h\nu \rightarrow {}^{1}DOM^{*} \rightarrow {}^{3}DOM^{*}$	(2.38)
$^{1}\text{DOM}^{*}_{(aq)} \rightarrow \text{DOM}^{+} + e^{-}_{(aq)}$	(2.39)
$DOM^{\bullet+} + O_2/H_2O \rightarrow HO^{\bullet} + DOM$	(2.40)
$O_2 + e^{(aq)} \rightarrow O_2^{\bullet-}$	(2.41)
$^{1}\text{DOM}^{*}_{(aq)} + R \rightarrow R^{\bullet} + \text{DOM}^{\bullet}$	(2.42)
$R^{\bullet} + DOM^{\bullet} + O_2/H_2O \rightarrow HO^{\bullet} + R + DOM$	(2.43)
$^{3}\text{DOM}^{*} + \text{O}_{2} \rightarrow \text{DOM}^{\bullet} + \text{O}_{2}^{\bullet-}$	(2.44)
$O_2^{\bullet-} + H^+ \leftrightarrow HO_2^{\bullet}$ $pK_a=4.8$	(2.5)
$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$	(2.7)
$HO_2 + O_2 + H_2O \rightarrow H_2O_2 + O_2 + OH^-$	(2.45)
$^{3}\text{DOM}^{*} + ^{3}\text{O}_{2} \rightarrow \text{DOM}^{\bullet+} + \text{O}_{2}^{\bullet-}$	(2.46)
$^{3}\text{DOM}^{*} + {}^{3}\text{O}_{2} \rightarrow {}^{1}\text{DOM} + {}^{1}\text{O}_{2}$	(2.47)
$^{1}O_{2} + H_{2}O \rightarrow ^{3}O_{2} + H_{2}O$	(2.48)
$^{1}\text{DOM} + {^{1}\text{O}_{2}} \rightarrow \text{DOM}^{\bullet+} + \text{O}_{2}^{\bullet-}$	(2.49)

Indirect photoreactions take place through initial excitation followed by energy transfer leading to formation of various photo-products namely ROS in the presence of dissolved oxygen. These free-radical species could react with various compounds e.g., amino acids (tryptophan) or fatty acids, etc.. If such materials are released from cells, they may be subjected to irradiation directly resulting in complete destruction and forming components of organic debris. Considering heterogeneous and polydisperse nature of HSs, generation of ROS could proceed selectively via light interaction with low molecular or high molecular fractions with respect to their degree of functionality and hydrophobic core (Vione et al., 2014). Since all natural waters contain colloidal matter either as living or non-living in nature, these compounds are also prone to photolysis and/or photo-initiated reactions under sunlit environmental conditions (Scully et al., 2003).

DOM has attracted attention of researchers in recent years due to its influence in water treatment. Especially for disinfection purposes in case that chlorination process is applied, DOM is the major precursor in the formation of DBPs most significantly trihalomethanes (Richardson et al., 2002; 2007). Furthermore, DOM can serve as substrates for the undesirable microbiological growth in pipelines promoting public health concern. Therefore, removal of DOM is considered as the main target of all water treatment facilities. Besides conventional treatment processes (coagulation and flocculation), oxidation based technologies i.e. UV-irradiation, or ozonation are also frequently applied (Parsons, 2004). Photocatalytic degradation of NOM and its representative compounds like HA and FA have been investigated by Bekbolet and colleagues (Bekbolet and Ozkosemen, 1996, Bekbolet et al., 1998; Uyguner and Bekbolet, 2005; Uyguner-Demirel and Bekbolet, 2011). The investigation of photocatalytic degradation of NOM was studied in detail by various reseachers (Espinoza and Frimmel, 2009) and several review papers were published about this subject (Matilainen and Sillanpää, 2010; Uyguner-Demirel et al., 2017).

Throughout the application of solar light initiated photolytic and photocatalytic destruction of bacteria, the release of organic matter can be observed and characterized (Carré et al., 2014; Castro-Alférez et al., 2016; Kiwi and Nadtochenko, 2005). The release of organic matter and DOM can be followed with UV-vis absorption spectra and with the application of specific fluorescence techniques like EEM. UV-vis parameters at 436 nm (Color₄₃₆) and 365 nm (UV₃₆₅) give information about the presence of color forming moieties and absorbance at 280 nm (UV₂₈₀) and 254 nm (UV₂₅₄) present the aromatic fractions (Uyguner and Bekbolet, 2005).

Specific visible (SCoA₄₃₆) and UV absorbance (SUVA₃₆₅, SUVA₂₈₀ and SUVA₂₅₄) values represent DOC normalized color forming and aromatic moieties (Uyguner and Bekbolet, 2005).

SUVA₂₅₄ value can be used to identify water sample's composition in terms of hydrophobicity and hydrophilicity. SUVA₂₅₄ < 3 indicates the presence of hydrophilic and aliphatic content however SUVA₂₅₄ > 4 represents the existence of hydrophobic and aromatic material (Edzwald et al., 1985).

Fluorescence spectroscopy techniques are quantitative analytical techniques can be used to characterize NOM due to their sensitivity and non-destructive features (Alberts and Takács, 2004; Senesi, 1990). Fluorescence occurs during the excitation of a substance by the absorption of a photon from UV-vis region and emission of light while returning to its ground state, and at specific wavelengths it describes the relation of absorbed and emitted photons (Valeur, 2002). NOM has internal fluorescence characteristics accompanying with changes in the emission maximum wavelength when exiting at different ones. Besides limited distinctiveness of excitation scan spectra and emission scan spectra, synchronous scan fluorescence spectroscopy maintains a constant between excitation and emission wavelengths; therefore, it is a useful tool to indicate differences and similarities between samples (Figure 2.16). Synchronous scan fluorescence spectrum at $\lambda_{emis}=470$ nm (FI_{sync470}) indicates the presence of HA (Uyguner and Bekbolet, 2005), and at $\lambda_{emis}=280$ nm (FI_{sync280}) reflects the presence of protein, especially tryptophan in the solution (Reynolds, 2003).

EEM fluorescence spectrophotometry can also be used to characterize different fluorophores in the water sample. In EEM fluorescence contour plots, natural organic matter exhibits fluorescence in excitation wavelength range of $\lambda_{exc} \sim 200$ to 500 nm and emission wavelength range of $\lambda_{emis} \sim 280$ to 600 nm (Baker et al., 2008). When the destruction of bacteria occurs, the presence of aromatic proteins I (λ_{exc} 220-250 nm and λ_{emis} 280-332 nm), aromatic proteins II (λ_{exc} 220-250 nm and λ_{emis} 332-380 nm), microbial by-products (λ_{exc} 250-470 nm and λ_{emis} 280-380 nm), humic-like substances (λ_{exc} 220-470 nm and λ_{emis} 380-580 nm), and fulvic-like substances (λ_{exc} 220-250 nm and λ_{emis} 380-580 nm) can be seen in EEM fluorescence plots (Figure 2.18) (Coble, 1996).

Since intrinsic fluorescence properties reflects information pertaining to structure, conformation, and heterogeneity of organic matter as well as dynamical properties related to intramolecular and intermolecular interactions, a fluorescence-derived index as fluorescence index (FI) is defined as the ratio of the emission intensity at λ_{emis} =450 nm to that at λ_{emis} =500 nm, following the excitation at λ_{exc} =370 nm (Bekbolet and Sen-Kavurmaci, 2015; McKnight et al., 2001). FI \leq 1.4 represents the presence of humic fluorophores of organic matter mainly derived from terrestrial sources. On the other hand, FI \geq 1.9 indicates the existence of organic material stemming from microbial origin (Sen-Kavurmaci and Bekbolet, 2014). Relating this information to DOM characterization and bacteria cell

destruction probably resulting in release of fluorophoric organic compounds and/or fractions would be crucial in understanding mechanism of microorganism inactivation.



Figure 2.18. Diagram of EEM fluorescence contour plot regions.

Bacteria inactivation by TiO₂ photocatalysis has been studied for elucidation of mechanistic details by many research groups as stated previously in Section 2.4.2. (Bekbolet, 1997; Bekbolet and Araz, 1996; Bekbolet and Tomruk, 1998; Bekbolet, 2007). Birben and colleagues presented that the presence of humic matter expressed a retardation effect on solar photocatalytic inactivation of *E. coli*. Regrowth of *E. coli* could not be assessed under specific experimental conditions. EEM fluorescence contour plots displayed the formation of new fluorophoric regions described as microbial by-products and aromatic proteins. Also, undoped TiO₂ P-25 expressed comparatively slower removal rates in comparison to doped specimens (Birben et al., 2017a).

Besides the already reported studies by Bekbolet and colleagues on *E. coli* photocatalytic inactivation, further interest was directed to use of a VLA photocatalyst such as Fe-doped TiO₂. TiO₂ source was P-25 serving as a base specimen and a novel sol-gel synthesized TiO₂ was also introduced expressing different physico-chemical and morphological properties. It should also be mentioned that the aforementioned VLA Fe-doped TiO₂ specimens were previously utilized for HA degradation under simulated solar light irradiation (Birben et al., 2017b). Therefore, solar light initiated photocatalytic treatment of *E. coli* by using Fe-doped TiO₂ species in various aqueous matrix conditions would bring further insight into bacteria inactivation in natural waters.

3. MATERIALS AND METHODS

3.1. Materials

3.1.1. Preparation of E. coli suspension

E. coli K12 strain, ATCC 23716 was used as an indicator bacterium. *E. coli* growth was provided by inoculation of one *E. coli* colony in 25 mL of LB broth. The suspension was incubated at 37° C and shaken at 90 rpm for 16-18 hours. Working *E. coli* solutions (10^{5-8} Colony Forming Unit (CFU)/mL) were prepared in various reaction media which was explained below Sections 3.1.4, 3.1.5, and 3.1.6. Cell samples were taken from *E. coli* suspension during the stationary phase of the bacteria growth. *E. coli* was freshly incubated in LB broth for each experiment set.

3.1.2. Luria Bertani Broth

Aqueous solution of LB broth was prepared by using 2 g tryptone, 1 g yeast extract and 2 g NaCl (Bertani, 1951; Singleton, 1997). LB broth was freshly prepared as a volume of 200 mL when required.

3.1.3. CHROMagarTMECC

CHROMagarTM ECC, which is a selective solid medium for *E. coli* and other coliforms was used. CHROMagarTM ECC was composed from 15 g/L agar, 8 g/L peptone and yeast extract, 5 g/L NaCl, and 4.8 g/L chromogenic mix, was used as a culture medium (CHROMagar, 2019). It provides easy plate reading by giving different colors to different bacteria. *E. coli* colonies demonstrate blue color while other coliform colonies have mauve color. Other bacteria colonies are either colorless, or growth has been inhibited.

3.1.4. Isotonic Solution

Saline solution designated as isotonic solution (IsoT) is used in the experiments to obtain optimal osmatic pressure condition to bacteria. IsoT as 0.85 % solution was prepared by adding 8.5 g NaCl (Sigma-Aldrich) into 1 L of distilled-deionized water and dissolved using magnetic stirrer to obtain complete dissolution (APHA, AWWA, WPCF, 2012).

3.1.5. Water Matrix Solution

Water matrix referred as WM, containing common anions and cations such as Cl⁻, NO₃⁻, SO₄²⁻, H₂PO₄⁻, HCO₃⁻/CO₃²⁻, Na⁺, K⁺, Ca²⁺, and Mg²⁺ was used as a natural surface water analogue and prepared according to the recipe from Smith and colleagues (Smith et al., 2002). Concentrations of anions and cations were verified and presented at Table A.1.

3.1.6. Humic Acid Solution

Commercial humic acid sodium salt as purchased from Aldrich was used as a DOM analogue. Aqueous stock solution of 1000 mg/L HA was prepared by dissolving 1.0 g HA in 1 L of ultrapure water (Millipore Milli-Q plus system, with resistivity of 18.2 M Ω cm at 25°C). Working HA solution's concentration was chosen as 50 mg/L and it was prepared by diluting the stock solution. In order to prepare 100 kDa molecular size fraction of 50 mg/L HA, sequential fractionation as prefiltration through 0.45 µm cellulose acetate membrane filter and subsequent ultrafiltration process using 100 kDa cut-off membrane filter (Amicon 8050 stirred cell unit) was applied. Spectroscopic characterization of 100 kDa HA was presented at Appendix B Section.

3.1.7. Photocatalyst Specimens

A total of six photocatalyst specimens were used as two undoped and four Fe-doped types. TiO₂ P-25 (Evonik) and synthesized TiO₂ (SynTiO₂) were used as bare TiO₂ specimens and their 0.25% and 0.50% Fe-doped species, namely as 0.25% Fe-TiO₂, 0.50% Fe-TiO₂, 0.25% Fe-SynTiO₂, and 0.50% Fe-SynTiO₂, were used as doped types.

TiO₂ P-25 which mainly composed of 80 % anatase and 20 % rutile crystal phases was supplied from Evonik and used as a standard photocatalyst specimen. TiO₂ exhibits surface area of 55 ± 5 m²/g, primary particle size as 30 nm and pH_{zpc} as 6.25. SynTiO₂ was prepared by applying a modified sol gel method using titanium (IV) isopropoxide (97 % Aldrich) as the precursor material (Turkten and Cinar, 2017). Fe-doped TiO₂ and SynTiO₂ photocatalyst specimens were prepared according to wet impregnation method using Fe(NO₃)₃·9H₂O (Merck) as the iron source (Yalcin et al., 2010). Excluding TiO₂ P-25, all of the prepared photocatalyst specimens were calcined at 500°C for 5 h. Physicochemical and morphological characteristic properties of all six photocatalysts specimens were previously reported and elucidated by crystallographic properties (XRD), band-gap energy (UV-DRS), surface morphology (SEM) and Brunauer-Emmett-Teller (BET) surface area, and presented in Appendix C Section (Birben et al., 2015, 2017; Turkten, 2016; Turkten and Cinar, 2019; Turkten et al., 2019; Yalcin et al., 2010). Photocatalyst loading was chosen as 0.25 mg/mL and kept constant for all experiments.

3.2. Methods

3.2.1. Experimental Set-Up

Solar light initiated photolytic and photocatalytic inactivation experiments were carried out using an Atlas-Suntest CPS+ solar simulator. The light source is an air-cooled Xenon lamp with a wavelength range of λ =290 nm to 800 nm and light intensity, I₀=250 W/m². The solar simulator is equipped with quartz and UV special glass filters to simulate typical outdoor sunlight irradiation. All experiments were carried out by using 150 mL cylindrical Pyrex reaction vessel the diameter and height of which is equal to 7.5 cm and 3.5 cm, respectively (Figure 3.1).



Figure 3.1. Solar simulator Atlas-Suntest CPS+ and reaction vessel.

3.2.2. Experimental Procedure

Photocatalyst concentration was used as 0.25 mg/mL in all solar light initiated photocatalytic disinfection experiments. An ultrasonic water bath was used to obtain a homogenous suspension of the working solution and the photocatalyst prior to the addition of *E. coli*. Sample volumes were kept constant for all experiments as 100 mL. In order to maintain a homogenized sample, all experiments were carried out under constant continuous stirring. Samples were subjected to treatment for specified reaction periods (0, 10, 20, 30, 40 and 60 minutes). Distilled-deionized water was used for volume correction to account for loss due to evaporation (<10 % in 60 min). Following each treatment period,

10 mL of sample was collected for bacteria enumeration processes and the remaining 90 mL sample was filtered through 0.45 μ m membrane filters to remove bacteria and TiO₂ specimens from the solution prior to analysis.

3.2.3. Analytical Methods

3.2.3.1. *E. coli* enumeration. After each treatment period, 10 mL composite sample was collected in 15 mL falcon tube. Extreme care was taken to sampling in order to ensure that the sample would be representing the whole reaction medium. Samples were further subjected to *E. coli* enumeration. Sampling was carried out under continuous mixing by pipetting 1 mL sample from 10 different locations of the pyrex reaction vessel. Samples were diluted with 0.85 % NaCl IsoT solution before inoculation. Appropriate sample dilutions $(10^{-2} \text{ to } 10^{-5})$ were selected based on irradiation time of each sample. Under prolonged irradiation periods, lower dilutions and high sample volumes were preferred. In the experiments where high *E. coli* concentration is expected, samples were diluted three times and low inoculation volumes were preferred like 10 or 20 µL. In case that the expected *E. coli* concentration would be low, samples were not diluted at all or simply diluted one-fold, and high inoculation volumes were used such as 50 or 100 µL. All experiments were performed as triplicate runs.

E. coli enumeration as CFU/mL was performed by viability test and calculations were made according to following equation presented in Standard Methods (Equation 3.1) (APHA, AWWA, WPCF, 2012).

$$CFU/mL = \frac{Colonies \ counted}{Volume \ plated \ (mL)} \times Dilution \ Factor$$
(3.1)

Drop plate method was chosen for the *E. coli* enumeration due to its practicability and to allow inoculating sufficient number of parallel cultures (Miles and Misra, 1938; Reed and Reed, 1948). CHROMagarTM ECC was used as a culture medium to provide high color contrast as could be visualized by blue color between *E. coli* and other coliform bacteria colonies having mauve colorization. Petri dish diameter was chosen as 9 cm. An appropriate dilution and volume of sample which inoculated on the medium was chosen according to expected bacteria inactivation rate. Inoculated petri dishes were kept in the incubator at 44°C for 24 h. Enumeration of *E. coli* colonies were carried out by visual inspection (Figure 3.3).



Figure 3.2. *E. coli* cultured CHROMagarTM ECC plates.
Top Left: 10⁻² dilution from initial *E. coli* solution, inoculation volume: 10 μL.
Top Right: 10⁻⁵ dilution from *E. coli* suspension, inoculation volume: 5 μL.
Middle Left: 10⁻³ dilution from initial *E. coli* solution, inoculation volume: 20 μL.
Middle Right: 10⁻³ dilution from t=0 *E. coli* solution, inoculation volume: 10 μL.
Bottom Left: 10⁻² dilution from t=0 *E. coli* solution, inoculation volume: 10 μL.

<u>3.2.3.2.</u> Total protein determination. Lowry method as developed by Lowry and colleagues was chosen as protein determination method (Lowry et al., 1951). It has been the most widely used protein estimation method in molecular biology due to its moderately constant sensitivity, applicability, and suitability for protein concentration in the range of 0.01-1.0 mg/mL.

The method consists of two distinct steps. In the first step, which is based on Biuret reaction, sample treated with the Lowry reagent that contains NaOH-Na₂CO₃-CuSO₄-Na₂C₄H₄O₆ and peptide bonds in the proteins react with copper under alkaline conditions to form a copper-protein complex. In the second step of the method, copper-protein complex is treated with the phenol reagent developed by Folin and Ciocalteu (Folin and Ciocalteu, 1927). The Folin-Ciocalteu reagent is a phosphotungustic-phosphomolybdic acid solution. The reagent is reduced by the copper-treated proteins to molybdenum blue which can be determined colorimetrically at λ =750 nm.

 $Protein + Cu^{2^{+}} \xrightarrow{OH^{-}} Tetradendrate Cu^{+} complex \xrightarrow{Folin-Ciocalteu reagent} Copper - protein complex$

The reaction mainly depends on the tyrosine and tryptophan content of the sample. The main advantage of the method is its sensitivity, but a wide range of substances can interfere with the method, including nitrogen containing buffers, drugs, nucleic acids, glucose, uric acid, guanine, xanthine, aromatic amines, unsaturated aromatic compounds, sulphides, sulphurous acid, H_2O_2 , ferrous ions, thymol, sulfosalicylic acid, picric acid (over 0.1 percent), glycine (over 0.5 percent), hydrazine (over 0.5 percent) and ammonium sulphate (over 0.15 percent) (Herbert et al., 1971; Lowry et al., 1951; Sapan et al., 1999; Waterborg, 2002).

<u>Procedure.</u> Samples were taken from the +4°C and brought to room temperature. Then, samples were vortexed, and 3.0 mL of sample was inserted into a 16x100 mm glass tube. Then, 4.2 mL of Lowry reagent was added into the tubes. Tubes were capped and vortexed, then incubated for 20 minutes at room temperature in dark conditions. After 20 minutes of incubation, 0.6 mL of diluted Folin-Ciocalteu Reagent was added into the tubes. Samples were vortexed and incubated for 35 minutes at room temperature in dark conditions. After incubation, molybdenum blue color formed which represented the concentration of reduced Folin-Ciocalteu Reagent measured at λ =750 nm. Bovine Serum Albumin (BSA) was used as a standard and calibration curve was plotted in the range of 0-100 mg/L (Appendix D).

<u>*The Lowry Reagent.*</u> The Lowry reagent was daily prepared by mixing Solution A, B and C with a volume ratio of 100:1:1 respectively.

Solution A (alkaline solution). 2.8598 g NaOH and 14.3084 g Na₂CO₃ were dissolved in 500 mL of distilled-deionized water.

Solution B. 0.9104 g CuSO4 was dissolved in 100 mL of distilled-deionized water.

Solution C. 2.8530 g Na₂C₄H₄O₆.2H₂O was dissolved in 100 mL of distilled-deionized water.

<u>The Folin-Ciocalteu reagent, 1N.</u> The reagent was prepared prior to use at the last five minutes of the first incubation time by mixing 5 mL of 2N Folin-Ciocalteu's phenol reagent and 5 mL of distilled-deionized water in an amber bottle.

<u>3.2.3.3. Total carbohydrate determination.</u> Phenol-sulfuric acid method was chosen as carbohydrate determination method. It is the most widely used colorimetric carbohydrate determination method. The method can be applied to detect all types of carbohydrates including monosaccharides, disaccharides, oligosaccharides, and polysaccharides (Dubois et al., 1956).

Colorimetric methods for detection of carbohydrates are based on mild oxidizing/reducing properties of monosaccharides. In phenol-sulfuric acid method monosaccharides act as reducing agents (Panagiotopoulos and Sempere, 2005). Addition of H₂SO₄ into phenol solutions containing carbohydrate provides enough heat under highly acidic conditions thus glycosidic bonds of the carbohydrate can completely hydrolyze and the condensation of dehydrated sugars with phenol groups generate a yellow-orange colored aromatic complex which is characteristic for the reaction (Figure 3.4) (Saha and Brewer, 1994; Panagiotopoulos and Sempere, 2005). The color produced is proportional to the amount of carbohydrate and stable for several hours. Absorbance of the solution is measured quantitatively at λ =490 nm to determine sugar concentration. This method is accurate to ±2 % under appropriate conditions (Dubois et al., 1956; An and Smith, 2005).



Figure 3.3. Schematic reaction of phenol-sulfuric acid method (Viel et al., 2018).

Despite the wide range of applicability of phenol-sulfuric acid method, broad range of substances can interfere with the reaction including cellulose fibres, other insoluble carbohydrates, NO_3^- and Fe^{3+} and glycoproteins (Dubois et al., 1956; Martens and Frankenberger, 1990; Rao and Pattabiraman, 1989). Moreover, investigators reported that when humic material is treated with H₂SO₄, it gives absorbance in 350-500 nm range and may interfere with the total carbohydrate determination (Sieburth and Jensen, 1969; Meadows and Campbell, 1978). Additionally, a strong acid usage may cause the production of some degradation products which can absorb the light in the same spectral band with carbohydrates, resulting in the overestimation of carbohydrate concentration (Josefsson et al. 1972; Dawson and Liebezeit, 1981). Researchers indicated that color development depends on the order of the reagent addition and sample matrix which could cause low reproducibility in environmental samples (Artem'yev, 1970; Gerchakov and Hatcher, 1972). Moreover, researchers reported that method gives various absorbance values for equal concentrations of different types of sugars, so it is more accurate when it is used for the samples that contain only one type of carbohydrate (Brummer and Cui, 2005; Josefsson et al., 1972).

<u>Procedure.</u> Samples were taken from +4°C and brought to room temperature. Then, samples were vortexed, and 1.0 mL of samples were inserted into 16x100 mm glass tube. Then, 1.0 mL of 5 % of phenol solution and 5 mL of concentrated H₂SO₄ were added rapidly and directly against the liquid surface to achieve good mixing. Samples were kept on stand for 10 minutes, and then they were vortexed and placed into thermoreactor for digestion at 32°C for 25 minutes. After digestion, samples were cooled down and generated yellow-orange color was measured at λ = 490 nm. Glucose was used as a standard and calibration curve was plotted in the range of 0-50 mg/L (Appendix E).

<u>5 % Phenol solution</u>. Phenol was supplied from Merck and 5 g was dissolved in 100 mL of distilleddeionized water.

<u>3.2.3.4. K² leakage determination.</u> K² leakage from bacterial cells via solar photocatalytic inactivation was determined by applying the flame photometric method outlined as 3500-K B in Standard Methods (Standard Methods for the Examination of Water and Wastewater, 2012). Perkin Elmer AAnalyst 300 Atomic Absorption Spectrometer was used. The method is based on the determination of potassium at λ =766.5 nm and reported as Total K, mg/L. Calibration range was selected as 0-2.0 mg/L and Potassium Inductively Coupled Plasma Certipur® solution (Merck) was used as a standard.

<u>3.2.3.5. Endotoxin determination.</u> Endotoxin detection was carried out using ENDOSAFE® Gel-Clot Limulus Amebocyte Lysate test (LAL). LAL test is based on the usage of LAL clotting protein derived from horseshoe crab *Limulus polyphemus amebocytes* for the qualitative detection of Gramnegative bacteria endotoxins by gel-clot method (Levin and Bang, 1968). LAL technique can detect as low as $10^{-6} \mu g/mL$ of endotoxin (Reinhold and Fine, 1971; Yin et al., 1972). When in-activated endotoxins are used, the test gives negative results. In addition, LAL gives reactions not only with the lipopolysaccharide (LPS) but also free lipid A fractions of LPS (Yin et al., 1972).

Samples were treated by buffered lysate in single test-vials in a thermoreactor at 37°C for 60 minutes. Presence of endotoxin was detected by the formation of visible clot in test tubes with respect to a selected limiting value as 0.125 (Endotoxin Units) EU/mL. It can be detected with momentary inversion of the test tube up to two minutes after the incubation period (WHO, 2012; Maji and Chatterjee, 2011).

<u>3.2.3.6.</u> UV-vis spectroscopic absorbance measurements. Absorbance recordings were taken in wavelength range of 200-600 nm by employing Perkin Elmer Lambda 35 UV-vis Spectrometer. Perkin Elmer quartz cuvettes with 1.0 cm path-length that were rinsed with distilled-deionized water prior to each analysis. UV-vis absorption spectra of the samples were measured in the indicated wavelength range using distilled-deionized water as a reference. The presence of color forming moieties and aromatic domains were measured by using specified absorbance measurements as wavelength of 436 nm, 365 nm, 280 nm, and 254 nm and designed as $Color_{436}$, UV_{365} , UV_{280} , and UV_{254} respectively. Specified UV-vis parameters were explained as follows:

Color₄₃₆: Absorbance at 436 nm, Color forming moieties

- UV₃₆₅ : Absorbance at 365 nm, Organic matter content
- UV_{280} : Absorbance at 280 nm, Aromaticity, Organic matter content
- UV_{254} : Absorbance at 254 nm, Surrogate parameter for DOC

Specific UV-vis parameters defined as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀ and SUVA₂₅₄ were calculated by normalization of specified UV-vis parameters to DOC (mg/L) according to following equations (Equations 3.2-3.5).

$SCoA_{436} = Color_{436}/DOC (L/m mg)$	(3.2)
$SUVA_{365} = UV_{365}/DOC (L/m mg)$	(3.3)
$SUVA_{280} = UV_{280}/DOC (L/m mg)$	(3.4)
$SUVA_{254} = UV_{254}/DOC (L/m mg)$	(3.5)

<u>3.2.3.7. Fluorescence spectroscopic measurements.</u> Fluorescence spectroscopic measurements in the synchronous scan mode were recorded by using Perkin Elmer LS 55 Luminescence Spectrometer. Luminescence spectrometer is equipped with a 150W Xenon arc lamp and a red sensitive photomultiplier tube. 1.0 cm path-length quartz cuvettes were rinsed with distilled-deionized water before and after each analysis to prevent the presence of any contamination.

Synchronous scan fluorescence data were obtained using a bandwidth of $\Delta\lambda$ =18 nm between excitation and emission monochromators and in an emission wavelength range of λ_{emis} =200 nm to 600 nm. Depending on the system characteristics, specified fluorescence intensities detected at λ_{emis} =280 nm and λ_{emis} =470 nm were elucidated and designated as FI_{sync280} and FI_{sync470}. Moreover, specific fluorescence intensities SFI_{sync470} and SFI_{sync280} were also calculated by normalization of maximum synchronous scan fluorescence intensities as FI_{sync280} and FI_{sync470} to DOC content of the sample (Equations 3.6 and 3.7).

$$SFI_{sync280} = FI_{sync280}/DOC$$
(3.6)
$$SFI_{sync470} = FI_{sync470}/DOC$$
(3.7)

FI values were calculated by using the ratio of the emission intensity at λ_{emis} =450 nm to that at λ_{emis} =500 nm, following the excitation at λ_{exc} =370 nm (Equation 3.8).

$$FI = \frac{\text{The emission intensity at }\lambda \text{emis}=450 \text{ nm following the excitation at }\lambda \text{exc}=370 \text{ nm}}{\text{The emission intensity at }\lambda \text{emis}=500 \text{ nm following the excitation at }\lambda \text{exc}=370 \text{ nm}}$$
(3.8)

EEM fluorescence profiles were obtained by simultaneous incremental changes in both excitation and emission wavelengths. A gradual increase of λ_{exc} from 200 nm to 500 nm and λ_{emis} from 200 nm to 600 nm were recorded. Three-dimensional contour plots were derived from data and modelled using MATLAB R2013a program. Fluorescence trend based selected excitation-emission regional information of EEM fluorescence contour plots were explained in detail in Section 2.4.

<u>3.2.3.8. Non-purgeable organic carbon measurements.</u> Non-purgeable organic carbon (NPOC) (mg Org C/L) contents of the samples were determined by Total Organic Carbon Analyzer Shimadzu TOC-VWP calibrated using potassium hydrogen phthalate with a concentration range of 0-25 mg Org C/L as the standard. NPOC mode sparging inorganic carbon contents prior to organic carbon determination was selected to determine the DOC contents. All NPOC measurements were represented with mg/L for simplicity reasons.

4. RESULTS AND DISCUSSION

In this study, four different aqueous matrix conditions referred to as follows; IsoT solution, WM solution, HA solution, and "HA and WM" solution were used, to investigate their different effects on solar photolytic and photocatalytic inactivation of *E. coli*. IsoT solution was used to obtain optimum osmotic pressure condition to bacteria in the absence of other inorganic and organic content and its results represented the baseline conditions for other matrixes. WM solution was used as a synthetic natural water representative containing common anions and cations and its composition was addressed in Table A.1. HA solution having 100 kDa molecular size fraction was used to introduce NOM in the system and observe the bacteria inactivation in the presence of organic matter. Spectroscopic characterization of 100 kDa HA was presented in Appendix B Section. "HA and WM" solution was prepared as a mixture of WM and HA solutions and used to mimic natural water conditions containing both inorganic and organic matter in the medium.

Solar photocatalytic inactivation treatments were carried out with mainly two types of photocatalyst as undoped and Fe-doped TiO₂ specimens. TiO₂ (Evonik) and SynTiO₂ were used as undoped photocatalyst specimens and 0.25% Fe-TiO₂, 0.50% Fe-TiO₂, 0.25% Fe-SynTiO₂ and 0.50% Fe-SynTiO₂ were used as their Fe-doped versions. Characteristic properties of all six photocatalysts were represented in Appendix C Section.

Solar photolytic and photocatalytic inactivation of *E. coli* could be successfully expressed by first order kinetic model, logarithmic reduction value (LRV) and percentage of reduction. Logarithmic reduction profile of bacteria inactivation indicated that kinetics could well be expressed by first order kinetic model. Calculations were made according to Chick's law equation (Equation 4.1) (Chick, 1908).

Rate (R) =
$$-dN/dt = kN$$
 or $\ln(N/N_0) = -kt$ or $N/N_0 = e^{-kt}$ (4.1)

Where,

R: first order rate (CFU/mL min) N: *E. coli* count, CFU/mL at time t N_o: initial *E. coli* count, CFU/mL t: irradiation time, min k: first order reaction rate constant, min⁻¹ Half-life ($t_{1/2}$, min) was calculated by the following Equation 4.2 as; $t_{1/2}=0.693/k$ (4.2)

LRV of bacteria was calculated according to following equation (Equation 4.3).

$$LRV = \log_{10}(N_0/N_{60})$$
(4.3)

Where,

N_o: initial *E. coli* count, CFU/mL as indicated above N₆₀: *E. coli* count, CFU/mL at irradiation time of 60 min

LRV could also be expressed as percentage reduction and the respective conversions were given in Table 4.1. (EPA, 1991).

Reduction, %
68
90
96.8
99
99.7
99.9
99.99

Table 4.1. Bacteria LRV in terms of percentage reduction.

As could be visualized in Table 4.1, LRV of 4 indicates 99.99 % bacteria reduction and it could go further as LRV of 5 and 6 expressing reductions as 99.999 and 99.9999 %.

4.1. Solar Photolytic Inactivation of E. coli

4.1.1. Solar Photolytic *E. coli* Inactivation Under Specified Reaction Conditions, Reductions and Kinetics

E. coli suspension in IsoT solution (N₀=5.46E+05 CFU/mL) was subjected to solar photolytic treatment. Upon irradiation period of 30 min, *E. coli* count decreased to 5.25E+02 CFU/mL and following further exposure of t_{irr}=60 min, *E. coli* count was reduced almost by half to 2.65E+01 CFU/mL. Under these conditions LRV of *E. coli* was 4.31 and reduction percentage was 99.99 %. In case when *E. coli* suspension in WM solution (N₀=4.61E+05 CFU/mL) was treated for an irradiation period of 30 min, bacteria count reduced to 2.67E+02 CFU/mL. Following light exposure of t_{irr}=60

min, *E. coli* count decreased to 2.30E+01 CFU/mL revealing LRV as 4.30 and reduction as 99.99 %. *E. coli* (N_0 =1.09E+06 CFU/mL) suspension was directly prepared in HA solution and exposed to irradiation for a period of 30 min revealing an *E. coli* count of 5.37E+03 CFU/mL. Upon light exposure for a duration of t_{irr}=60 min, *E. coli* cell count further decreased to 2.50E+02 CFU/mL revealing LRV as 3.64 and reduction as 99.9 %. *E. coli* suspension with an initial count of N_0 =7.09E+05 CFU/mL was prepared in a solution composed of "HA and WM" components resembling natural waters was further subjected to irradiation for 30 min. Under these conditions, *E. coli* count decreased to 2.98E+04 CFU/mL. Upon light exposure of t_{irr}=60 min, *E. coli* count slightly decreased to 2.63E+03 CFU/mL revealing LRV as 2.43 and reduction as 99 %.

Based on these results, the effect of solution matrix for *E. coli* reduction could be displayed in terms of LRV of *E. coli* in an increasing order as: "HA and WM" < HA < WM < IsoT. Irradiation time dependent *E. coli* inactivation profiles in the presence of different solution matrixes were presented in a comparative manner (Figure 4.1). First order kinetic model parameters of solar photolytic treatment of *E. coli* were also presented in Table 4.2.



Figure 4.1. Irradiation time dependent *E. coli* inactivation upon solar photolytic treatment in different solution matrixes.

Solar photolytic treatment of *E. coli* in IsoT solution resulted in a rather fast reduction with $k=0.184 \text{ min}^{-1}$ and half-life as 3.76 min. On the other hand, in the presence of both "HA and WM", reaction rate decreased significantly revealing $k=0.0920 \text{ min}^{-1}$ and half-life as 7.53 min. The effect of reaction medium could be expressed in terms of inactivation rate constant (k, min⁻¹) in a decreasing order of IsoT > WM > HA > "HA and WM". With respect to the minor differences in initial bacteria

counts, inactivation rates (CFU/mL min) followed a decreasing trend as; 1.42E+05 > 1.00E+05 > 8.02E+04 > 6.52E+04 for HA, IsoT, WM, and "HA and WM", respectively.

Matrix	First	IDV	Deduction 0/			
Maurx	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min	LKV	Keuuciioii, %	
<i>E. coli</i> in IsoT	0.184	3.76	1.00E+05	4.31	99.99	
<i>E. coli</i> in WM	0.174	3.98	8.02E+04	4.30	99.99	
<i>E. coli</i> and HA	0.130	5.33	1.42E+05	3.64	99.9	
<i>E. coli</i> in "HA and WM"	0.0920	7.53	6.52E+04	2.43	99	

Table 4.2. Solar photolytic treatment of *E. coli*: Inactivation parameters.

Saline solution (0.85 % NaCl) is an isotonic solution with pH=7.36 which provides a stable osmotic pressure on cell membrane of the bacteria and prevents the osmotic lysis of the cell (Tortora et al., 2018). Isotonic solution was chosen as the base experimental medium instead of phosphate buffer solution to prevent the inhibitory effect of phosphate species to the reaction medium upon photocatalysis (Long et al., 2017; Lyon et al., 2005). Even though the presence of this optimal osmotic conditions were assessed, cell counts of *E. coli* in IsoT solution showed high inactivation efficiency than expected. Sichel and colleagues reported that 0.7 log bacteria inactivation was achieved after t_{irr} =90 min of solar photolytic treatment (Sichel et al., 2007). Likewise, Birben and co-workers showed that almost 1 log *E. coli* reduction was attained at t_{irr} =60 min of solar photolytic treatment (Birben et al., 2017a). In this study, *E. coli* in IsoT solution showed 4 log bacteria reduction with the fastest reduction rate among other mediums, this high LRV could be explained with the combined effect of mechanistic stress factors.

E. coli in WM solution showed almost similar percent reduction with *E. coli* in IsoT suspension. High bacteria reduction could be explained with the presence of common anions and cations as Cl⁻, NO_3^- , SO_4^{2-} , PO_4^{3-} , HCO_3^-/CO_3^{2-} , Na^+ , K^+ , Ca^{2+} , and Mg^{2+} in WM. Nitrate ions are reactive in natural waters, once they are irradiated at λ =320 nm HO[•] forms in water (Equations 2.22-2.30). Other anions and cations could react with HO[•] and form radical species that may enhance the inactivation rate of bacteria. The effect of individual anions and cations in the photolytic inactivation conditions was reported that only in the presence of HCO_3^- an enhanced inactivation rate of bacteria was achived with ~7 log inactivation at t_{irr} =90 min. It was higher than the inactivation attained in the absence of any anion reaching 7 log bacteria reduction in ~135 min (Rincón and Pulgarin, 2004).

In the presence of HA solution, *E. coli* reduction was 99.9 % after t_{irr} =60 min. This value was lower than IsoT and WM media results (99.99 % for *E. coli* in IsoT solution and WM solution

conditions). Although HA acts like an exogenous photosensitizer (Cooper et al., 1988a), no significant effect was observed for bacteria inactivation. Presence of HA could cause lower bacteria inactivation rates in two ways; high NPOC content of HA solution as 4.78 mg/L and reduced light transmittance to bacteria due to HA. Similar findings were also reported in previous studies (Birben et al., 2017a, 2021b; Cantwell et al., 2008; Maraccini et al., 2016; Uyguner-Demirel et al., 2020).

The addition of WM into HA medium caused a slight retardation effect on bacteria reduction values. At the pH condition of the experimental solution as pH=6, surface charges of both *E. coli* and HA are negative. The cations present in the WM solution could be adsorbed on the surfaces of bacteria and organic matter which can decrease the available surface area giving reaction with ROS formed by anions in the WM. Moreover, photosensitization ability of HA could possibly be reduced by the adsorbed cations onto its surface. The observed retardation effect of reaction medium was also presented by Uyguner-Demirel and colleagues (Birben et al., 2021b; Uyguner-Demirel et al., 2020).

4.1.2. Organic Matter Formation and Removal Upon Solar Photolytic Treatment of *E. coli* in IsoT Solution

UV-vis absorption spectra of emerged organic matter in IsoT matrix in comparison to sole IsoT solution were presented in Figure 4.2. A consistent decreasing trend of UV-vis spectral features was observed under all irradiation periods. Moreover, UV-vis spectra of sole 0.85 % IsoT solution also expressed similar characteristics with the findings of Tong and colleagues (Tong et al., 2020).



Figure 4.2. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon solar photolytic treatment.

After addition of *E. coli* into IsoT solution, increased absorbances were recorded between λ =200 nm and 240 nm could be related to presence of organic moieties originating from EPS of bacteria. On the other hand, irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-240 nm excluding the presence of dense aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) that should be evident in longer wavelengths.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm which indicates the presence of proteins, especially tryptophan contents (Reynolds, 2003) in the sample (Figure 4.3). A significant increase of the fluorescence intensities (FI_{sync}) was recorded upon irradiation period of 40 min expressing the formation of fluorophores by cell damage and a decreasing trend was detected at the end of 60 min irradiation period. Accordingly, under initial conditions, FI_{sync280} was 25.1 following an increasing trend up to 20 min, then a sharp decrease followed by a sharp increase to 52.7 upon irradiation period of 40 min at which both UV₂₈₀ and UV₂₅₄ expressed the highest values as 0.0318 and 0.0464, respectively. These results would indicate the release of organics mainly composed of UV-absorbing centers also exhibiting fluorophores. Furthermore, FI_{sync470} displayed quite low presence changing in the range of 0.472-5.79.



Figure 4.3. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon solar photolytic treatment.

Due to the non-existence and possibly subsequent degradation of color forming moieties of the released organic matrix upon irradiation and the usage of IsoT solution, which was also a colorless solution, Color₄₃₆ and UV₃₆₅ values attained upon 30 min irradiation time were very low ≤ 0.0001 that could be regarded as insignificant (Figure 4.4). Under initial conditions, UV₂₈₀ and UV₂₅₄ were

recorded as 0.0238 and 0.0332 respectively. Upon treatment, UV_{280} variations expressed irradiation time dependent inconsistent trend in the range of 0.0149 to 0.0318. The highest UV_{280} was observed upon irradiation period of 40 min at which 62.6 % was removed at the end of t_{irr}=60 min. On the other hand, following a similar trend to UV_{280} , UV_{254} displayed variations 0.0232 to 0.0464. The highest UV-vis parameters were recorded upon irradiation period of 40 min.

From a general view, NPOC expressed variations in the range of 2.35-9.26 mg/L (Figure 4.4). The highest NPOC release was attained upon irradiation period of 30 min. NPOC content increased between initial (2.35 mg/L) and $t_{irr}=30 \min (9.26 \text{ mg/L})$ due to the release of organics from damaged and/or destructed *E. coli* cells, followed by a sharp decrease at $t_{irr}=40 \min (3.66 \text{ mg/L})$ and again followed by a decreasing trend during solar photocatalytic treatment.



Figure 4.4. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon solar photolytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.5. As the most significant parameter, SUVA₂₅₄ displayed variations in the range of 0.251-1.41 L/m mg during photolytic treatment expressing differences in both UV₂₅₄ and NPOC removals. SUVA₂₈₀ was recognized as the second significant specific UV-vis parameter varied in between 0.161 L/m mg and 1.01 L/m mg being the lowest for t_{irr}=30 min and the highest for initial solution. Accordingly, SCoA₄₃₆ expressed changes in between ≤ 0.0001 L/m mg and 0.172 L/m mg. Due to almost non-existence of Color₄₃₆ at t_{irr}=30 min, SCoA₄₃₆ could not be calculated. Contrary to very low SFI_{sync470} in the range of 0.108-1.58, predominant SFI_{sync280} expressed variations in the range of 3.58-14.4.



Figure 4.5. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT upon solar photolytic treatment.

Characteristic properties of organic matter under initial condition and upon solar photolytic inactivation of *E. coli* in IsoT solution exposed to irradiation for 60 min were presented in Table 4.3.

 Table 4.3. Characteristic properties of organic matter present in solar photolytic treatment of *E. coli* in IsoT solution.

Irradiation	uV-vis Spectroscopic Parameters, cm ⁻¹ NPOC, SUVA ₂₅ .	SUVA ₂₅₄ ,	FI	SEI	EI					
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L L/m mg	mg/L	L/m mg	Γ1 _{sync280}	SF1 _{sync280}	ГІ
Initial	0.0031	0.0057	0.0238	0.0332	2.35	1.41	25.1	10.7	1.44	
60 min	0.0019	0.0028	0.0199	0.0288	2.95	0.976	39.0	13.2	2.09	

Upon irradiation period of 60 min at which 4.31 log reduction of *E. coli* in IsoT solution was attained (Table 4.2), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0019, UV₃₆₅: 0.0028, UV₂₈₀: 0.0199, UV₂₅₄: 0.0288, and NPOC as 2.95 mg/L. Edzwald and colleagues reported that SUVA₂₅₄ < 3 indicated the presence of hydrophilic and aliphatic content of the sample (Edzwald et al., 1985). Initial and t_{irr}=60 min SUVA₂₅₄ values were calculated as 1.41 L/m mg and 0.976 L/m mg, respectively, representing that initial and remaining organic matter after t_{irr}=60 min could be considered having hydrophilic and aliphatic character.

FI_{sync280} increased at t_{irr}=60 min due to the released protein content of *E. coli*. SFI_{sync280} displayed slight increase at t_{irr}=60 min in accordance with the increased NPOC due to the released organic material from bacteria. Moreover, FI_{sync470}, which represents synchronous fluorescence intensity at λ_{emis} =470 nm and indicates the presence of humic-like fluorophores (Uyguner and Bekbolet, 2005),

were detected under initial and $t_{irr}=60$ min conditions as 0.472 and 2.02, respectively. Since the treatment was conducted in the absence of any humic material, $FI_{sync470}$ did not show any detectable peak as expected. A slight increase of $FI_{sync470}$ could be due to the released organics from damaged *E. coli* cells expressing complex aromatic structure. Accordingly, $SFI_{sync470}$ of initial and $t_{irr}=60$ min conditions were calculated as 0.201 and 0.683, respectively. Slightly increased $FI_{sync470}$ and NPOC contents caused an increase in $SFI_{sync470}$ at the end of 60 min irradiation period. From a general perspective, as $FI \ge 1.9$ represents the presence of microbially derived organic material, after $t_{irr}=60$ min, FI was calculated as 2.09 which complies with the previous findings of the researchers (Sen-Kavurmaci and Bekbolet, 2014). Bearing in mind the presence of extracellular organics, the results also implied that upon solar simulated irradiation conditions cell destruction could also be attained over an irradiation period of 60 min at which 99.99 % reduction was attained.

In accordance with the UV-vis absorbance and synchronous scan fluorescence spectroscopic analyses, EEM fluorescence contour plots of organic matter during irradiation time dependent *E. coli* inactivation in IsoT solution under solar photolytic treatment conditions were also followed and illustrated in Figure 4.6. It should be indicated that both emergence and removal mechanisms were prevailing under the specified conditions.



Figure 4.6. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon solar photolytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I (λ_{exc} 220-250 nm and λ_{emis} 280-332 nm), II (λ_{exc} 220-250 nm and λ_{emis} 332-380 nm), and IV (λ_{exc} 250-470 nm and λ_{emis} 280-380 nm) expressing that the fluorophores were mainly originated from aromatic proteins and microbial by-products under all irradiation conditions. Throughout the irradiation periods, released microbial by-products and proteins from damaged and destructed *E. coli* cells showed a decreasing trend in accordance with the solar photolytic degradation of released organic matter along with reduction of *E. coli* cells. The observed release of microbial by-products and proteins as a result of photolytic treatment of *E. coli* in IsoT solution was expected in accordance with previous studies of Birben and colleagues (Birben et al., 2017a, 2021b; Uyguner-Demirel et al., 2020). However, removal of these chromophoric groups and fluorophores could not be observed under the specified irradiation conditions most probably due to the inefficiency of the energy provided by simulated solar radiation and resistivity of these groups to degradation.

4.1.3. Mechanistic Evaluation of *E. coli* Inactivation Upon Solar Photolytic Treatment in IsoT Solution

Upon solar irradiation total K, protein, and carbohydrate contents present under initial and $t_{irr}=60$ min conditions as inactivation products of solar photolytic treatment of *E. coli* in IsoT solution were illustrated in Table 4.4.

Solor photolytic treatment	E. coli in IsoT solution			
Solar photolytic treatment	Initial	60 min		
Total K, mg/L	0.72	0.85		
Protein, mg/L	4.84	6.64		
Carbohydrate, mg/L	1.23	11.6		

Table 4.4. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in IsoT solution upon solar photolytic treatment.

Total K content of sole IsoT solution was detected as 0.042 mg/L however initial solution displayed higher total K which measured as 0.72 mg/L. High total K content of initial solution could be due to the membrane filtration applied prior to total K measurement. In membrane filtration step, reverse osmosis exerts to overcome osmotic pressure. Applied pressure is greater than osmotic pressure of solution to provide effective filtration. When bacteria exposed to increased external osmotic pressure, cell goes to dehydration and release solutes, namely osmolytes, including inorganic ions, mainly K⁺ ion, and organic molecules (Wood, 2015). Application of membrane filtration

generating increased osmotic pressure on *E. coli* cell wall could be the cause of K⁺ ion release and detected as total K content in initial solution. Therefore, initial total K content (0.72 mg/L) could be regarded as a background concentration for starting solutions following application of membrane filtration, irrespective of the minor differences between initial bacteria counts. After t_{irr}=60 min, total K content showed a slight increase. In regular conditions, K⁺ ion transportation in bacteria occurs via K⁺ ion channels and transporter proteins present in cytoplasmic membrane as passive transport mechanisms or Na⁺-K⁺ pumps by active transport; however, after solar photolytic inactivation process, damaged *E. coli* cell wall increases its permeability resulting in enhanced K⁺ ion release from the cell (Beagle and Lockless, 2020; Tortora et al., 2018).

Initial protein concentration of E. coli in IsoT solution prior to solar photolytic treatment was detected as 4.84 mg/L. Besides the inner protein content of bacteria, other protein containing structures are present on outer membrane of the cell wall as porin proteins, on cytoplasmic membrane as transporter proteins, and outside of the cell wall as flagella, fimbriae, and pili. Fimbriae and pili have different structures and functions than flagella. Fimbriae are shorter, thinner, and straighter than flagella and provide surface adhesion ability to bacteria, also it could be found over the entire surface. Pili are longer than fimbriae and each bacterium contains only one or two pili used in the DNA transfer from one bacterium to another. While flagella contain flagellar proteins like flagellin; fimbriae and pili consist of pilin protein. Moreover, degradative enzymes and transporter proteins are founding in periplasmic space defined as the region between outer membrane and cytoplasmic membrane (Tortora et al., 2018). It could be indicated that, the initial protein content of E. coli in IsoT solution could be due to the presence of these protein containing structures present on and outside of the bacterial plasma membrane and cell wall. Furthermore, protein content was also evidenced by EEM fluorescence contour plot under initial conditions prior to irradiation (Figure 4.6). At the end of 60 min irradiation period, protein content showed an increase (4.84-6.64 mg/L) due to release of proteins from damaged E. coli cells. Increased protein concentration in solution at tirr=60 min could also be verified by increased FI_{sync280} (Table 4.3).

Initial carbohydrate concentration was detected as 1.23 mg/L. It could be indicated that, the presence of glycocalyx or EPS highly composed of polysaccharides which synthesized inside the bacteria cell and secreted to the external part of the cell wall could be the cause of detected initial carbohydrate concentration. Due to released carbohydrate content from damaged bacteria, and carbohydrate containing cell structures, an increase was observed in carbohydrate concentration via bacteria inactivation upon irradiation period of 60 min as 11.6 mg/L. Increased NPOC contents significantly supported increased protein and carbohydrate contents at t_{irr} =60 min (Table 4.3).

4.1.4. Organic Matter Formation and Removal Upon Solar Photolytic Treatment of *E. coli* in WM Solution

UV-vis absorption spectra of organic matter in WM solution were displayed in Figure 4.7. Sole WM solution displayed a lower trend from initial and solar photolytic treatment samples. Different from IsoT solution conditions, the use of WM solution resulted in different absorbance characteristics wavelength at $\lambda < 250$ nm. Especially between $\lambda = 200$ nm and 210 nm, a shoulder effect was detected for all solutions originating from the strong absorption spectrum of NO₃⁻ having its λ_{max} at approximately 200 nm (Krishnan and Guha, 1934; Edwards at el., 2001). Increased absorbances were recorded between λ =200 nm and 240 nm following the addition of *E. coli* into WM solution, which could be revealing the presence of organic moieties originating from bacteria. On the other hand, irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-260 nm in the predominant absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) that should be evident in longer wavelengths. UV-vis absorption spectra of organic matter in WM solution displayed a declining trend without any absorption peaks complying UV-vis absorption spectra of humic substance which was represented in Section 2.4. (Figure 2.16a). A slightly increasing trend in between λ =240 nm and 280 nm could be distinguished under all irradiation period conditions, which could be due to released organic aromatic groups from bacteria. The existence of amino acids having aromatic moieties originating from the presence of EPS or inner content of lysed E. coli cells could be the cause of detected absorbance variations in the indicated region (Birben et al., 2021a).



Figure 4.7. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed intense peaks at $\lambda_{emis}=280 \text{ nm}$ and $\lambda_{emis}=420 \text{ nm}$ (Figure 4.8). A significant increase of FI_{sync} was recorded upon t_{irr}=40 min expressing the formation of fluorophores by cell damage and a decreasing trend was detected at 60 min of irradiation period. Similar emergence and removal of fluorescence intensities between irradiation period of 40 min and 60 min was also recorded for solar photolytic treatment of *E. coli* in IsoT solution (Figure 4.3). Maximum fluorescence intensity after $\lambda_{emis}=300 \text{ nm}$ indicates the presence of aromatic hydrocarbons (Dujmov and Sučević, 1990). An intense peak formed at $\lambda_{emis}=420 \text{ nm}$ could be originating from released organic content from bacteria. The intensities at $\lambda_{emis}=420 \text{ nm}$ displayed variations but decreased at t_{irr}=60 min showing the degradation of aromatic content in the sample. In the absence of HA, FI_{sync470} displayed quite low changes in the range of 0.805-2.46 however predominant FI_{sync280} expressed variations in the range of 10.8-37.9.



Figure 4.8. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

From a general perspective, NPOC expressed variations in the range of 7.82-15.8 mg/L (Figure 4.9). NPOC content displayed a fluctuating trend approximately as 2 mg/L until t_{irr} =20 min (9.29-9.81 mg/L) and increased significantly at t_{irr} =40 min (15.8 mg/L) due to the release of organics from damaged and destructed *E. coli* cells, followed by a sharp decrease after t_{irr} =40 min of solar photolytic treatment (7.82 mg/L). All UV-vis parameters displayed quite low presence however in the absence and subsequent degradation of color forming moieties of the released organic matter, Color₄₃₆ and UV₃₆₅ expressed lower values than UV₂₈₀ and UV₂₅₄. Highest UV₂₈₀ and UV₂₅₄ were recorded at t_{irr} =40 min as 0.0240 and 0.0338, respectively as could also be considered as similar to t_{irr} =60 min (0.0232 for UV₂₈₀; 0.0332 for UV₂₅₄).



Figure 4.9. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.10. SUVA₂₅₄ displayed variations in between 0.211-0.425 L/m mg during photolysis expressing differences in both UV₂₅₄ and NPOC removals. The highest SUVA₂₅₄ was calculated for irradiation period of 60 min. As the second significant parameter, SUVA₂₈₀ varied in between 0.145 and 0.297 L/m mg, being the lowest for t_{irr}=10 min and the highest for t_{irr}=60 min conditions. Furthermore, SFI_{sync470} and SFI_{sync280} expressed variations in the range of 0.057-0.156 and 1.17-3.86, respectively. The predominant fluorophoric group was accepted as SFI_{sync280}.



Figure 4.10. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

Characteristic properties of organic matter under initial condition and upon solar photolytic inactivation of *E. coli* in WM solution exposed to irradiation for 60 min were presented in Table 4.5.

Table 4.5. Characteristic properties of organic matter present in solar photolytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis S	pectroscop	troscopic Parameters, cm ⁻¹			SUVA ₂₅₄ ,	FLours280	EI	FI _{sync280}	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	SF1sync280	ГІ			
Initial	0.0009	0.0011	0.0173	0.0248	9.29	0.267	10.9	1.17	1.36		
60 min	0.0044	0.0070	0.0232	0.0332	7.82	0.425	22.1	2.82	1.54		

Upon irradiation period of 60 min at which 4.30 log reduction of *E. coli* in WM solution was attained (Table 4.2), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0044, UV₃₆₅: 0.0070, UV₂₈₀: 0.0232, UV₂₅₄: 0.0332, and NPOC as 7.82 mg/L.

In the absence of HA containing solution matrix, initial and irradiation period of 60 min SUVA₂₅₄ values were calculated as 0.257 L/m mg and 0.425 L/m mg, respectively. Besides the increase of SUVA₂₅₄ at the end of 60 min irradiation period, remaining organic matter could be considered having hydrophilic and aliphatic character (Edzwald et al., 1985).

FI_{sync280} also increased due to released protein content from *E. coli* cells. Relative to increased FI_{sync280} and decreased NPOC content at t_{irr}=60 min, SFI_{sync280} displayed an increase at 60 min of irradiation period. Detected FI_{sync470} were low in the absence of HA as 0.843 and 0.960 for initial and t_{irr}=60 min, respectively. This solar photolytic treatment was conducted in the absence of any humic material, therefore; FI_{sync470} did not exhibit any detectable peak. However, at λ_{emis} =420 nm FI_{sync} data displayed intense peaks for initial, t_{irr}=10 min, 20 min, and 40 min solutions due to released organic content from damaged bacteria cells. Moreover, SFI_{sync470} of initial and t_{irr}=60 min conditions were calculated almost insignificant as 0.0908 and 0.123, respectively. Besides slightly increased SFI_{sync470}, it displayed a decrease after t_{irr}=40 min (Figure 4.10) which means that released organic material from bacteria was degraded the end of the irradiation period of 60 min.

 $FI \le 1.4$ represents the presence of humic fluorophores originating from the presence of humic matter and $FI \ge 1.9$ indicates the existence of microbially derived organic material (Sen-Kavurmaci and Bekbolet, 2014). Although this solar photolytic treatment set was carried out in the absence of HA solution, FI was calculated as 1.54 after irradiation period of 60 min. It could be attributed that after *E. coli* reduction attained as 99.99 % at the end of 60 min irradiation period, released organic

content from damaged bacteria degraded and caused detection of a FI value between humic fluorophores and microbially derived fluorophores.

In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment were also followed and displayed in Figure 4.11.



Figure 4.11. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV indicating that the fluorophores were mainly originated from proteins and microbial by-products under all irradiation conditions. Following further irradiation times, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photolytic degradation of organic matter along with reduction of *E. coli*. However, complete removal of these chromophoric groups and fluorophores could not be observed under the specified irradiation conditions most probably due to the inefficiency of the energy provided by simulated solar radiation and resistivity of these groups to degradation. EEM fluorescence contour plots of organics observed under IsoT (Figure 4.6) and WM conditions could be considered as similar excluding regional fluorescence intensities.

4.1.5. Mechanistic Evaluation of *E. coli* Inactivation Upon Solar Photolytic Treatment in WM Solution

Upon irradiation, as inactivation products of *E. coli* cells in WM solution upon solar photolytic treatment, total K, protein, and carbohydrate contents released under initial and t_{irr} =60 min conditions were shown in Table 4.6.

	E. coli in WM solution			
Solar photolytic treatment	Initial	60 min		
Total K, mg/L	3.92	4.08		
Protein, mg/L	1.46	7.55		
Carbohydrate, mg/L	1.53	14.0		

 Table 4.6. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon solar photolytic treatment.

Total K content showed a slight increase at t_{irr} =60 min expressing K⁺ ion release from the bacteria cells. Solar photolytic inactivation of *E. coli* in WM solution resulted in a total K content of 4.08 mg/L. Initial total K concentration could be the result of K⁺ ion containing components present in WM solution which resulted in a concentration of 3.84 mg/L. Therefore, in the presence of WM, total K content of WM solution could be considered as a baseline concentration. In comparison to the very low total K attained in the presence of IsoT as 0.85 mg/L (Table 4.4) it could be indicated that almost similar amount of K⁺ ion was released upon irradiation for 60 min.

Initial protein concentration of *E. coli* in WM solution prior to solar photolytic treatment was lower than the protein content attained in the presence of IsoT solution (4.84 mg/L) as presented in Table 4.4. The reason could be attributed to sole presence of EPS in IsoT solution in the absence of any interfering components like WM solution. Moreover, upon irradiation for 60 min 7.55 mg/L protein release was attained being higher than the protein content released in the presence of IsoT solution (6.64 mg/L) (Table 4.4). In a comparative manner, it could be indicated that both under initial conditions and upon $t_{irr}=60$ min, carbohydrate contents were higher than *E. coli* treatment in IsoT solution (Table 4.4). Total carbohydrate content was 14.0 mg/L upon irradiation period of 60 min. Based on the attained concentrations, released protein and carbohydrate contents increased due to damaged bacteria cells. It should also be indicated that the WM components could certainly affect the findings occurring through various competing photo-initiated reactions as indicated previously at Equations 2.13-2.37, resulting in diminished effective light irradiation for direct *E. coli* inactivation.
4.1.6. Organic Matter Formation and Removal Upon Solar Photolytic Treatment of *E. coli* in HA Solution

UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.12 and 4.13. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-550 nm due to the presence of aromatic domains, UV absorbing centers as well as color forming moieties (all specified UV-vis parameters as Color₄₃₆, UV₃₆₅, UV₂₈₀ and UV₂₅₄). On the other hand, sole HA solution representing as HA initial (HA_i) expressed distinctly different trend in wavelength region below λ =230 nm at which ~0.582 absorbance was recorded for all solutions. Upon decreasing wavelength, HA_i expressed lower absorbance values contrary to the increasing trend of released organic matter due to cell destruction. However, obtained UV-vis absorption spectra of HA_i complying with UV-vis absorption spectra of humic substance which was represented in Section 2.4. (Figure 2.16a).



Figure 4.12. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed two intense peaks at λ_{emis} =400 nm and 470 nm, and a minor peak at λ_{emis} =280 nm (Figure 4.13). While peaks at λ_{emis} =280 nm specific to proteins, peaks at λ_{emis} =400 nm and 470 nm are characteristic to HA and in resemblance with the findings of Uyguner and Bekbolet (Reynolds, 2003; Uyguner and Bekbolet, 2005). An irradiation time dependent sequential decreasing order especially in FI_{sync470} was observed throughout light exposure. On the other hand, FI_{sync470} and FI_{sync280} displayed changes in the range of 35.4-58.6 and 6.38-9.60, respectively. In the presence of HA solution, FI_{sync470} accepted as the predominant

fluorophoric group. Moreover, $FI_{sync280}$ value of HA_i could not be detected however after the addition of bacteria into HA solution (initial), $FI_{sync280}$ increased to 7.70 indicating the presence of fluorophores reflecting proteinaceous content in the solution. After irradiation period of 60 min, $FI_{sync280}$ decreased (6.49) due to degradation of released protein content from *E. coli* cells.



Figure 4.13. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

NPOC content increased up to $t_{irr}=30 \text{ min } (4.78-17.3 \text{ mg/L})$ due to the release of the organics from the damaged and destructed *E. coli* cells, followed by a sharp decrease at $t_{irr}=40 \text{ min } (9.95 \text{ mg/L})$ and again followed by an increasing trend during solar photolytic treatment (17.4 mg/L) (Figure 4.14).



Figure 4.14. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

All UV-vis parameters expressed high presence relative to treatments conducted in the absence of HA solution (Figures 4.4, 4.9, and 4.14). Especially Color_{436} and UV_{365} displayed evident existence due to presence of colorful HA solution. Similar to previous solar photolytic inactivation treatments of *E. coli* in IsoT and WM solutions (Figures 4.4 and 4.9), the most significant UV-vis parameter was visualized as UV_{254} showing variations in between 0.4917-0.5000. However, specified UV-vis parameters did not express any significant difference between irradiation periods, therefore; it could be indicated that upon solar photolytic treatment, no significant degradation of organic content and color forming moieties was achieved.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.15. In contradiction to solar photolytic treatment sets of *E. coli* in IsoT and WM solutions (Figures 4.5 and 4.10) higher SUVA values were calculated in the presence of HA solution. As the most pronounced parameter, SUVA₂₅₄ displayed variations in the range of 2.83-10.5 L/m mg during solar photolysis expressing differences in both UV₂₅₄ and NPOC removals. The highest SUVA₂₅₄ was attained from initial condition.

Furthermore, besides very low SFI_{sync280} in the range of 0.372-0.965, SFI_{sync470} expressed variations in the range of 2.03-12.3. In the presence of HA solution, calculated SFI_{sync280} values were almost insignificant however SFI_{sync470} was accepted as the predominant fluorophoric group due to its remarkable presence in comparison to SFI_{sync280}. Moreover, according to insignificant presence of FI_{sync280} related to HA_i, SFI_{sync280} would be irrelevant compared to SFI_{sync470}.



Figure 4.15. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

Characteristic properties of organic matter under initial condition and upon solar photolytic inactivation of *E. coli* in HA solution exposed to irradiation for 60 min were presented in Table 4.7.

Table 4.7. Characteristic properties of organic matter present in solar photolytic treatment of *E. coli* in HA solution.

Irradiation time, min	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC,	SUVA ₂₅₄ ,	EI	SEI	EI
	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1sync470	SF1sync470	ГІ
Initial	0.0761	0.1630	0.4196	0.4997	9.01	5.55	56.3	6.25	1.08
60 min	0.0721	0.1573	0.4079	0.4933	17.4	2.83	35.4	2.03	1.10

Upon 60 min solar photolytic treatment of *E coli* in the presence of HA solution at which 3.64 log *E. coli* reduction was achieved (Table 4.2), released organic matter expressed following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0721, UV₃₆₅: 0.1573, UV₂₈₀: 0.4079, UV₂₅₄: 0.4933, and NPOC as 17.4 mg/L. After 60 min of irradiation period, NPOC expressed an increase (9.01-17.4 mg/L) due to present humic matter in reaction medium and released cytoplasmic organic content from damaged *E. coli* cells.

SUVA₂₅₄ of initial and t_{irr}=60 min conditions were calculated as 5.55 and 2.83 L/m mg, respectively. SUVA₂₅₄ < 3 indicated the presence of hydrophilic and aliphatic content whereas SUVA₂₅₄ > 4 revealed hydrophobic and aromatic substances in solution (Edzwald et al., 1985). It could be indicated that while initial solution displayed hydrophobic and aromatic features, upon t_{irr} =60 min organic matrix expressed hydrophilic and aliphatic characteristics. Following 60 min of irradiation period, decrease in SUVA₂₅₄ could be related to the degradation of aromatic organics during light exposure.

 $FI_{sync470}$ was significantly evident as 56.3 in initial condition due to the presence of HA solution and displayed a decrease to 35.4 at t_{irr}=60 min due to degradation of organics in the reaction medium. Calculated $SFI_{sync470}$ displayed a decrease at the end of 60 min irradiation period (6.25-2.03) in accordance with increased NPOC contents (9.01-17.4 mg/L).

Moreover, $FI_{sync280}$ under initial and $t_{irr}=60$ min conditions were 7.70 and 6.49, respectively. In the presence of HA solution, quite low $FI_{sync280}$ values were recorded in comparison to IsoT (Table 4.3) and WM solution conditions (Table 4.5); masking effect of HA on protein content could be the cause of low $FI_{sync280}$ values. $SFI_{sync280}$ were 0.855 and 0.372 under initial and $t_{irr}=60$ min conditions respectively. In contrast to high $SFI_{sync470}$ as being quite low $SFI_{sync280}$ could be considered as

insignificant. However, both $FI_{sync280}$ and $SFI_{sync280}$ displayed a slight decrease at the end of 60 min irradiation period most probably due to the degradation of released proteinaceous matter from *E. coli*.

 $FI \le 1.4$ is attributed to presence of humic fluorophores (Sen-Kavurmaci and Bekbolet, 2014). In the presence of HA solution as a reaction medium, FI of initial and t_{irr}=60 min was calculated as 1.08 and 1.10, respectively complying with the findings of researchers.

In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment were also followed and displayed in Figure 4.16.



Figure 4.16. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

EEM fluorescence contour plots displayed the presence of Regions III (λ_{exc} 220-250 nm and λ_{emis} 380-580 nm), and V (λ_{exc} 220-470 nm and λ_{emis} 380-580 nm) indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Presence of HA caused a masking effect over the Region I, II and IV owing to the quenching of fluorophores originating from protein like materials by humic-like materials (Wang et al., 2015). Following further irradiation periods

fulvic-like and humic-like content of the sample showed a decreasing trend in accordance with the solar photolytic degradation of HA and reduction of *E. coli*. In accordance with previous findings of researchers, obtained EEM fluorescence contour plots during solar photolytic traetment of *E. coli* in HA solution displayed an expected degradation trend (Birben et al., 2017a, 2021b; Uyguner-Demirel et al., 2020).

4.1.7. Mechanistic Evaluation of *E. coli* Inactivation Upon Solar Photolytic Treatment in HA Solution

As inactivation products upon solar photolytic treatment of *E. coli* in the presence of HA solution, total K, protein, and carbohydrate contents present under initial and t_{irr} =60 min conditions were presented in Table 4.8.

 Table 4.8. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon solar photolytic treatment.

Solar photolytic treatment	<i>E. coli</i> and HA			
Solai photolytic treatment	Initial	60 min		
Total K, mg/L	0.69	0.72		
Protein, mg/L	9.70	14.6		
Carbohydrate, mg/L	3.25	4.77		

In the absence of WM solution, low total K contents were detected similar to *E. coli* in IsoT solution solar photolytic treatment (Table 4.4). Total K content showed a slight increase after 60 min irradiation period, expressing released K⁺ ion amount from destructed *E. coli* cells. Besides the reduction of bacteria as 99.9 %, total K content did not show a significant increase at t_{irr} =60 min. The reason could be attributed to possible adsorption of released K⁺ ions from bacteria onto negatively charged, ruptured *E. coli* cell wall particles and negatively charged moieties of HA, then consequently eliminated with membrane filtration before total K analysis.

Due to the presence of HA solution, whose protein and carbohydrate concentrations were measured as 9.71 mg/L and 2.63 mg/L, respectively; initial contents of both protein (9.70 mg/L) and carbohydrate (3.25 mg/L) were higher than those achieved by solar photolytic treatment carried out in IsoT (Table 4.4) and WM solutions (Table 4.6). After t_{irr} =60 min, both protein and carbohydrate concentrations displayed an increase (14.6 mg/L for protein; 4.77 mg/L for carbohydrate) due to released protein and carbohydrate contents from damaged and/or destructed *E. coli* cells.

4.1.8. Organic Matter Formation and Removal Upon Solar Photolytic Treatment of *E. coli* in "HA and WM" Solution

Following a similar fashion, UV-vis absorption and synchronous scan fluorescence spectra of organic matter present in "HA and WM" solution were presented in Figures 4.17 and 4.18, respectively. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-550 nm due to the presence of aromatic domains, UV absorbing centers as well as color forming moieties (Color₄₃₆, UV₃₆₅, UV₂₈₀ and UV₂₅₄). Main difference could be visualized in between HA_i and "HA and WM" solution. Different from HA_i, HA solution together with WM solution ("HA and WM" solution) displayed a rather different absorbance features at λ < 240 nm with a shoulder between λ =200-210 nm. The reason could be attributed to the presence of NO₃⁻ in WM solution. Due to strong absorption spectrum of NO₃⁻, which having its λ_{max} at approximately 200 nm, solar *E. coli* inactivation conducted in WM solution containing reaction media expressed different absorption trend in comparison to the conditions in the presence of IsoT and HA solution (Krishnan and Guha, 1934; Edwards at el., 2001). Similar UV absorption pattern was also visualized via solar photolytic treatment of *E. coli* in sole WM solution (Figure 4.7). On the other hand, upon irradiation, minor variations could be recorded in wavelength range of 230-500 nm.



Figure 4.17. UV-vis absorption spectra of organic matter. Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed two intense peaks at λ_{emis} =390 nm and 470 nm, and a minor peak at λ_{emis} =280 nm, representing the presence of fluorophores originating from both humic matter and protein content of the solutions (Figure 4.18).

Moreover, both in "HA and WM" solution and after addition of bacteria into the reaction solution (initial), increased fluorescence intensities were observed relative to sole HA_i condition at $\lambda_{emis} < 430$ nm. An irradiation time dependent decreasing order in FI_{sync470} data was observed, except a minor increase between t_{irr}=20 min (35.3) and t_{irr}=30 min (36.4) conditions.

In the presence of HA solution, dominant $FI_{sync470}$ displayed changes in the range of 28.4-58.6 however $FI_{sync280}$ showed minor presence with variations in between 4.43-16.3 (Figure 4.19). On the other hand, $FI_{sync280}$ of HA_i solution could not be detected similar to previous HA solution conditions (Figure 4.14).



Figure 4.18. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

In presence of "HA and WM" solution, NPOC expressed changes in between 4.65-7.13 mg/L (Figure 4.19). In accordance with the UV₂₅₄ variations, NPOC content increased up to t_{irr} =10 min (4.65-6.55 mg/L) and slightly decreased upon t_{irr} =20 min (6.40 mg/L). Following further irradiation period of 30 min NPOC significantly increased (7.13 mg/L) and gradually decreased up to t_{irr} =60 min (5.29 mg/L). The highest NPOC release was attained upon irradiation period of 30 min.

Similar to previously presented solar photolysis of *E. coli* inactivation in sole HA solution (Figure 4.14), all UV-vis parameters expressed higher presence than treatments conducted in IsoT and WM solutions (Figures 4.4 and 4.9). As the most significant specified UV-vis parameter, UV_{254} expressed variations in the range of 0.4143-0.5050, being the highest for "HA and WM" solution and the lowest for t_{irr}=60 min condition. UV_{254} values were comparatively higher from UV_{280} as expected

under all conditions. On the other hand, Color₄₃₆ and UV₃₆₅ displayed evident presence due to the use of HA solution as a solution matrix component. However, specified UV-vis parameters did not show any significant difference between initial and t_{irr} =60 min conditions, revealing that there was no significant degradation of organic content and color forming moieties during solar light exposure period of 60 min. Similar findings were also recorded for solar photolytic treatment of *E. coli* in sole HA solution as well (Figure 4.14). In accordance with undetected FI_{sync280} of HA_i solution, SFI_{sync280} could not be calculated either.



Figure 4.19. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.20. As the most evident parameter, SUVA₂₅₄ displayed variations in between 5.88-10.8 L/m mg during 60 min of solar photolytic treatment, expressing differences in both UV₂₅₄ and NPOC removals. The highest and the lowest SUVA₂₅₄ were recorded for HA_i solution and t_{irr}=30 min condition, respectively. SUVA₂₈₀, which was the second significant specific UV-vis parameter calculated, showed differences in the range of 4.8-9.1 L/m mg. On the other hand, under all conditions, calculated SCoA₄₃₆ and SUVA₃₆₅ were comparatively lower than SUVA₂₈₀ and SUVA₂₅₄ as expected.

Furthermore, contrary to predominant $SFI_{sync470}$ varied in the range of 4.87-12.6, quite low $SFI_{sync280}$ expressed variations in the range of 0.937-2.38. Due to the use of HA solution as an aqueous reaction medium component, calculated $SFI_{sync470}$ values displayed significant presence contrary to $SFI_{sync280}$.



Figure 4.20. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

For comparative evaluation purposes, characteristic properties of organic matter under initial condition and upon solar photolytic inactivation of *E. coli* in "HA and WM" solution for 60 min of solar light exposure were presented in Table 4.9.

 Table 4.9. Characteristic properties of organic matter present in solar photolytic treatment of *E. coli* in "HA and WM" solution.

Irradiation time, min	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC,	SUVA ₂₅₄ ,	EI	SEI	EI
	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	Гl
Initial	0.0651	0.1385	0.3560	0.4302	6.29	6.84	51.3	8.15	1.18
60 min	0.0591	0.1291	0.3382	0.4143	5.29	7.83	28.4	5.37	1.23

Upon 60 min solar photolytic treatment of *E coli* in the presence of "HA and WM" solution, at which 2.43 log *E. coli* reduction was achieved (Table 4.2), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0591, UV₃₆₅: 0.1291, UV₂₈₀: 0.3382, UV₂₅₄: 0.4143, and NPOC as 5.29 mg/L.

SUVA₂₅₄ of initial and t_{irr} =60 min solutions were calculated as 6.84 L/m mg and 7.83 L/m mg, respectively. According to Edzwald and colleagues, SUVA₂₅₄ > 4 indicated the presence of hydrophobic and aromatic substances in solution (Edzwald et al., 1985). Therefore, it could be indicated that throughout the solar irradiation duration of 60 min, no significant degradation of organic content was recorded which could also be supported with data obtained from specified UV-vis measurements (Figure 4.19). Moreover, it could be concluded that when HA solution used with

WM solution, an inhibitory effect could be detected in both *E. coli* reductions and organic matter degradation in comparison to data obtained from solar photolytic inactivation of *E. coli* in sole HA solution (Tables 4.2, 4.7, and 4.9; Figures 4.1, 4.15, and 4.20).

 $FI_{sync470}$ displayed significant presence under initial condition due to the existence of HA solution and decreased nearly by half at t_{irr}=60 min via the degradation of organic content in the solution. $SFI_{sync470}$ expressed a decrease upon t_{irr}=60 min. On the other hand, initial and t_{irr}=60 min conditions of $FI_{sync280}$ were recorded as 12.5 and 9.85, respectively. According to $FI_{sync280}$ and NPOC data, $SFI_{sync280}$ of initial and t_{irr}=60 min runs were calculated as 1.98 and 1.86, respectively. Decreased $FI_{sync280}$ at t_{irr}=60 min could possibly be due to degradation of released protein content of *E. coli*. At t_{irr} =60 min, FI was calculated as 1.23 revealing the presence of humic fluorophores which comply with the presence of HA solution as a reaction matrix component. Similar findings were also reported by Birben and colleagues under similar experimental conditions (Birben et al., 2021b).

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in "HA and WM" solution during solar photolytic treatment were presented in Figure 4.21.



Figure 4.21. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

EEM fluorescence contour plots expressed the presence of Regions III, and V like *E. coli* inactivation in the presence of HA solution (Figure 4.16). Presence of HA solution caused a masking effect over the Region I, II and IV owing to the quenching of fluorophores originating from protein like materials by humic-like materials (Wang et al., 2015). Following further irradiation periods fulvic-like and humic-like content of the sample showed a decreasing trend in accordance with the solar photolytic degradation of humic matter and reduction of *E. coli*. Similar findings were reported by Uyguner-Demirel and colleagues under similar experimental conditions (Birben et al., 2021b; Uyguner-Demirel et al., 2020).

4.1.9. Mechanistic Evaluation of *E. coli* Inactivation Upon Solar Photolytic Treatment in "HA and WM" Solution

As inactivation products of solar photolytic *E. coli* inactivation in "HA and WM" solution, total K, protein, and carbohydrate contents of initial and t_{irr}=60 min solutions were presented in Table 4.10.

Solor photolytic treatment	<i>E. coli</i> in "HA and WM"			
Solar photolytic treatment	Initial	60 min		
Total K, mg/L	4.53	4.46		
Protein, mg/L	7.57	13.8		
Carbohydrate, mg/L	2.63	4.27		

Table 4.10. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon solar photolytic treatment.

Initial total K content was detected as higher than solar photolytic inactivation of *E. coli* in IsoT (0.72 mg/L), WM (3.92 mg/L), and HA (0.69 mg/L) solutions due to the presence of both "HA and WM" as matrix components (Tables 4.4., 4.6, and 4.8). Total K content of sole "HA and WM" solution was detected as 4.15 mg/L, therefore; for solar photolytic treatments conducted in "HA and WM" solution, total K concentration (4.15 mg/L) could be accepted as baseline value of the solution. After t_{irr} =60 min, total K content showed a slight decrease which could be related with low bacteria LRV as 2.43 (Table 4.2). It could be indicated that, excessive amount of K⁺ ion in the solution matrix could be used by bacteria to compensate K⁺ ion leakage and maintain inner osmotic pressure thus retardation of cell wall damage could be occurred.

Initial protein and carbohydrate concentrations of sole "HA and WM" solution were 9.40 mg/L and 3.78 mg/L, respectively. The reason of high initial protein (7.57 mg/L) and carbohydrate (2.63

mg/L) contents of *E. coli* in complex solution matrix could be attributed to the co-presence of "HA and WM" as solution components. Owing to released cytoplasmic organic matter from bacteria, and EPS present outside of the cell wall, *E. coli* in "HA and WM" solution upon solar photolytic treatment displayed increased protein and carbohydrate contents after irradiation period of 60 min as 13.8 mg/L and 4.27 mg/L, respectively. Besides high protein and carbohydrate contents of initial and solar light exposure duration of 60 min conditions, comparatively higher concentrations were obtained in the case of *E. coli* and sole HA solution exposed to solar photolytic treatment (Table 4.8).

4.1.10. Comparative Evaluation of Different Experimental Matrixes on Solar Photolytic Inactivation Mechanism of *E. coli*

Total K, protein, and carbohydrate concentrations, and bacteria enumeration results of solar photolytic treatment of *E. coli* in different solution matrixes in comparison with initial and t_{irr}=60 min conditions were given in Figure 4.22.



Figure 4.22. Initial and $t_{irr}=60$ min total K, protein, and carbohydrate contents, and bacteria enumeration results of *E. coli* inactivation upon solar photolytic treatment in various aqueous matrixes.

Total K contents expressed variations in between 0.69-4.53 mg/L being the lowest for initial condition of *E. coli* in HA solution and the highest for initial sample of *E. coli* in "HA and WM" solution matrix. Total K content displayed increased concentrations with the effect of present K⁺ ion content in WM solution. Total K concentrations were plotted against bacteria count data in order to detect if there is any correlation between these parameters, initial conditions showed a moderate

correlation, as R^2 =0.73, except the presence of IsoT solution, whereas t_{irr}=60 min runs displayed a strong correlation, as R^2 =0.99, excluding the presence of WM solution. Solar photolytic treatments carried out with IsoT and WM solutions showed lower protein contents both for initial and t_{irr}=60 min samples, and higher carbohydrate contents at t_{irr}=60 min than sole HA and "HA and WM" solutions.

Covering all conditions, protein contents varied in between 1.46-14.6 mg/L being the lowest for initial sample of *E. coli* in WM solution and the highest for t_{irr} =60 min condition of *E. coli* in HA solution. Similarly, carbohydrate contents varied in between 1.23-14.0 mg/L being the lowest for initial condition of *E. coli* in IsoT solution and the highest for t_{irr} =60 min solution of *E. coli* in WM solution. In the presence of HA solution, increased protein and decreased carbohydrate contents were detected at t_{irr} =60 min (14.6 mg/L protein and 4.77 mg/L carbohydrate for *E. coli* in HA solution; 13.8 mg/L protein and 4.27 mg/L carbohydrate for *E. coli* in HA solution; it could be attributed to the formation of HA and protein complexes due to their opposite charges at the pH of the solution (Tan et al., 2008). It could also be indicated that the presence of HA solution. Since, *E. coli* cell counts decreased in the presence of sole HA solution to 2.50E+02 CFU/mL and 2.63E+03 CFU/mL for *E. coli* in "HA and WM" solution upon irradiation period of 60 min. Accordingly, "HA and WM" condition affected *E. coli* inactivation more negatively than the presence of sole HA solution.

Released protein and carbohydrate concentrations and bacteria LRV of solar photolytic treatment of *E. coli* in different solution matrixes were given in Figure 4.23.



Figure 4.23. Released protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon solar photolytic treatment in various aqueous matrixes.

Released protein and carbohydrate contents were calculated by subtracting initial from concentrations obtained at tirr=60 min condition. Released protein content was detected as the lowest in E. coli in IsoT (1.80 mg/L), however it could be due to the degradation of released proteins throughout the 60 min irradiation period. While protein concentrations of the samples were directly proportional to HA contents, released protein concentrations did not display excessive differences in various solution matrixes. However, presence of HA solution showed inverse proportionality to released carbohydrate concentrations. It could be concluded that both carbohydrate concentrations after 60 min of irradiation period (4.77 mg/L for E. coli and HA solution; 4.27 mg/L for E. coli in "HA and WM" solution) and released carbohydrate concentrations (1.52 mg/L for E. coli and HA solution; 1.65 mg/L for E. coli in "HA and WM" solution) of the samples were inversely proportional to HA contents. Similar E. coli LRV calculated in the presence of IsoT (4.31) and WM (4.30) solutions however solar photolytic treatment conducted with sole HA (3.64) and "HA and WM" (2.43) solutions displayed decreased LRV of E. coli. In order to investigate the possible correlation between E. coli LRV and released organic content, both released protein and carbohydrate concentrations were plotted against LRV individually. Bacteria LRV and released protein concentration showed a strong correlation as $R^2=0.84$, except for WM solution, also LRV and released carbohydrate concentration expressed a strong correlation as R²=0.97, excluding the presence of sole HA solution.

Upon solar photolytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV_{280} and NPOC under initial and t_{irr} =60 min conditions were given in Figure 4.24. Protein concentrations and UV_{280} displayed an increased trend in the presence of HA solution contrary to, $FI_{sync280}$ showing reverse relation with HA content. Covering all conditions, $FI_{sync280}$ varied in between 6.46-39.0 being the lowest for t_{irr} =60 min sample of *E. coli* in HA solution and the highest for t_{irr} =60 min solution of *E. coli* in IsoT solution. Similarly, UV_{280} data displayed differences in the range of 0.0173-0.420 being the lowest for initial condition of *E. coli* in WM solution and the highest for initial sample of *E. coli* in HA solution. Lower $FI_{sync280}$ values were attained for either sole HA solution and "HA and WM" solution conditions, which could be derived from the masking effect of humic-like fluorophores over protein-like fluorophores due to the quenching of the fluorophores. Masking effect of fluorophores having HA origin over protein like materials were previously detected with EEM fluorescence contour plots (Figures 4.16 and 4.21).

Furthermore, protein concentrations were plotted against $FI_{sync280}$ data to evaluate any correlation between these two protein related parameters. As a result, a strong correlation, as $R^2 > 0.97$, was detected under all conditions except for the presence of sole WM solution.



Figure 4.24. Initial and $t_{irr}=60$ min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC and bacteria enumeration results upon solar photolytic treatment of *E. coli* in various aqueous matrixes.

NPOC contents were detected as higher than carbohydrate concentrations under all conditions, except *E. coli* in IsoT and WM solutions at t_{irr} =60 min ones (11.6 mg/L carbohydrate and 2.95 mg/L NPOC for *E. coli* in IsoT solution; 14.0 mg/L carbohydrate and 7.82 mg/L NPOC for *E. coli* in WM solution). The highest NPOC content (17.4 mg/L) was detected in *E. coli* in HA solution after t_{irr} =60 min of treatment. In comparison with sole HA solution, reduced *E. coli* inactivation, carbohydrate concentration and NPOC contents were recorded upon use of both "HA and WM" as solution matrix constituents. Therefore, it could be indicated that in the presence of "HA and WM" solution, solar photolytic process expressed selectivity towards degradation of humic matter rather than bacteria.

Moreover, carbohydrate concentrations were plotted against NPOC contents in order to detect if there is any correlation between these parameters. While initial conditions displayed a strong correlation as R^2 =0.99, excluding the presence of sole WM solution, t_{irr}=60 min runs showed a moderate correlation as R^2 =0.69, except the presence of "HA and WM" solution.

4.2. Solar Photocatalytic Inactivation of E. coli

4.2.1. TiO₂ Solar Photocatalytic Inactivation of E. coli

4.2.1.1. TiO₂ solar photocatalytic *E. coli* inactivation under specified reaction conditions, reductions, and kinetics. E. coli suspension in IsoT solution (No=7.95E+05 CFU/mL) was subjected to TiO2 solar photocatalysis. Upon irradiation period of 30 min, E. coli count decreased to 6.18E+02 CFU/mL. Following further exposure of $t_{irr}=60$ min, a drastic decrease was attained as < 10 CFU/mL. Under these conditions, LRV was 5.90 and reduction was 99.999 %. E. coli suspension in WM solution (N₀=6.41E+05 CFU/mL) treated for an irradiation period of 30 min, expressed a decrease of 2.50E+03 CFU/mL. Following light exposure of t_{irr}=60 min, E. coli count declined to 1.33E+02 CFU/mL revealing LRV as 3.73 and reduction as 99.9 %. E. coli (No=1.39E+06 CFU/mL) suspension was prepared in HA solution and exposed to irradiation for a period of 30 min, E. coli count was 5.66E+05 CFU/mL. Upon light exposure of tirr=60 min, E. coli count further decreased to 3.85E+04 CFU/mL revealing LRV as 1.45 and reduction as 90 %. E. coli (No=7.32E+05 CFU/mL) suspension was prepared in a solution composed of "HA and WM" components and was subjected to irradiation for 30 min, E. coli count decreased to 1.65E+05 CFU/mL. Upon light exposure of tirr=60 min, E. coli count decreased to 5.50E+04 CFU/mL revealing LRV as 1.08 and reduction as 90 %. Irradiation time dependent E. coli inactivation profiles in the presence of various solution matrixes were presented in Figure 4.25. First order kinetic model parameters rate constant (k, min⁻¹), half-life ($t_{1/2}$, min), and rate, (R, CFU/mL min) of TiO₂ solar photocatalytic inactivation of *E. coli* were presented in Table 4.11.



Figure 4.25. Irradiation time dependent *E. coli* inactivation upon TiO₂ solar photocatalytic treatment in various solution matrixes.

Based on these results, the effect of solution matrix could be displayed in terms of LRV of *E*. *coli* in an increasing order as: "HA and WM" < HA < WM < IsoT. TiO₂ solar photocatalytic treatment of *E*. *coli* in IsoT solution resulted in a rather fast reduction with k=0.248 min⁻¹ and half-life as 2.79 min. In "HA and WM" solution, a reaction rate constant of k=0.0488 min⁻¹ and half-life as 14.2 min were attained. The effect of reaction medium could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; IsoT > WM > HA > "HA and WM". Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) also followed the same trend as; 1.97E+05 > 9.44E+04 > 7.26E+04 > 3.22E+04 for IsoT, WM, HA, and "HA and WM", respectively.

Motrix	First or	IDV	Daduction 0/			
Maurix	k, min ⁻¹ $t_{1/2}$, min		R, CFU/mL min	LKV	Reduction, %	
<i>E. coli</i> in IsoT	0.248	2.79	1.97E+05	5.90	99.999	
<i>E. coli</i> in WM	0.132	5.25	9.44E+04	3.73	99.9	
<i>E. coli</i> and HA	0.0672	10.3	7.26E+04	1.45	90	
<i>E. coli</i> in "HA and WM"	0.0488	14.2	3.22E+04	1.08	90	

Table 4.11. TiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution showed the highest bacteria reduction percentage among other media as 99.999 %. Besides the combined effect of mechanistic stress factors as light and stirring, the addition of TiO₂ into the system provided higher LRV than *E. coli* in IsoT solution under photolytic conditions. The presence of TiO₂ particles caused external ROS formation and increased the detrimental effect of formed internal ROS.

TiO₂ solar photocatalytic treatment of *E. coli* in WM solution displayed comparatively lower percentage reduction than *E. coli* in IsoT solution. Lower bacteria reduction could be explained by the presence of common anions and cations in WM solution. As mentioned before in Section 2.5, WM components as mono and multivalent common anions (Cl⁻, NO₃⁻, SO₄²⁻, PO₄³⁻ and HCO₃⁻) could be adsorbed onto positively charged surface sites of TiO₂ thereby reduced the inactivation rate by decreased light absorption efficiency. Furthermore, their adsorption could create a negatively charge layer on the photocatalyst which causes the repulsion between TiO₂ and negatively charged *E. coli* (Abdullah et al., 1990; Bekbolet et al., 1998; Gogniat et al., 2006; Rincón and Pulgarin, 2004; Uyguner and Bekbolet, 2009; Uyguner-Demirel et al., 2013). Moreover, these anions could also be removed by *E. coli* intake via passive transport (Rincón and Pulgarin, 2004). Adsorbed anions onto the surface of TiO₂ hinders the surface interaction and reduces bacteria inactivation rate directly. Marugan and colleagues showed that inorganic matter present in the solution could serve as nutrients for the bacteria and help to maintain their survival as well (Marugán et al., 2010). In the presence of HA, a retardation effect was observed on *E. coli* decline expressed as a reduction of 90 % after 60 min of irradiation time. Decreased *E. coli* inactivation rate could be attributed to the presence of HA acting both as a ROS scavenger and a TiO₂ surface blocker, which could cause TiO₂ aggregation and prevent the contact of *E. coli* and photocatalyst (Birben et al., 2017a; Planchon et al., 2013). Under solution pH=6.5 conditions, HA possessed negatively charged centers due to deprotonation of carboxylic groups, TiO₂ surface was neutral due to the presence of equal number of oppositely charged sites (pH=pH_{zpc}) and as always *E. coli* surface was negative. As a consequence of surface interactions prevailing between HA and TiO₂, HA through reconformational changes inhibited the surface-active sites of photocatalyst which resulted in reduced surface interaction between *E. coli* cells and TiO₂ (Birben et al., 2021b). Similar findings were also presented by Marugán and colleagues (Marugán et al., 2008, 2010).

When reaction matrix changed to "HA and WM" solution, the lowest *E. coli* LRV was attained as 1.08. Combined effect of "HA and WM" resulted in decreased *E. coli* inactivation rates. These findings could be attributed to co-presence of HA and WM which played beneficial roles for bacteria as a nutrient source and increased the resistance towards photocatalysis. It could be concluded that TiO_2 expressed selectivity towards organic matrix rather than *E. coli* in matrixes containing HA (Birben et al., 2021b).

4.2.1.2. Organic matter formation and removal upon TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were illustrated in Figures 4.26 and 4.27. A general trend of a continuously declining absorbance with respect to increasing wavelength was attained under all irradiation periods. Besides significantly similar UV absorbance changes obtained from solar photolytic treatment of *E. coli* in IsoT solution (Figure 4.2), TiO₂ solar photocatalytic treatment of *E. coli* in the same aqueous matrix displayed quite low variations in the wavelength range of λ =205-240 nm excluding the minor presence of dense aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) (Figure 4.26). The highest absorption was detected between the stated wavelength range at t_{irr}=60 min. A very narrow wavelength region in between λ =210-220 nm revealed the presence of organic moieties with changing absorption upon increasing irradiation periods. On the other hand, absorbances detected at λ < 210 nm were characteristic to IsoT solution present as an aqueous medium (Tong et al., 2020).

Synchronous scan fluorescence spectra of emerged organic matter displayed a specific intensity peak at λ_{emis} =280 nm (Figure 4.27) followed by a tailing was noticed. FI_{sync} recorded at λ_{emis} =280 nm expressed a consistent irradiation time dependent decreasing order. The reason of the indicated

decrease in FI_{sync} could be related to the degradation of these fluorophores upon irradiation for the specified periods during TiO₂ photocatalysis. Moreover, the presence of fluorophoric groups expressing very minor intensities at $\lambda_{emis} \approx 470$ nm could also be encountered. On the other hand, contrary to minor presence of FI_{sync470} expressing very low intensities as 0.0857-0.695, predominant FI_{sync280} displayed changes in the range of 21.8-48.5 (Figure 4.28).



Figure 4.26. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.



Figure 4.27. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.

NPOC expressed variations in the range of 2.17-8.69 mg/L (Figure 4.28). NPOC content increased up to $t_{irr}=20 \text{ min } (2.17-8.08 \text{ mg/L})$ due to the release of organics from damaged and/or

destructed *E. coli* cells, followed by a sharp decrease at $t_{irr}=30 \text{ min} (3.35 \text{ mg/L})$ and again followed by an increasing trend during solar photocatalytic treatment. All UV-vis parameters displayed quite low values. The highest UV₂₅₄ displayed variations in the range of 0.0162-0.0287 the highest for $t_{irr}=10 \text{ min} (0.0287)$ and the lowest for $t_{irr}=60 \text{ min}$. Color₄₃₆ excluding t=0 and $t_{irr}=10 \text{ min}$, and UV₃₆₅ of $t_{irr}=20 \text{ min}$ and 60 min conditions were detected quite low as ≤ 0.0003 due to the non-existence and subsequent degradation of color forming moieties of the released organic matrix upon irradiation and the usage of IsoT solution as an aqueous matrix. Accordingly, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.29.



Figure 4.28. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.



Figure 4.29. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.

SUVA₂₅₄ displayed variations during TiO₂ photocatalysis in between 0.186-1.08 L/m mg, expressing differences in both UV₂₅₄ and NPOC removals. SUVA₂₈₀ expressed noticeable presence in the range of 0.091-0.791 L/m mg being the highest for t=0 and the lowest for t_{irr}=60 min. However, due to almost insignificant presence of Color₄₃₆ and UV₃₆₅, calculated SCoA₄₃₆ and SUVA₃₆₅ of initial, t_{irr}=20 min, 30 min, 40 min and 60 min conditions displayed quite low presence as \leq 0.009. Furthermore, SFI_{sync470} displayed quite insignificant presence in the range of 0.010-0.302 however SFI_{sync280} varied in the range of 2.51-22.3. The key fluorophoric group was accepted as SFI_{sync280}.

For simplicity purposes, based on the irradiation time dependent variations in UV-vis and fluorescence spectroscopic properties along with NPOC data, characteristic properties of organic matter under initial condition, upon introduction of TiO_2 (t=0) and TiO_2 solar photocatalytic treatment for irradiation period of 60 min were presented in Table 4.12.

Table 4.12. Characteristic properties of organic matter present in TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC,	SUVA ₂₅₄ ,	FI	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1 _{sync280}	SF1 _{sync280}	ГІ
Initial	0.0000	0.0002	0.0143	0.0209	2.17	0.963	48.5	22.3	1.64
t=0	0.0009	0.0026	0.0182	0.0248	2.30	1.08	45.8	19.9	1.64
60 min	0.0000	0.0000	0.0079	0.0162	8.69	0.186	21.8	2.51	1.06

No remarkable difference was detected between the initial and t=0 conditions of TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution revealing insignificant surface interactions. Upon t_{irr} =60 min at which 5.90 log reduction of *E. coli* in IsoT was attained (Table 4.11), released organic matter expressed the following UV-vis parameters (cm⁻¹); UV₂₈₀: 0.0079, UV₂₅₄: 0.0162, and NPOC as 8.69 mg/L. Color₄₃₆ and UV₃₆₅ of samples upon t_{irr} =60 min could not be noticed due to absence of color forming moieties in the structure of the organic matrix released to the aqueous medium. Quite high NPOC content at t_{irr} =60 min expressed very low SUVA₂₅₄ as 0.186 L/m mg representing that the remaining organic matter could still be expressing hydrophilic and aliphatic character (Edzwald et al., 1985). A decrease in FI_{sync280} was attained due to degraded protein content of *E. coli* debris. SFI_{sync280} displayed a decrease at t_{irr} =60 min in accordance with the increased NPOC content due to the released organic material from bacteria. On the other hand, under initial, t=0, and t_{irr} =60 min conditions, minor FI_{sync470} of initial, t=0, and t_{irr} =60 min conditions were calculated as 0.273, 0.302, and 0.010, respectively. In the absence of HA, FI_{sync470} and SFI_{sync470} were detected as considerably lower

than $\lambda_{emis}=280$ nm of the same parameters as expected. Insignificant presence of fluorophoric regions at $\lambda_{emis}=470$ nm indicated that the organic matrix was mainly composed of aromatic moieties that were attributed to the presence of aromatic amino acids as further evidenced by EEM fluorescence contour plots (Figure 4.30). Although reaction matrix did not contain any humic matter, FI of t_{irr}=60 min was calculated as 1.06. This very low FI might specify the fluorophores present in the remaining organic matter due to complete reduction of bacteria under the specified conditions.

Irradiation time dependent EEM fluorescence contour plots of organic matter emerged and removed during *E. coli* inactivation in IsoT solution by TiO_2 solar photocatalytic treatment were displayed in Figure 4.30.



Figure 4.30. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV indicating that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Throughout the irradiation periods, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of released organic matter and reduction of *E. coli*. Moreover, different from the solar photolytic treatment of *E. coli* in IsoT solution (Figure 4.6), fluorescence intensities were detected as lower than photolytic process.

<u>4.2.1.3.</u> Mechanistic evaluation of *E. coli* inactivation upon TiO_2 solar photocatalytic treatment in <u>IsoT solution</u>. Upon irradiation, as inactivation products of TiO_2 solar photocatalytic treatment of *E. coli* in IsoT solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.13.

Table 4.13. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in IsoT solution upon TiO₂ solar photocatalytic treatment.

TiO ₂ solar photocatalytic	E. coli in IsoT solution					
treatment	Initial	t=0	60 min			
Total K, mg/L	1.76	0.91	0.96			
Protein, mg/L	3.06	2.83	4.20			
Carbohydrate, mg/L	0.891	1.19	1.61			

Total K content showed an immediate adsorptive removal at t=0, as 48 %; and displayed almost no difference till the end of 60 min irradiation period. Stable total K content between t=0 and t_{irr}=60 min conditions could be due to both by adsorption of released K⁺ ions onto oppositely charged ruptured *E. coli* cell wall particles and/or onto the negatively charged sites of TiO₂ particles.

Initial protein content of *E. coli* in IsoT solution was determined as 3.06 mg/L. Zhang and colleagues reported that flagellar assembly of *E. coli* did not detach during photocatalytic process (Zhang et al., 2019). Even if it was not possible to fully detach flagellar assembly with solar photocatalytic treatment, adsorbed TiO₂ particles onto flagella, fimbriae or pili could cause splitting of these structures, which could be the cause of high initial protein concentrations. At the end of t_{irr} =60 min treatment period, both protein and carbohydrate contents displayed increasing trends. Protein contents were plotted against FI_{sync280} and NPOC data in order to detect if there could be any relationship between these parameters, and strong correlations were detected as R²=0.94 and R²=0.97,

respectively. Furthermore, a strong correlation was detected as R²=0.84 when carbohydrate contents plotted against NPOC data.

4.2.1.4. Organic matter formation and removal upon TiO₂ solar photocatalytic treatment of *E. coli* in WM solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in WM solution were illustrated in Figures 4.31 and 4.32. Although humic-like trend was observed under all irradiation periods, UV absorption was completely diminished at $\lambda > 250$ nm. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =205-260 nm lacking the dominant presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) that should be evident at longer wavelengths. On the other hand, absorbances detected at $\lambda < 210$ nm were characteristic to NO₃⁻ which was a component of WM solution (Krishnan and Guha, 1934; Edwards at el., 2001).



Figure 4.31. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm (Figure 4.32). An irradiation time dependent decreasing order in FI_{sync280} data was observed up to t_{irr}=40 min (40.8-15.9) and intensity showed a slight increase at t_{irr}=60 min (19.1) of solar photocatalytic treatment indicating that degraded fluorophores could be reemerged during specified irradiation period. Contrary to the synchronous scan fluorescence spectra attained in IsoT solution (Figure 4.27), the presence of fluorophores was evident at λ_{emis} > 320 nm displaying a tailing effect rather than a well-defined peak. In the absence of humic matter in the solution matrix, insignificant FI_{sync470} varying in between 0.206-2.98 were detected originating from released organics

from damaged *E. coli* cells. However, predominant $FI_{sync280}$ expressed changes in the range of 19.1-52.2 due to released aromatic protein content from bacteria (Figure 4.33). The main fluorophoric group was accepted as $FI_{sync280}$.



Figure 4.32. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

NPOC content increased up to $t_{irr}=30 \text{ min } (1.51-11.2 \text{ mg/L})$ with respect to *E. coli* inactivation (6.41E+05 CFU/mL for t=0; 2.50E+03 CFU/mL for $t_{irr}=30 \text{ min}$) resulting in release of organics from damaged and destructed *E. coli* cells, followed by a sharp decrease at $t_{irr}=40 \text{ min } (4.99 \text{ mg/L NPOC}; 5.17E+02 CFU/mL$ *E. coli*) and slightly increased during solar photocatalytic treatment (Figure 4.33).



Figure 4.33. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

All UV-vis parameters displayed minor presence however, upon 30 min of irradiation period at which bacteria count decreased to 2.50E+03 CFU/mL (6.41E+05 CFU/mL *E. coli* for t=0) (Figure 4.25), the highest values were attained as Color₄₃₆: 0.0047, UV₃₆₅: 0.0081, UV₂₈₀: 0.0317, UV₂₅₄: 0.0457, and NPOC: 11.2 mg/L due to released organic matter from damaged bacteria cells. In the non-existence and subsequent degradation of color forming moieties of the released organic matter upon irradiation and the usage of WM solution, which was also a colorless solution, attained Color₄₃₆ and UV₃₆₅ values were lower than UV₂₈₀ and UV₂₅₄. Following prolonged irradiation periods, a decreasing trend could be visualized for both specified UV-vis parameters and NPOC contents.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.34. As the most significant parameter, SUVA₂₅₄ displayed variations in the range of 0.408-2.03 L/m mg being the lowest for t_{irr} =30 min and the highest for initial conditions during TiO₂ solar photocatalytic treatment indicating differences in both UV₂₅₄ and NPOC removals. Moreover, SUVA₂₈₀ was recognized as the second significant specific UV-vis parameter varied in the range of 0.283-1.45 L/m mg being the lowest for t_{irr} =30 min and the highest for initial conditions.

Furthermore, besides very low presence of $SFI_{sync470}$ changing in between 0.0184-1.78, $SFI_{sync280}$ showed significant existence varying in the range of 2.05-33.2. As expected quite low $SFI_{sync470}$ were calculated in accordance with almost insignificant $FI_{sync470}$ it could be most probably due to absence of humic matter as a solution matrix component.



Figure 4.34. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

For comparison purposes, characteristic properties of organic matter under initial condition, upon introduction of TiO_2 (t=0) and at t_{irr}=60 min of TiO_2 solar photocatalytic inactivation of *E. coli* in WM solution medium were presented in Table 4.14.

Table 4.14. Characteristic properties of organic matter present in TiO₂ solar photocatalytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC, SU	SUVA ₂₅₄ ,	FI	SEI 200	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV_{254}	mg/L	L/m mg	1 ⁻¹ sync280	ST Isync280	1.1
Initial	0.0031	0.0040	0.0219	0.0307	1.51	2.03	50.2	33.2	1.46
t=0	0.0035	0.0042	0.0236	0.0311	1.67	1.86	52.2	31.3	1.46
60 min	0.0027	0.0049	0.0216	0.0330	5.13	0.643	19.1	3.72	1.90

Upon irradiation period of 60 min TiO₂ solar photocatalytic treatment at which 3.73 log reduction of *E. coli* in WM was attained (Table 4.11), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0027, UV₃₆₅: 0.0049, UV₂₈₀: 0.0216, UV₂₅₄: 0.0330, and NPOC as 5.13 mg/L. As expressed previously, the negligible presence of both color forming moieties and UV₃₆₅ were evidently noticed. Similar UV-vis spectroscopic parameters were recorded for initial, t=0 and t_{irr} =60 min irradiation periods revealing insignificant surface interactions.

As presented at Figure 4.34, all specific UV-vis parameters expressed quite low importance. But more importantly SUVA₂₅₄, which brings an insight to composition of the samples in terms of hydrophobicity and hydrophilicity, was calculated as 0.643 L/m mg at t_{irr} =60 min represented that the remaining organic matter could still be considered as having hydrophilic and aliphatic character (Edzwald et al., 1985).

 $FI_{sync280}$ decreased to 19.1 after 60 min of irradiation period (50.2 for initial) as a result of degraded protein content. $SFI_{sync280}$ also displayed a decrease at t_{irr} =60 min in accordance with the increased NPOC due to the released organic material from decaying bacteria cells. Although an inconsistent trend was attained (Figure 4.33), $FI_{sync470}$ were quite insignificant as 2.27, 2.98, and 0.289 for initial, t=0, and t_{irr} =60 min conditions, respectively. Due to the degradation of released organic content from damaged *E. coli* cells, $FI_{sync470}$ decreased at t_{irr} =60 min as expected. $SFI_{sync470}$ values of initial, t=0, and t_{irr} =60 min conditions were calculated as 1.50, 1.78, and 0.0563, respectively.

On the other hand, FI value calculated under $t_{irr}=60$ min period was 1.90 most probably due to the presence of microbially derived organic material due to the prevailing destructive reaction conditions.

In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution upon TiO_2 solar photocatalytic treatment were also followed and displayed in Figure 4.35.



Figure 4.35. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

EEM fluorescence counter plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all conditions in a decreasing manner with respect to increasing exposure to irradiation. Moreover, contrary to the solar photolytic (Figure 4.6) and TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution (Figure

4.30), regional fluorescence intensities were lower in the presence of aqueous medium composed of WM most probably due to lower UV₂₅₄ and NPOC contents (Table 4.3, 4.12 and 4.14).

4.2.1.5. Mechanistic evaluation of *E. coli* inactivation upon TiO_2 solar photocatalytic treatment in <u>WM solution</u>. As inactivation products of TiO_2 solar photocatalytic treatment of *E. coli* in WM solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min solutions were presented in Table 4.15.

TiO ₂ solar photocatalytic	E. coli in WM solution					
treatment	Initial	t=0	60 min			
Total K, mg/L	4.23	4.09	4.19			
Protein, mg/L	0.961	0.710	8.15			
Carbohydrate, mg/L	1.06	1.64	1.80			

Table 4.15. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon TiO₂ solar photocatalytic treatment.

Total K content displayed a similar trend with the solar photocatalytic treatment inactivation kinetics of *E. coli* in IsoT solution (Table 4.13); at t=0, total K showed a slight adsorptive removal as 3.3 % and displayed almost no difference upon irradiation period of 60 min. Stable total K content between t=0 and t_{irr}=60 min of irradiation duration could be caused from the adsorption of released K⁺ ions onto the ruptured *E. coli* cell wall particles having opposite surface charge and/or negatively charged sites of TiO₂ particles due to the preferential electrostatic interactions.

Upon TiO₂ solar photocatalytic treatment *E. coli* in WM solution, both protein and carbohydrate content expressed increased concentrations at the end of the 60 min treatment period. Dissimilar to the solar photolytic and photocatalytic treatment of *E. coli* in IsoT solution (Table 4.4 and 4.13), protein concentration showed a drastic increase after 60 min of irradiation (0.961 mg/L for initial; 8.15 mg/L for t_{irr}=60 min). Significantly increased protein content after 60 min of irradiation period was also detected in solar photolytic treatment of *E. coli* in WM solution (Table 4.6). Carbohydrate content did not display a significant increase at t_{irr}=60 min similar to IsoT solution condition (Table 4.13). However, it could be indicated that upon t_{irr}=60 min in IsoT and WM solutions, attained carbohydrate contents with the use of TiO₂ were higher than the ones obtained from solar photolytic treatment sets (Tables 4.4, 4.6 and 4.13). A strong correlation was detected as R²=1.00 between protein and FI_{sync280} as well as between protein and NPOC data. Carbohydrate contents were also plotted against NPOC and a moderate correlation was obtained as R²=0.50.

4.2.1.6. Organic matter formation and removal upon TiO₂ solar photocatalytic treatment of *E. coli* in HA solution. UV-vis absorption (Figure 4.36) and synchronous scan fluorescence (Figure 4.37) spectral features of organic matter in reaction medium were presented. It should be noted that the absorbance of HA_i could be visualized as following a rather different trend in $\lambda < 235$ nm region as expected. Since sample designated as "initial" represented "HA and *E. coli*" solution followed by 0.45 µm filtration, organic matter exudates from *E. coli* would also be encountered in the UV-vis spectra. Upon introduction of TiO₂ (t=0) into the solution, absorbances displayed an increase especially between wavelength range of λ =215 nm and 385 nm. The reason could be attributed to mechanistic effect of TiO₂ on *E. coli* cell wall, TiO₂ particles adsorbed on bacteria cells could cause cell wall damage and consequently release of cytoplasmic organic content to the reaction medium that could be resulting in increased UV light absorption.



Figure 4.36. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-500 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) originating mainly from the structural properties of HA present either as remaining unreacted sub-fractions or degraded under irradiation via oxidation reactions, and also originating from the organic matter released by destructed *E. coli* cells via photocatalysis. Due to overlapping absorbance behavior, no discrimination would be assessed in between the compositional variations. On the other hand, as different from solar photolytic treatment of *E. coli* in HA solution (Figure 4.12), irradiation time dependent sequential degradation of organic matter was achieved with TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed a high intensity peak at λ_{emis} =370 nm and two minor peaks at λ_{emis} =440 nm and 480 nm indicating the presence of aromatic domains (Dujmov and Sučević, 1990) (Figure 4.37). However, HA_i and initial conditions displayed a rather different trend. After the introduction of TiO₂ into the solution (t=0), a drastic increase in fluorescence intensities was detected. The reason could be attributed to immediate interaction of photocatalyst with HA and *E. coli* cell surface, which could cause a mechanistic harm to bacteria via cell wall damage or degrade some EPS onto the cell wall. Moreover, released extracellular and/or intracellular organic matter from bacteria via mechanistic action of TiO₂ could behave like "in-situ generated humic-like matter" and contribute to enhanced intensity peaks of aromatic domains (Uyguner-Demirel et al., 2018).

On the other hand, due to masking effect of humic fluorophores on protein fluorophores, measured intensities at λ_{emis} =280 nm were quite low. After t_{irr}=10 min, an irradiation time dependent decreasing order in FI_{sync} data was observed with a significant decrease after irradiation period of 30 min. Furthermore, FI_{sync470} and FI_{sync280} displayed changes in the range of 1.24-99.9 and 7.81-91.5, respectively (Figure 4.38). FI_{sync280} of HA_i could not be detected however after the addition of bacteria into HA solution i.e. the "initial" condition, FI_{sync280} increased to 7.81 indicating the presence of proteinaceous components in solution.



Figure 4.37. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

Along with the presence of HA solution, NPOC expressed considerable variations in the range of 4.78-20.9 mg/L (Figure 4.38). NPOC of sole HA solution (HA_i) and HA together with *E. coli*

(initial) was measured as 4.78 and 7.95 mg/L, respectively. After the introduction of bacteria, increased NPOC content was detected which could be due to mainly EPS of *E. coli*. At t=0 condition with the addition of TiO₂ into reaction medium, NPOC displayed a slight decrease as 6.59 mg/L. NPOC content increased from t_{irr}=10 min to 20 min (8.46-20.9 mg/L) due to the release of organics from damaged and/or destructed *E. coli* cells and along with the degradation of released cytoplasmic organic content from bacteria decreased at t_{irr}=30 min (13.2 mg/L), and showed another slight increase between t_{irr}=30 min and 40 min (15.2 mg/L), followed by a decreasing trend during solar photocatalytic treatment (5.68 mg/L).



Figure 4.38. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

All UV-vis parameters expressed high presence due to the use of HA solution as a reaction medium (Figure 4.38). After t=0 condition, all specified UV-vis parameters followed a sequential decrease in comparison to solar photolytic treatment of *E. coli* in HA solution (Figure 4.14), which indicates the effectiveness of TiO₂ onto degradation of organic content in the solution. As the most significant parameter, UV_{254} showed changes in the range of 0.0509-0.5397, being the highest for t=0 condition and the lowest for t_{irr}=60 min condition. Moreover, at t_{irr}=60 min Color₄₃₆ was measured as 0.0033 due to subsequent degradation of color forming moieties originating from humic matter and released cytoplasmic organic content of bacteria.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.39. SUVA₂₅₄ displayed variations in between 0.722-10.5 L/m mg during TiO₂ solar photocatalytic treatment showing differences in both UV₂₅₄ and

NPOC removals. On the other hand, due to almost insignificant presence of $Color_{436}$ (0.0033) and UV_{365} (0.0081) at t_{irr}=60 min (Figure 4.38), calculated SCoA₄₃₆ and SUVA₃₆₅ displayed quite low presence as 0.039 and 0.118 L/m mg, respectively.



Figure 4.39. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

Furthermore, $SFI_{sync470}$ and $SFI_{sync280}$ expressed variations in the range of 0.337-15.2 and 0.982-10.3, respectively. In comparison to solar photolytic inactivation of *E. coli* in HA solution (Figure 4.15) relatively high $SFI_{sync280}$ values were calculated in the presence of TiO_2 as a bare photocatalyst specimen. According to undetected $FI_{sync280}$ of HA_i, $SFI_{sync280}$ could not be calculated either. Although $SFI_{sync280}$ of initial sample (0.982) and $SFI_{sync470}$ of irradiation period of 60 min condition (0.337) could be regarded as insignificant.

For comparative evaluation purposes, characteristic properties of organic matter under initial condition, upon introduction of TiO_2 (t=0) and at the end of TiO_2 solar photocatalytic inactivation of *E. coli* in HA solution for 60 min of solar light exposure were presented in Table 4.16.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC, SUVA	SUVA ₂₅₄ ,	EI	SEI	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1 _{sync470}	SF1 _{sync470}	ГІ
Initial	0.0724	0.1571	0.4017	0.4787	7.95	6.02	56.8	7.14	1.17
t=0	0.0819	0.1857	0.4738	0.5397	6.59	8.19	99.9	15.2	1.12
60 min	0.0033	0.0081	0.0351	0.0509	5.68	0.896	1.24	0.218	1.29

Table 4.16. Characteristic properties of organic matter present in TiO₂ solar photocatalytic treatment of *E. coli* in HA solution.

Upon 60 min of TiO₂ solar photocatalytic treatment of *E coli* in the presence of HA solution at which 1.45 log *E. coli* reduction was achieved (Table 4.11), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0033, UV₃₆₅: 0.0081, UV₂₈₀: 0.0351, UV₂₅₄: 0.0509, and NPOC as 5.68 mg/L. Due to degradation of both already present organic content originating from used HA solution and emerged organic matter from damaged bacteria, all specified UV-vis parameters and NPOC displayed a decrease after 60 min of irradiation period.

SUVA₂₅₄ of initial, t=0, and t_{irr}=60 min conditions were calculated as 6.02, 8.19, and 0.896 L/m mg, respectively. Edzwald and colleagues reported that SUVA₂₅₄ < 3 indicates the presence of hydrophilic and aliphatic content however SUVA₂₅₄ > 4 reveals the existence of hydrophobic and aromatic substances in solution (Edzwald et al., 1985). Therefore, it could be indicated that initial and t=0 solutions displayed hydrophobic and aromatic features while t_{irr}=60 min condition expressed hydrophilic and aliphatic characteristics due to degradation of aromatic organics throughout 60 min of solar photocatalytic treatment process. The use of HA solution as reaction medium caused relatively high NPOC and SUVA₂₅₄ at the beginning of the solar photocatalytic treatment (for initial and t=0 conditions) as expected.

 $FI_{sync470}$ displayed a significant increase between the initial (56.8) and t=0 (99.9) conditions and decreased at t_{irr} =60 min (1.24) owing to degradation of organic content. According to recorded $FI_{sync470}$ and NPOC data, $SFI_{sync470}$ was calculated as 7.14, 15.2, and 0.218 for initial, t=0 and t_{irr} =60 min solutions, respectively. Moreover, $FI_{sync280}$ of initial, t=0 and t_{irr} =60 min conditions were recorded as 7.81, 68.2, and 18.2, respectively. Similar with $FI_{sync470}$, $FI_{sync280}$ demonstrated a drastic increase between initial and t=0 samples, and due to degradation of released protein content resulted in a significant decrease in $FI_{sync280}$ at t_{irr} =60 min. $SFI_{sync280}$ of initial, t=0, and t_{irr} =60 min runs were calculated as 0.982, 10.3, and 1.96, respectively.

In the presence of HA, quite low $FI_{sync280}$ and $SFI_{sync280}$ values were detected in comparison with solar photolytic and photocatalytic treatment of *E. coli* in IsoT solution and WM conditions (Tables 4.3, 4.5, 4.12 and 4.14). It could be indicated that masking effect of HA on protein content could be the main cause of detected low $FI_{sync280}$ and $SFI_{sync280}$ values.

Furthermore, FI of initial, t=0, and t_{irr} =60 min conditions were calculated as 1.17, 1.12, and 1.29, respectively. According to literature, FI \leq 1.4 is attributed to presence of humic fluorophores (Sen-Kavurmaci and Bekbolet, 2014). All calculated FI values indicated the presence of fluorophores originating from present HA solution matrix as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in HA solution by TiO_2 solar photocatalytic treatment were displayed in Figure 4.40.



Figure 4.40. EEM fluorescence contour plots of organic matter. Irradiation time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like organic fractions. Although microbial by-products and protein-like fluorophores were also still evident in solution, the presence of humic-like components might largely quenched protein-like ones (Wang et al., 2015). Following further irradiation periods, fulvic-like and humic-like fluorophores showed a decreasing trend in accordance with the solar photocatalytic degradation of HA, and reduction of *E. coli* counts.
However, alongside almost removed fluorescence intensities of Regions III, and V; Region I, which represents the presence of aromatic protein content, expressed remaining fluorescence intensities after 60 min of irradiation period. The reason could be attributed to remained high amount of *E. coli* (3.85E+04 CFU/mL) (Figure 4.25) in the solution after selected irradiation period. On the other hand, beside reduced intensities of EEM fluorescence contour plots at Regions III, and V, degradation of HA content could also be followed via UV-vis absorbance spectra (Figure 4.36), synchronous scan fluorescence spectra (Figure 4.37), NPOC contents (Figure 4.38).

<u>4.2.1.7. Mechanistic evaluation of *E. coli* inactivation upon TiO_2 solar photocatalytic treatment in <u>HA solution</u>. As a result of bacteria inactivation upon TiO_2 solar photocatalytic treatment of *E. coli* in HA solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.17.</u>

Table 4.17. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon TiO₂ solar photocatalytic treatment.

TiO ₂ solar photocatalytic	E. coli and HA				
treatment	Initial	t=0	60 min		
Total K, mg/L	0.53	0.49	0.66		
Protein, mg/L	13.4	11.0	3.75		
Carbohydrate, mg/L	2.77	2.92	2.24		

Total K content showed an immediate adsorptive removal as 7.5 % along with the addition of photocatalyst into the solution (t=0) and slightly increased at t_{irr} =60 min condition expressing K⁺ ion release from damaged bacteria cells. Detected low total K contents displayed similarity with solar photolytic treatment of *E. coli* in HA solution (Table 4.8), therefore it could be concluded that the presence of HA solution caused a significant inhibition of total K content detection.

Previous solar photolytic and photocatalytic treatments carried out with different solution matrixes (Tables 4.4, 4.6, 4.8, 4.10, 4.13, and 4.15) showed increased protein and carbohydrate concentrations after t_{irr} =60 min however TiO₂ solar photocatalytic treatment of *E. coli* in the presence of HA solution displayed decreasing trends for both protein (13.4-3.75 mg/L) and carbohydrate (2.77-2.24 mg/L) contents. Protein content showed a significant decrease (72 %) while carbohydrate concentration displayed almost no change (< 1 %). Both protein concentrations and FI_{sync280} (Figure 4.38), and carbohydrate and NPOC contents (Table 4.16) showed no correlation with each other. On the other hand, when protein concentrations plotted against NPOC a strong correlation was obtained

as R^2 =0.85. It could be indicated that decreasing trend of NPOC were related to the preferential decrease of protein content in comparison to carbohydrate content.

4.2.1.8. Organic matter formation and removal upon TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.41 and 4.42. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) mainly originating from humic sub-fractions. UV-vis absorption spectra showed a drastic decrease after the introduction of photocatalyst into the reaction medium (t=0), which could be due to immediate adsorptive removal of present aromatic domains and color forming moieties via membrane filtration prior to UV-vis spectroscopic measurements. Similar to TiO₂ solar photocatalytic inactivation of *E. coli* in HA solution, irradiation time dependent sequential degradation of organic matter was achieved starting from t_{irr}=10 min (Figure 4.36). On the other hand, due to the use of NO₃⁻ containing WM solution as a component of aqueous reaction medium, different absorbance characteristics were observed at λ < 240 nm as expected.



Figure 4.41. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intensity peak at λ_{emis} =370 nm for t=0 and λ_{emis} =470 nm for HA_i, "HA and WM", initial and t_{irr}=10 min conditions (Figure 4.42). All irradiation period scans demonstrated minor peaks at λ_{emis} =280 nm, except HA_i and "HA and WM" solution. Obtained synchronous scan fluorescence spectra of HA_i was characteristic

to HA solution and in resemblance with the findings of Uyguner and Bekbolet (Uyguner and Bekbolet, 2005). Moreover, between wavelength range λ_{emis} =300-480 nm, various peaks could be visualized indicating the presence of aromatic domains in solutions (Dujmov and Sučević, 1990). The peaks obtained were in accordance with the presence and absence of the bacteria in the reaction medium. After introduction of TiO₂ into reaction solution, a significant increase in FI_{sync} data was observed, which could be originating from released cytoplasmic organic content of bacteria due to the mechanistic effect of TiO₂ on cell wall structure. From a general perspective, an irradiation time dependent decreasing order in FI_{sync470} data was observed.

On the other hand, dominant $FI_{sync470}$ and comparatively low $FI_{sync280}$ displayed changes in the range of 2.78-186.3 and 4.20-63.6, respectively (Figure 4.43). $FI_{sync280}$ of HA_i could not be detected however $FI_{sync280}$ obtained in "HA and WM" solution was detected as 4.20. Along with the presence of HA solution as an aqueous matrix component, the main fluorophoric group was recognized as $FI_{sync470}$.



Figure 4.42. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

NPOC expressed variations in the range of 4.78-21.5 mg/L (Figure 4.43). NPOC content slightly decreased between HA_i (4.78 mg/L) and "HA and WM" (4.38 mg/L) solution, after the addition of bacteria into the solution (initial) slightly increased (7.88 mg/L) and displayed 26.5 % immediate adsorptive removal after the addition of photocatalyst into the reaction matrix (t=0) (5.79 mg/L). At t_{irr} =10 min, NPOC showed a significant increase (16.5 mg/L) and followed by a decreasing trend up

to $t_{irr}=30 \text{ min (12.3 mg/L)}$, afterwards showed a sharp increase at $t_{irr}=40 \text{ min (21.5 mg/L)}$ and decreased until the end of solar photocatalytic treatment of 60 min (10.5 mg/L).

All UV-vis parameters expressed high presence due to the use of HA solution as a reaction medium component (Figure 4.43). Except t=0 condition, all specified UV-vis parameters followed a sequential decrease in comparison to solar photolytic treatment of *E. coli* in "HA and WM" solution (Figure 4.19). Similar to solar photocatalytic inactivation of *E. coli* in sole HA solution in the presence of TiO₂ (Figure 4.38), the use of TiO₂ expressed quite effective organic matter removal in comparison to solar photolytic treatment (Figure 4.19). As the most significant parameter, UV₂₅₄ showed changes in the range of 0.1035-0.5000, being the highest for HA_i solution and the lowest for t_{irr}=60 min condition. According to undetected FI_{sync280} of HA_i solution, its SFI_{sync280} could not be calculated either.



Figure 4.43. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

Based on NPOC contents, all specific parameters as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.44. As the most apparent parameter, SUVA₂₅₄ displayed variations in between 0.730-10.5 L/m mg during TiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals. Furthermore, due to quite low presence of Color₄₃₆ (0.0069) and UV₃₆₅ (0.0175) at t_{irr}=60 min condition (Figure 4.43), calculated SCoA₄₃₆ and SUVA₃₆₅ displayed quite low presence as 0.066 and 0.167, respectively. On the other hand, SFI_{sync470} and SFI_{sync280} expressed changes in the range of 0.265-39.0 and 0.871-11.0, respectively. The key fluorophoric group was recognized as SFI_{sync470} in the presence of HA solution.



Figure 4.44. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

For comparative evaluation purposes, characteristic properties of organic matter present under initial condition, upon introduction of TiO_2 (t=0) and at the end of TiO_2 solar photocatalytic inactivation of *E. coli* in "HA and WM" solution for 60 min of solar light exposure were presented in Table 4.18.

Table 4.18. Characteristic properties of organic matter present in TiO2 solar photocatalytic treatmentof *E. coli* in "HA and WM" solution.

Irradiation	UV-vis S	Spectroscop	pic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	TI	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	ГІ
Initial	0.0664	0.1409	0.3614	0.4351	7.88	5.52	52.1	6.61	1.23
t=0	0.0248	0.0601	0.1867	0.2229	5.79	3.85	61.7	11.0	1.28
60 min	0.0069	0.0175	0.0734	0.1035	10.5	0.986	2.78	0.265	1.31

Upon irradiation period of 60 min TiO₂ solar photocatalytic treatment of *E coli* in "HA and WM" solution, 1.08 log *E. coli* reduction was achieved (Table 4.11), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0069, UV₃₆₅: 0.0175, UV₂₈₀: 0.0734, UV₂₅₄: 0.1035, and NPOC as 10.5 mg/L. SUVA₂₅₄ value at initial and t_{irr}=60 min conditions were detected as 5.52 and 0.986 L/m mg, respectively. Obtained SUVA₂₅₄ complied with the content of the solution matrix and indicate the degradation of organics at the end of 60 min irradiation period. The use of HA as solution matrix caused relatively high initial NPOC and SUVA₂₅₄ as expected. NPOC content of t_{irr}=60 min showed an increase due to HA content and released organic material from bacteria. Obtained FI_{sync280} of initial, t=0 and t_{irr}=60 min solutions were detected as 14.4, 63.6, and 21.8,

respectively. $FI_{sync470}$ and $FI_{sync280}$ displayed similar patterns with treatment conducted in sole HA solution (Figure 4.38 and Table 4.16), with an increase at t=0 and a significant decreased at t_{irr}=60 min conditions. Furthermore, $SFI_{sync280}$ of initial, t=0, and t_{irr}=60 min runs were calculated as 1.83, 11.0, and 2.08, respectively. FI at t_{irr}=60 min was calculated as 1.31 indicating the presence of fluorophores originating from HA containing solution matrix as expected.

EEM fluorescence counter plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO_2 solar photocatalytic treatment were displayed in Figure 4.45.



Figure 4.45. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

EEM fluorescence counter plots displayed the presence of Regions III, and V like solar photolytic E. coli inactivation in "HA and WM" solution (Figure 4.21). Presence of HA solution caused a masking effect over the Region I, II and IV owing to the quenching of the fluorophores of protein like materials by humic-like materials (Wang et al., 2015). Following further irradiation periods, fulvic-like and humic-like components of the sample showed a decreasing trend in accordance with the solar photocatalytic degradation of HA and reduction of E. coli (Figures 4.25 and 4.43). EEM fluorescence contour plots of E. coli in the presence of both HA and WM components showed similarities with solar photocatalytic *E. coli* inactivation in sole HA solution (Figure 4.40). Cell enumerations of E. coli in HA solution (3.85E+04 CFU/mL), and E. coli in "HA and WM" solution (5.55E+04 CFU/mL) after 60 min of irradiation were detected as relatively close to each other although HA contents showed differences (Table 4.16 and 4.18). Following treatment of E. coli in "HA and WM" solution, less HA degradation was observed than sole HA conditions which could also be followed via reduced intensities at Region III and V of EEM contour plots, UV-vis absorption spectra, and synchronous scan fluorescence spectra (Figures 4.36, 4.37 and 4.40). It could be attributed that the presence of WM solution as an experimental media content could cause the formation of WM solution dependent regional profiles in EEM fluorescence contour plots. Recorded data showed that retardation effect exerted by WM components was only perceived by HA rather than directly on *E. coli* most probably due to repulsive electrostatic forces.

<u>4.2.1.9. Mechanistic evaluation of *E. coli* inactivation upon TiO_2 solar photocatalytic treatment in <u>"HA and WM" solution.</u> As inactivation products of TiO_2 solar photocatalytic treatment of *E. coli* in "HA and WM" solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.19.</u>

TiO ₂ solar photocatalytic	E. coli in "HA and WM"					
treatment	Initial	t=0	60 min			
Total K, mg/L	4.51	4.49	4.07			
Protein, mg/L	13.9	7.83	5.72			
Carbohydrate, mg/L	3.93	2.24	1.48			

Table 4.19. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon TiO₂ solar photocatalytic treatment.

Total K content showed a slight decrease between initial and $t_{irr}=60$ min conditions. An insignificant adsorptive removal of total K (< 0.5 %) was attained at t=0 condition. Total K decrease at the end of 60 min irradiation period could be related to the low bacteria LRV as 1.08 (Table 4.11).

It could be indicated that, *E. coli* cells remaining intact and/or having undamaged cell walls after 60 min treatment period could not display K^+ ion release. Moreover, additional K^+ ion contents originating from WM solution, could compensate released K^+ ion content via leakage from the cells, and thereby protect bacteria from destruction via the retardation of cell wall damage.

Upon TiO₂ solar photocatalytic treatment, *E. coli* in "HA and WM" solution displayed a decrease in both protein (13.9-5.72 mg/L) and carbohydrate (3.93-1.48 mg/L) contents at the end of t_{irr} =60 min period. Similar findings were also detected from TiO₂ solar photocatalytic treatment of *E. coli* in sole HA solution (Table 4.17). Contrary to the data of *E. coli* in HA solution, NPOC increased after t_{irr} =60 min for *E. coli* in "HA and WM" solution (Tables 4.16 and 4.18). Moreover, EEM counter plots attained at t_{irr} =60 min displayed the presence of humic-like and fulvic-like fluorophores (Figure 4.45), therefore; it could be concluded that EEM counter plots and NPOC data correlated well with each other instead of protein/carbohydrate contents and NPOC. No correlation was detected either between FI_{sync280} and protein contents or between NPOC and protein/carbohydrate concentrations.

4.2.1.10. Comparative evaluation of different experimental matrixes on TiO₂ solar photocatalytic inactivation mechanism of E. coli. As TiO2 solar photocatalytic inactivation products, total K, protein, and carbohydrate concentrations along with bacteria enumerations in different solution matrixes were compiled from Figure 4.25, Tables 4.13, 4.15, 4.17, 4.19 and presented in Figure 4.46. Total K contents varied in between 0.49-4.51 mg/L being the lowest for t=0 sample of E. coli in HA solution and the highest for initial sample of E. coli in "HA and WM" solution. Total K content in WM solution displayed higher presence as expected due to the existence of medium constituents. No significant difference of total K contents was determined in between initial, t=0, and t_{irr}=60 min conditions upon solar photocatalysis of E. coli in WM, HA and "HA and WM" matrixes. Protein and carbohydrate contents of E. coli in IsoT solution indicated increased values at tirr=60 min (4.20 mg/L protein; 1.61 mg/L carbohydrate) contrary to the decrease attained in total K content (0.96 mg/L). Furthermore, total K contents were plotted against E. coli cell counts in order to detect if there could be any correlation between these parameters, initial and t=0 conditions showed a strong correlation, as $R^2 > 0.83$, except the presence of IsoT solution, however t_{irr}=60 min runs displayed no correlation between these two parameters. Covering all conditions protein contents varied in between 0.710-13.9 mg/L being the lowest for t=0 sample of E. coli in WM solution and the highest for initial sample of E. coli in "HA and WM" condition. Similarly, carbohydrate contents varied in between 0.89-3.93 mg/L being the lowest for initial condition of E. coli in IsoT solution and the highest for initial sample of E. coli in "HA and WM" solution.



Figure 4.46. Initial, t=0, and t_{irr} =60 min total K, protein, carbohydrate contents, and bacteria enumeration results of *E. coli* inactivation upon TiO₂ solar photocatalytic treatment in various aqueous matrixes.

The highest protein concentration at $t_{irr}=60$ min was determined in the presence of WM components (8.15 mg/L). While protein contents of *E. coli* in WM solution initial (0.96 mg/L) and t=0 min (0.71 mg/L) conditions revealed lowest ones, at $t_{irr}=60$ min the highest protein release was detected. In sole IsoT solution, initial (0.89 mg/L) and t=0 (1.19 mg/L) conditions displayed lower carbohydrate contents than other experimental conditions. Contrary to the inactivation profiles in IsoT and WM solutions expressing increasing trend in both protein and carbohydrate contents after 60 min irradiation, samples containing sole HA solution displayed decreased organic matter content. It could be indicated that, the presence of HA solution caused an enhancement for both protein and carbohydrate contents of initial and t=0 conditions. *E. coli* inactivation at the end of 60 min irradiation time decreased in the presence of HA solution, and the use of sole HA solution as reaction matrix affected *E. coli* inactivation more negatively than the presence of "HA and WM" solution.

Released and degraded protein and carbohydrate contents and bacteria LRV of TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes were given in Figure 4.47. Released protein and carbohydrate concentrations of *E. coli* in IsoT and WM conditions were calculated by subtracting initial data from the concentrations obtained at t_{irr} =60 min condition (Figure 4.47 Left). While released protein and carbohydrate concentrations of IsoT and WM matrixes could be calculated, in HA and "HA and WM" solutions, degradation of protein and carbohydrate contents should also be encountered. Released protein content was lower for *E. coli* in IsoT solution (1.14 mg/L). However, almost no difference was detected between released carbohydrate contents for both IsoT (0.72 mg/L) and WM (0.74 mg/L) solutions. LRV of *E. coli* was inversely proportional to protein release for IsoT (5.9) and WM (3.7) matrixes. No direct correlation was detected between bacteria LRV and released protein and carbohydrate contents for *E. coli* in IsoT and WM solution conditions.



Figure 4.47. Released (Left) and degraded (Right) protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon TiO₂ solar photocatalytic treatment in various aqueous matrixes.

Degraded protein and degraded carbohydrate concentrations of *E. coli* in HA and "HA and WM" conditions were calculated by subtracting detected concentrations of t_{irr} =60 min conditions from initial condition data (Figure 4.47 Right). Solar photocatalytic treatments containing HA solution showed a decrease of protein and carbohydrate contents. It could be due to extra organic matter coming from HA and high bacteria survival. Due to low *E. coli* inactivation, protein and carbohydrate content arising from bacteria were inhibited, therefore it could be indicated that the protein and carbohydrate contents (Figure 4.23), it could be attributed to the use of TiO₂ showing selectivity towards organic matrix rather than bacteria. It could be concluded that LRV of *E. coli* is directly proportional to protein degradation but inversely proportional to carbohydrate degradation.

Upon solar photocatalytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV_{280} , and NPOC under initial, t=0, and t_{irr} =60 min conditions were given in Figure 4.48. Protein concentrations and UV_{280} displayed an increase in the presence of HA solution, like solar photolytic treatment (Figure 4.24). Similar data were obtained for $FI_{sync280}$ of initial and t=0 conditions in IsoT and WM matrixes. The highest $FI_{sync280}$ (68.2) and UV_{280} (0.4738) were measured in HA solution upon introduction of TiO₂ (t=0) into the solution matrix. A strong correlation was detected between protein and $FI_{sync280}$ data as $R^2 > 0.81$ except for sole HA solution.



Figure 4.48. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and bacteria enumeration results of TiO₂ solar photocatalytic treatment of *E. coli* in various aqueous matrixes.

NPOC contents were detected as higher than carbohydrate concentrations for all conditions. The highest NPOC content (10.5 mg/L) and the lowest *E. coli* inactivation efficiency (5.55E+04 CFU/mL) was obtained in case that "HA and WM" solution was used upon $t_{irr}=60$ min exposure. The second high NPOC was detected in IsoT solution at $t_{irr}=60$ min (8.69 mg/L) with complete bacteria inactivation signified by LRV as 5.90 (Table 4.11).

It could be concluded that final (t_{irr} =60 min) NPOC content of "HA and WM" solution mostly originated from HA present either as remaining unreacted and/or degraded sub-fractions under irradiation via oxidation reactions rather than *E. coli* inactivation. Moreover, a strong correlation between carbohydrate and NPOC contents as R² > 0.82 were detected, excluding the presence of sole WM solution.

4.2.2. SynTiO₂ Solar Photocatalytic Inactivation of E. coli

4.2.2.1. SynTiO₂ solar photocatalytic *E. coli* inactivation under specified reaction conditions, reductions, and kinetics. *E. coli* suspension (N₀=4.37E+05 CFU/mL) was prepared in IsoT solution and was subjected to solar photocatalytic treatment using SynTiO₂. Upon irradiation period of 30 min, *E. coli* count decreased to 1.90E+02 CFU/mL. Following light exposure of t_{irr} =60 min, complete *E. coli* inactivation was detected. Under these conditions LRV was 5.64 and reduction was 99.999 %. Upon preparation of *E. coli* suspension in WM solution expressing an initial count as N₀=6.33E+05 CFU/mL, for an irradiation period of 30 min, *E. coli* count decreased to 6.00E+02 CFU/mL. Upon light exposure of t_{irr} =60 min, *E. coli* count decreased to 2.85E+01 CFU/mL revealing LRV as 4.35 and reduction as 99.99 %. *E. coli* (N₀=1.69E+05 CFU/mL) suspension was prepared in HA solution for an irradiation period of 30 min, *E. coli* count decreased to 1.67E+02 CFU/mL. Following light exposure of t_{irr} =60 min, *E. coli* count reached to 3.35E+01 CFU/mL revealing LRV as 3.70 and reduction as 99.99 %. *E. coli* (N₀=5.31E+05 CFU/mL) suspension was prepared in a solution of "HA and WM" components and was subjected to an irradiation period of 30 min in which *E. coli* count decreased to 2.21E+03 CFU/mL. Upon light exposure of t_{irr} =60 min, *E. coli* count 4.67E+02 CFU/mL revealing LRV as 3.06 and reduction as 99.9 %.

Based on these results, the effect of solution matrix could be displayed in terms of LRV of *E*. *coli* in an increasing order as: "HA and WM" < HA < WM < IsoT. *E. coli* inactivation profiles and first order kinetic parameters were presented in Figure 4.49 and Table 4.20.



Figure 4.49. Irradiation time dependent *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in various solution matrixes.

SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution resulted in a rather fast reduction with k=0.254 min⁻¹ and half-life as 2.73 min. In the presence of "HA and WM", a reaction rate constant of k=0.117 min⁻¹ and half-life as 5.92 min were attained. The effect of reaction medium could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; IsoT > WM > HA > "HA and WM". Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) were calculated as follows; 1.14E+05 > 1.11E+05 > 6.21E+04 > 2.54E+04 for WM, IsoT, "HA and WM", and HA, respectively.

Matrix	First o	IDV	Reduction, %		
Maurix	k, min ⁻¹ $t_{1/2}$, min R, CFU/mL min				LKV
E. coli in IsoT	0.254	2.73	1.11E+05	5.64	99.999
<i>E. coli</i> in WM	0.180	3.85	1.14E+05	4.35	99.99
<i>E. coli</i> and HA	0.150	4.62	2.54E+04	3.70	99.9
<i>E. coli</i> in HA and WM	0.117	5.92	6.21E+04	3.06	99.9

Table 4.20. SynTiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution showed the highest bacteria reduction percentage among other media, as 99.999 %. Similar *E. coli* LRV was achieved with TiO₂ solar photocatalytic treatment (Table 4.11).

Due to constituents present in WM solution used in SynTiO₂ solar photocatalytic treatment, *E. coli* displayed lower % reduction than *E. coli* in IsoT solution in correlation with the findings attained upon use of TiO₂ in IsoT and WM solutions (Table 4.11).

In the presence of HA, decreased *E. coli* cell counts resulted in reduction as 99.9 % upon use of SynTiO₂ photocatalyst specimen as being 2.5 times higher than LRV upon use of TiO₂ (Table 4.11). It could be indicated that while TiO₂ displayed selectivity for HA, SynTiO₂ was more selective towards *E. coli*.

Upon SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution, showed the same bacteria reduction percentage with HA, as 99.9 % however slightly lower LRV was attained in "HA and WM" solution. Although similar data were obtained with use of sole HA solution, these bacteria inactivation values were detected way higher than those with TiO₂ (Table 4.11). The diverse efficiencies of these TiO₂ specimens could confirm that SynTiO₂ had higher selectivity to *E. coli* rather than organic matrix. Moreover, Turkten and Cinar, having worked with the same SynTiO₂, achieved lower organic matter removals than TiO₂, in accordance with the assumption of SynTiO₂ selectivity against bacteria (Turkten and Cinar, 2017). 4.2.2.2. Organic matter formation and removal upon SynTiO₂ solar photocatalytic treatment of *E*. *coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter remaining in solution were presented in Figures 4.50 and 4.51. Irradiation time dependent absorbance changes at $\lambda < 280$ nm were significantly similar to each other in the absence of aromatic domains and color forming moieties. Similar UV spectral features were also attained from solar photolytic (Figure 4.2) and TiO₂ solar photocatalytic treatments of *E*. *coli* in IsoT solution (Figure 4.26).



Figure 4.50. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm. (Figure 4.51). The major peak was accompanied with a tailing effect in lower intensities in longer emission wavelengths as compared to the main peak intensity at λ_{emis} =280 nm as FI_{sync}≈50. Further evaluation would also consider minor peak intensity at λ_{emis} =470 nm to maintain a basis for comparison upon use of HA. From a general perspective, an irradiation time dependent decreasing order in FI_{sync280} data was observed. On the other hand, contrary to minor presence of FI_{sync470} expressing almost insignificant intensities in the range of 0.0216-0.532, predominant FI_{sync280} displayed changes in between 31.5-53.1 (Figure 4.52).

All UV-vis parameters displayed very low values (Figure 4.52). The highest UV₂₅₄ was attained upon $t_{irr}=60 \text{ min } (0.0341)$ as could also be considered as similar to $t_{irr}=20 \text{ min } (0.0332)$ although variations were insignificant. UV₂₈₀ varied in between 0.0164-0.0218 being the lowest for $t_{irr}=40 \text{ min}$ and the highest for initial samples. Due to non-existence and subsequent degradation of color forming moieties of the released organic matrix upon irradiation and the usage of IsoT solution, quite low

Color₄₃₆ and UV₃₆₅ values were attained. NPOC content showed a slight decrease between t=0 and t_{irr} =10 min (4.16-4.09 mg/L) and increased up to t_{irr} =40 min (5.65 mg/L) due to the release of organics from damaged and destructed *E. coli* cells, followed by a sharp decrease at t_{irr} =60 min (4.25 mg/L).



Figure 4.51. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon SynTiO₂ solar photocatalytic treatment.



Figure 4.52. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon SynTiO₂ solar photocatalytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated (Figure 4.53). SUVA₂₅₄ displayed variations (0.520-0.827 L/m mg) during SynTiO₂ solar photocatalytic treatment with respect to varying trends in both UV₂₅₄ and NPOC removals. Almost insignificant presence of color forming moieties also reflected in insignificant SCoA₄₃₆

profiles in accordance with slight variations in NPOC data determined under all conditions as 3.82-5.26 mg/L. Most important difference could be related to SFI_{syn280} as almost decreasing by half ratio. Besides SFI_{sync470} displayed quite insignificant presence in the range of 0.00519-0.0621, SFI_{sync280} as the main flurophoric group in the absence of HA, expressed variations in between 6.37-13.9.



Figure 4.53. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial conditions, upon introduction of SynTiO₂ (t=0) and upon SynTiO₂ solar photocatalytic inactivation of *E. coli* in IsoT solution exposed to irradiation for 60 min were presented in Table 4.21.

of E. coli in IsoT solution. UV-vis Spectroscopic Parameters, cm⁻¹ Irradiation NPOC. SUVA₂₅₄, FI FIsync280 SFI_{sync280} time, min mg/L L/m mg Color₄₃₆ UV365 UV₂₈₀ UV_{254} 0.0016 0.0036 0.0218 0.0316 1.39 Initial 3.82 0.827 53.1 13.9 t=049.0 0.0013 0.0025 0.0183 00267 4.16 0.642 11.8 1.63

4.25

0.802

31.5

7.41

1.90

0.0341

60 min

0.0024

0.0045

0.0199

 Table 4.21. Characteristic properties of organic matter present in SynTiO₂ solar photolytic treatment of *E. coli* in IsoT solution.

There was no significant difference between the slightly increasing order of UV-vis spectroscopic parameters recorded under all conditions. Upon irradiation period of 60 min at which 5.64 log reduction of *E. coli* in IsoT solution (Table 4.20) was attained, released organic matter expressed the very low UV-vis parameters, along with organic contents NPOC as 4.25 mg/L. SUVA₂₅₄ value at t_{irr} =60 min as 0.802 L/m mg represents that the remaining organic matter could be

considered having hydrophilic and aliphatic character (Edzwald et al., 1985). Owing to the degradation of protein content, $FI_{sync280}$ displayed a decreasing trend at the end of 60 min irradiation period. $SFI_{sync280}$ value showed slight decrease at t_{irr} =60 min in accordance with the increased NPOC due to the released organic material from bacteria. According to very low $FI_{sync470}$ and NPOC data, calculated $SFI_{sync470}$ were insignificant under all conditions. At t_{irr} =60 min, FI was calculated as 1.90, in accordance with the microbially derived organic content of the experiment solution.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in IsoT solution during SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.54.



Figure 4.54. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Throughout 60 min irradiation period, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of released organic matter and reduction of *E. coli*. However, at $t_{irr}=60$ min fluorescence intensities were detected as higher than the ones of TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution (Figure 4.30). Furthermore, decreasing intensities at Region I and II representing the presence of aromatic proteins could also be correlated with the irradiation time dependent decreasing order of FI_{sync280} data (Figures 4.51 and 4.52).

<u>4.2.2.3. Mechanistic evaluation of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in IsoT solution. As bacteria inactivation products total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples of SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution were presented in Table 4.22.</u>

SynTiO ₂ solar photocatalytic	E. coli in IsoT solution					
treatment	Initial	t=0	60 min			
Total K, mg/L	1.18	0.97	1.07			
Protein, mg/L	3.54	3.95	5.86			
Carbohydrate, mg/L	2.34	3.15	45.0			

Table 4.22. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation inIsoT solution upon SynTiO2 solar photocatalytic treatment.

Total K content displayed immediate adsorptive removal as 17.8 % at t=0, which could be resulting from K⁺ ion adsorption onto negatively charged *E. coli* and/or TiO₂ surfaces. At the end of 60 min irradiation period, total K content slightly increased due to released K⁺ ions from damaged bacteria cells.

E. coli in IsoT solution upon SynTiO₂ solar photocatalytic treatment displayed increasing trend in both protein and carbohydrate contents. Protein contents were plotted against $FI_{sync280}$ and NPOC in order to detect if there is any correlation between these parameters; while protein contents and $FI_{sync280}$ displayed a strong correlation as R²=1.00, protein concentrations and NPOC showed a moderate correlation as R²=0.60. Carbohydrate concentration showed a drastic increase after 60 min of irradiation period (45.0 mg/L) that could be due to an unexpected condition. Various researchers reported that the use of strong acid may cause production of some degradation products which can absorb light in the same spectral band with carbohydrates, resulting in overestimation of carbohydrate concentration (Josefsson et al. 1972; Dawson and Liebezeit, 1981). Moreover, Rao and Pattabiraman indicate that the presence of glycoproteins could be troublesome for detection of carbohydrate content (Rao and Pattabiraman, 1989). No significant correlation was detected between carbohydrate concentrations and NPOC data.

4.2.2.4. Organic matter formation and removal upon SynTiO₂ solar photocatalytic treatment of *E*. *coli* in WM solution. UV-vis absorption spectra of organic matter in reaction medium were presented in Figure 4.55. Irradiation time dependent absorbance changes displayed minor differences from each other in UV wavelength range of λ =200-280 nm in the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) that should be measured in longer wavelengths. Initial solution expressed higher absorption at $\lambda > 210$ nm however t_{irr}=60 min condition displayed lower absorption at $\lambda < 240$ nm indicating that light absorbing moieties could be emerged and removed throughout the irradiation period.

Different from *E. coli* in IsoT solution conditions, the use of WM solution resulted in the detection of different absorbance characteristics at $\lambda < 250$ nm. Especially between $\lambda=200-210$ nm a shoulder effect was detected for all samples originating from the strong absorption spectrum of NO₃⁻ having its λ_{max} at approximately 200 nm (Krishnan and Guha, 1934; Edwards at el., 2001). Similar absorption behaviors could also be visualized with previous solar photolytic (Figure 4.7) and TiO₂ photocatalytic treatments carried out with WM solution (Figure 4.31).



Figure 4.55. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm, although, a second peak at λ_{emis} =420 nm was observed for t=0 min condition (Figure 4.55). The emergence of the fluorophores at emission wavelength of 420 nm could be related to the conformational changes in the released organic matter upon use of SynTiO₂ expressing different surface properties in comparison to bare TiO₂ (Figure 4.32 and Appendix C). Furthermore, FI_{sync} data displayed a dramatic increase at t_{irr}=10 min displaying a broad spectrum followed which a reducing trend was attained during solar photocatalytic treatment. Fluorescence spectral shape resembled a mono peak at λ_{emis} =280 nm style as depicted previously in case of IsoT solution (Figure 4.51). Furthermore, very low FI_{sync470} showed changes in between 0.220-4.32 however dominant FI_{sync280}, in the absence of HA solution as an aqueous matrix component (Figure 4.57).



Figure 4.56. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

From a general perspective, NPOC expressed variations in the range of 1.81-11.7 mg/L (Figure 4.57). NPOC content significantly increased between initial (1.81 mg/L) and t=0 (7.17 mg/L) conditions which could be most probably due to mechanistic adverse effect of SynTiO₂ on *E. coli* cells, at t_{irr}=20 min (4.44 mg/L) NPOC declined and again increased until t_{irr}=30 min (11.7 mg/L) due to release of organics from damaged and destructed *E. coli* cells, and followed by a declining trend until t_{irr}=40 min (9.93 mg/L), further prolonged irradiation duration showed a slight increase up to 10.8 mg/L. The lowest NPOC content was determined under initial condition (1.81mg/L) representing the amount of organic matter present in WM solution following immediate filtration through 0.45 µm membrane filter. Therefore, approximately 10 mg/L NPOC was present in reaction

medium upon solar photocatalysis using SynTiO₂. On the other hand, UV_{254} varied in between 0.0302-0.1080 was detected as the most significant UV-vis parameter.



Figure 4.57. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.58. After the introduction of SynTiO₂ into the system, all specific UV-vis parameters expressed a drastic decrease. As the most significant parameter, SUVA₂₅₄ displayed changes (0.280-5.97 L/m mg) during solar photocatalytic treatment in relation to concurrent differences in both UV₂₅₄ and NPOC removals. At t_{irr}=60 min, SUVA₂₅₄ displayed a significant decrease as 95.3 %.



Figure 4.58. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

Due to non-existence and subsequent degradation of color forming moieties of the released organic matter upon irradiation, calculated SCoA₄₃₆ and SUVA₃₆₅ were quite low. However, SUVA₂₈₀ as the secondly significant parameter, expressed variations in between 0.213-5.34 L/m mg being the lowest for t_{irr} =60 min and the highest for initial conditions. Furthermore, specific fluorescence parameter as SFI_{sync470} and SFI_{sync280} expressed variations in the range of 0.0221-2.68 and 2.21-11.9, respectively. The main fluorophoric group was accepted as SFI_{sync280} and the most remarkable variation was noticed for SFI_{sync280} being highest under 20 min of irradiation (11.9).

Characteristic properties of organic matter present under initial and t=0 conditions and following irradiation for 60 min SynTiO₂ solar photocatalytic inactivation of *E. coli* in WM solution were presented in Table 4.23.

 Table 4.23. Characteristic properties of organic matter present in SynTiO₂ solar photolytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC,	SUVA ₂₅₄ ,	DI	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	F1 _{sync280}	SF1 _{sync280}	ГІ
Initial	0.0319	0.0452	0.0966	0.108	1.81	5.97	17.9	9.90	1.48
t=0	0.0127	0.0215	0.0480	0.0580	7.17	0.809	15.8	2.21	1.33
60 min	0.0034	0.0076	0.0230	0.0302	10.8	0.280	45.6	4.22	1.74

Upon irradiation period of 60 min SynTiO₂ solar photocatalytic treatment at which 4.35 log reduction of *E. coli* in WM was attained (Table 4.20), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0034, UV₃₆₅: 0.0076, UV₂₈₀: 0.0230, UV₂₅₄: 0.0302, and NPOC as 10.8 mg/L. SUVA₂₅₄ value was calculated as 0.280 L/m mg upon t_{irr}=60 min represents that the remaining organic matter could be considered having hydrophilic and aliphatic character (Edzwald et al., 1985).

 $FI_{sync280}$ increased at the end of $t_{irr}=60$ min, which could be related to the release of *E. coli* protein content. $SFI_{sync280}$ value displayed a decrease at $t_{irr}=60$ min in accordance with increased NPOC due to released organic material from bacteria. On the other hand, $FI_{sync470}$ displayed almost insignificant presence as 4.33, 2.16, and 0.908 for initial, t=0, and $t_{irr}=60$ min conditions, respectively. Owing to the degradation of released organic content from damaged *E. coli* cells, $FI_{sync470}$ expressed a decrease at $t_{irr}=60$ min. $SFI_{sync470}$ values of initial, t=0, and $t_{irr}=60$ min samples were calculated as 2.39, 0.301, and 0.0841, respectively. Furthermore, FI at $t_{irr}=60$ min was calculated as 1.74 which could mostly be related to the presence of microbially derived organic matter in solution (Sen-Kavurmaci and Bekbolet, 2014). Despite calculated FI was between the indication values of humic and microbially derived organics, it could most probably stemming from microbial origin. Moreover, the presence of microbially derived organic content in t_{irr} =60 min solution could also be supported with EEM contour plots represented in following Figure 4.59.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM during SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.59. EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions.



Figure 4.59. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

Following further irradiation periods, released microbial by-products and protein content from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli* which could be visualized by reduced fluorescence intensities of related regions. Furthermore, similar to previous solar photolytic (Figure 4.11) and TiO₂ solar photocatalytic treatment carried out in WM solution (Figure 4.35) lower regional fluorescence intensities were detected than IsoT solution conditions (Figures 4.6, 4.30, and 4.54). On the other hand, fluorescence intensities were detected as higher than TiO₂ solar photocatalytic treatment of *E. coli* in WM solution conditions (Figure 4.35).

<u>4.2.2.5. Mechanistic evaluation of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in WM solution. Upon bacteria reduction, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples of SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution were presented in Table 4.24.</u>

		J				
SynTiO ₂ solar photocatalytic	E. coli in WM					
treatment	Initial	t=0	60 min			
Total K, mg/L	4.38	4.26	4.12			
Protein, mg/L	2.35	2.85	14.8			
Carbohydrate, mg/L	2.28	1.89	1.83			

Table 4.24. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon SynTiO₂ solar photocatalytic treatment.

Total K content displayed a decreasing trend differing from *E. coli* behavior in WM solution under solar photolytic (Table 4.6) and photocatalytic conditions (Table 4.15). Immediate adsorptive removal of total K as 2.7 % was calculated at t=0 condition. Furthermore, decreased total K content after 60 min of irradiation period could be related to adsorption of released and/or present K⁺ ions in WM solution onto both negatively charged *E. coli* and SynTiO₂ surface sites.

Upon SynTiO₂ solar photocatalysis, *E. coli* in WM solution displayed different trends for protein and carbohydrate evolution upon cell destruction. While protein content showed a significant increase (2.35-14.8 mg/L), carbohydrate content slightly decreased at the end of 60 min irradiation duration (2.28-1.83 mg/L). A similar inverse relation was also detected by fluorescence intensities as expressed by $FI_{sync280}$ and $FI_{sync470}$ (Figure 4.57). Both $FI_{sync280}$ and protein contents increased with a strong correlation as R²=0.99. Protein contents and NPOC displayed a moderate correlation expressed as R²=0.68 however carbohydrate and NPOC contents showed a strong correlation as R²=0.91. 4.2.2.6. Organic matter formation and removal upon SynTiO₂ solar photocatalytic treatment of *E*. *coli* in HA solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in HA solution were presented in Figures 4.60 and 4.61. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of dense aromatic domains as well as color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀). Despite the visualization of irradiation time dependent sequential degradation of organic matter, differences were more pronounced in the previous set of TiO₂ solar photocatalytic treatment (Figure 4.36).



Figure 4.60. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon SynTiO₂ solar photocatalytic treatment.



Figure 4.61. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =370 nm and two minor peaks at λ_{emis} =440 nm and 480 nm representing the presence of aromatic constituents in solution matrix (Dujmov and Sučević, 1990) (Figure 4.61). A relevant decrease in synchronous fluorescence intensities were observed following 40 min of irradiation. On the other hand, FI_{sync470} and FI_{sync280} displayed changes in the range of 68.4-99.7 and 8.03-44.0, respectively irrespective of the exposure periods. The predominant fluorophoric group referred to FI_{sync470} in the presence of HA as the solution matrix was evident under all conditions along with FI_{sync370}. FI_{sync280} of sole HA_i would not be expected, however FI_{sync280} of the initial solution increased to 8.03 indicating the presence of protein content in the *E. coli* cell suspension (Figure 4.62).

From a general view, NPOC content expressed variations in the range of 4.74-23.1 mg/L. A drastic increase in NPOC content was attained upon $t_{irr}=10 \text{ min } (4.74-23.1 \text{ mg/L})$ due to the release of organics from damaged and/or destructed *E. coli* cells followed by a sharp decrease at $t_{irr}=20 \text{ min}$ (15.1 mg/L) and again displayed an increasing trend as observed during SynTiO₂ solar photocatalytic treatment (19.9 mg/L) (Figure 4.62).



Figure 4.62. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon SynTiO₂ solar photocatalytic treatment.

Along with the use of HA solution, all UV-vis parameters expressed high presence. After t=0 condition, all specified UV-vis parameters followed a decreasing trend however obtained decrease was rather low in comparison to TiO₂ solar photocatalytic treatment of *E. coli* in HA solution (Figure 4.38). UV₂₅₄, which was the most significant parameter displayed variations in the range of 0.4201-0.5283, being the highest for t=0 condition and the lowest for t_{irr}=60 min condition. Moreover,

especially Color_{436} expressed very minor decrease at the end of 60 min light exposure in comparison to almost total removal obtained with the use of TiO₂ as a bare photocatalyst specimen. According to minor removal of NPOC and UV-vis parameters, it could be concluded that low organic content degradation was attained throughout 60 min of irradiation duration. Obtained data could support the interpretation was made about SynTiO₂ having higher selectivity against *E. coli* rather than HA in Section 4.2.2.1.

Based on NPOC data attained under all conditions, specific UV-vis parameters as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.63. Most importantly, SUVA₂₅₄ displayed variations in between 2.11-9.76 L/m mg during SynTiO₂ solar photocatalytic treatment indicating concomitant differences in both UV₂₅₄ and NPOC removals. The highest SUVA₂₅₄ was attained for sole HA solution.

Furthermore, specific fluorescence parameters as predominant $SFI_{sync470}$ and quite low $SFI_{sync280}$ expressed variations in the range of 3.44-15.6 and 0.929-2.47, respectively. Obtained $SFI_{sync280}$ values were almost insignificant. According to insignificant presence of $FI_{sync280}$ related to HA_i , $SFI_{sync280}$ would be irrelevant in comparison to $SFI_{sync470}$.



Figure 4.63. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon SynTiO₂ solar photocatalytic treatment.

For comparative evaluation purposes, characteristic properties of organic matter present in solutions under initial condition, upon introduction of SynTiO₂ (t=0) and SynTiO₂ solar

photocatalytic inactivation of *E. coli* in HA solution exposed to irradiation for 60 min were presented in Table 4.25.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	FI	SEL	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	ГІ
Initial	0.0756	0.1629	0.4334	0.5150	8.65	5.95	75.0	8.67	1.08
t=0	0.0875	0.1836	0.4511	0.5283	14.6	3.62	114	7.80	1.10
60 min	0.0608	0.1358	0.3466	0.4201	19.9	2.11	68.4	3.44	1.16

Table 4.25. Characteristic properties of organic matter present in SynTiO₂ solar photolytic treatment of *E. coli* in HA solution.

Upon t_{irr}=60 min of SynTiO₂ solar photocatalytic treatment of *E coli* in HA solution, at which 3.70 log *E. coli* reduction (Table 4.20) was achieved, released organic matter expressed the following UV-vis parameters; Color₄₃₆: 0.0608, UV₃₆₅: 0.1358, UV₂₈₀: 0.3466, UV₂₅₄: 0.4201, and NPOC as 19.9 mg/L.

SUVA₂₅₄ of initial and following t_{irr} =60 min conditions of irradiation were calculated as 5.95 and 2.11 L/m mg, respectively. Detected decrease in SUVA₂₅₄ indicated the degradation of aromatic organics during 60 min of solar photocatalysis. The use of HA solution as reaction matrix caused relatively high initial NPOC and SUVA₂₅₄ as expected.

 $FI_{sync470}$ displayed a significant increase at t=0 condition (75.0-114) and after 60 min of irradiation decreased below initial value (68.4) emphasizing degradation of released cytoplasmic organic matter from *E. coli*. According to changes in FI_{sync470} and NPOC content, calculated SFI_{sync470} displayed a decrease after t_{irr}=60 min. On the other hand, FI_{sync280} measurements of initial, t=0 and t_{irr}=60 min conditions were detected as 8.03, 14.6 and 44.1, respectively. FI_{sync280} demonstrated a significant increase at the end of 60 min solar light exposure due to the released protein content from *E. coli* cells. SFI_{sync280} attained under initial, t=0, and t_{irr}=60 min conditions were calculated as 0.929, 2.13, and 2.21, respectively. After irradiation period of 60 min, FI was calculated as 1.16 indicating the presence of fluorophores emerging from humic matter present in experimental solution matrix as expected.

In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in HA solution during SynTiO₂ solar photocatalytic treatment were also followed and displayed in Figure 4.64.



Figure 4.64. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in HA solution upon SynTiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities expressed a slight decrease throughout the irradiation periods revealing that organic matter having HA origin showed only minor degradation during t_{irr} =60 min. Besides EEM fluorescence contour plots showing almost no change of humic-like and fulvic-like fluorophores, high *E. coli* LRV as 3.70 was attained upon 60 min of irradiation period which also supported the selectivity of SynTiO₂ towards *E. coli*. It could be concluded that specified UV-vis parameters, NPOC data, and EEM fluorescence contour plots were complied with each other about degradation of organic content in solution. Moreover, EEM fluorescence contour plots displaying low organic matrix removal were

similar to the findings of Turkten and Cınar (Turkten and Cinar, 2017). Although released and degraded microbial by-products and protein-like fluorophores were present in solution, they could not exactly be identified due to possible masking effect of humic-like components (Wang et al., 2015).

<u>4.2.2.7. Mechanistic evaluation of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in HA solution. Upon cell destruction of *E. coli* in HA solution via SynTiO₂ solar photocatalysis, released total K, protein, and carbohydrate contents under initial, t=0, and t_{irr}=60 min conditions were presented in Table 4.26.</u>

E. coli and HA SynTiO₂ solar photocatalytic treatment Initial t=060 min Total K, mg/L 0.61 0.73 0.64 Protein, mg/L 8.82 7.93 11.1 Carbohydrate, mg/L 2.51 2.61 25.3

Table 4.26. Total K, protein, and carbohydrate contents of time dependent E. coli inactivation inHA solution upon SynTiO2 solar photocatalytic treatment.

Total K content of *E. coli* in HA solution by solar photocatalytic treatment with SynTiO₂ displayed a slight increase between initial and t_{irr} =60 min conditions expressing K⁺ ion release from damaged bacteria cells. Measured low total K concentrations showed similarity with both solar photolytic (Table 4.8) and TiO₂ solar photocatalytic treatment sets of *E. coli* in the presence of HA solution (Table 4.17).

Upon SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution, increase in both protein and carbohydrate contents were observed at the end of 60 min solar light exposure. Carbohydrate concentration showed significant increase (2.51-25.3 mg/L) while protein concentration displayed a slight increase (8.82-11.1 mg/L). A plausible explanation of this remarkable increase of carbohydrate concentration could be emerging from nondegraded humic sub-fractions along with released carbohydrate from *E. coli* cells upon solar photocatalysis. Moreover, high carbohydrate concentrations could be related with EEM counter plots having high intensities for fulvic-like and humic-like regions as Region III and V (Figure 4.64). FI_{sync280} and NPOC data were plotted against protein concentrations to detect if there was any correlation, and neither FI_{sync280} nor NPOC showed any correlation with protein contents. On the other hand, NPOC and carbohydrate concentrations displayed a moderate correlation with each other as R²=0.72. 4.2.2.8. Organic matter formation and removal upon SynTiO₂ solar photocatalytic treatment of *E*. *coli* in "HA and WM" solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter remaining in "HA and WM" solution including *E. coli* cell inactivation products were presented in Figures 4.65 and 4.66, respectively. Irradiation time dependent absorbance changes were different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) mainly originating from humic sub-fractions as also recorded sole HA spectral features. Irradiation time dependent sequential degradation of organic matter could be visualized throughout 60 min of irradiation period. In contradiction to previous set of TiO₂ solar photocatalytic treatment (Figure 4.36), no evident difference was observed between initial and t=0 conditions revealing insignificant surface interactions. On the other hand, absorbances detected at λ < 240 nm were characteristic to NO₃⁻ containing "HA and WM" solution.



Figure 4.65. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =380 nm as was also observed in sole HA_i solution (Figures 4.61 and 4.66). Besides the intense peaks at λ_{emis} =380 nm, two minor peaks at λ_{emis} =440 nm and 480 nm were detected as well. Similar peak wavelengths presenting the presence of aromatic hydrocarbons were also detected for *E. coli* in HA solution upon SynTiO₂ solar photocatalytic treatment (Figure 4.61). Similar to solar photolytic treatment of *E. coli* in "HA and WM" solution (Figure 4.18), "HA and WM" solution and initial condition, showed increased fluorescence intensities relative to sole HA_i solution at λ_{emis} < 450 nm. After the addition of photocatalyst into solution matrix, a significant increase in FI_{sync} data was observed relative to sole HA_i, "HA and WM", and initial conditions. It could be indicated that the

mechanistic effect of SynTiO₂ on bacteria cell wall could cause damaged cell wall structure and release of cytoplasmic organics resulting in increased FI_{sync} data. Similar findings could also be visualized with the use of TiO₂ (Figure 4.42). From a general perspective, FI_{sync} data did not follow an irradiation time dependent decreasing order. $FI_{sync380}$ data showed a decreasing trend between t_{irr} =10 min and 30 min conditions and followed by an increasing trend at 60 min of solar light exposure revealing that the re-emergence of removed fluorophoric groups following further irradiation periods.

Furthermore, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 44.2-94.9 and 6.73-48.0, respectively (Figure 4.67). $FI_{sync280}$ of HA_i could not be detected however $FI_{sync280}$ in "HA and WM" solution was detected as 6.73. Due to the use of HA solution, the dominant fluorophoric group was accepted as $FI_{sync470}$.



Figure 4.66. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

Along with the presence of "HA and WM" solution as reaction matrix, NPOC expressed variations in the range of 4.87-18.3 mg/L (Figure 4.67). NPOC content showed an insignificant decrease between HA_i (4.87 mg/L) and "HA and WM" solution (4.77 mg/L) and increased up to $t_{irr}=20 \text{ min}$ (18.3 mg/L) due to release of organics from damaged and destructed *E. coli* cells, afterwards remained stable up to $t_{irr}=30 \text{ min}$ and followed by a decreasing trend until the end of 60 min solar light exposure duration (10.1 mg/L). According to almost remained stable *E. coli* enumerations at $t_{irr}=40 \text{ min}$ (6.75E+02 CFU/mL) and 60 min (4.67E+02 CFU/mL) (Figure 4.49), it

could be indicated that decreased NPOC content after $t_{irr}=30$ min could be related with degradation of both HA content and released cytoplasmic organic content from destructed *E. coli* cells.

Specified UV-vis parameters expressed a high presence due to the use of HA solution as a reaction matrix component. UV₂₅₄, as the most evident parameter, displayed changes in between 0.4626-0.3149. Excluding slightly increased UV₂₅₄ at irradiation period of 40 min, UV₂₅₄ showed a decreasing trend throughout the treatment, however obtained decrease could be considered as insignificant in comparison with TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution (Figure 4.43). Similar lessened decrease in specified UV-vis parameters was also recorded upon solar photocatalytic *E. coli* inactivation in sole HA solution the presence of SynTiO₂ in comparison to use of TiO₂ as a bare photocatalyst specimen (Figures 4.38 and 4.62).



Figure 4.67. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

Based on NPOC contents attained under all solar irradiation periods, specific UV-vis parameters as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.68. as the most significant specific UV-vis parameter, SUVA₂₅₄ expressed changes in the range of 1.76-9.50 L/m mg during SynTiO₂ solar photocatalytic treatment indicating concomitant differences in both UV₂₅₄ and NPOC removals. SUVA₂₅₄ showed a decreasing trend up to t_{irr} =30 min and increased at the end of t_{irr} =60 min. On the other hand, calculated SFI_{sync470} and SFI_{sync280} expressed variations in the range of 2.78-15.7 and 1.41-4.75, respectively. Due to the presence of HA

solution, $SFI_{sync470}$ displayed evident presence and accepted as the predominant fluorophoric group in compared to $SFI_{sync280}$.



Figure 4.68. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter present under initial and t=0 conditions, and t_{irr} =60 min of solar light exposure attained upon SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution were presented in Table 4.27.

Table 4.27. Characteristic properties of organic matter present in SynTiO₂ solar photolytic treatment of *E. coli* in "HA and WM" solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	DI	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	ГІ
Initial	0.0548	0.1242	0.3433	0.4155	7.61	5.46	78.3	10.3	1.17
t=0	0.0511	0.1176	0.3185	0.3821	9.21	4.15	82.3	8.94	1.19
60 min	0.0312	0.0784	0.2458	0.3149	10.1	3.12	49.0	4.85	1.26

Upon $t_{inr}=60 \text{ min of SynTiO}_2$ solar photocatalytic treatment of *E coli* in "HA and WM" solution, 3.06 log *E. coli* reduction was achieved (Table 4.20), and released organic matter expressed the following UV-vis parameters; Color₄₃₆: 0.0312, UV₃₆₅: 0.0784, UV₂₈₀: 0.2458, UV₂₅₄: 0.3149, and NPOC as 10.1 mg/L. According to calculated SUVA₂₅₄, it could be indicated that initial and t=0 solutions displaying aromatic and hydrophobic features however after $t_{irr}=60$ min, organic content degraded and expressed mostly aliphatic and hydrophilic characteristics (Edzwald et al., 1985).

 $FI_{sync470}$ and $SFI_{sync470}$ displayed a decrease at t_{irr} =60 min. However, $FI_{sync280}$ of initial, t=0, and t_{irr} =60 min conditions were detected as 24.8, 29.3, and 48.0, respectively and $SFI_{sync280}$ were calculated as 3.26, 3.18, and 4.75, respectively, showing an increased trend. FI of t_{irr} =60 min condition was calculated as 1.26 indicating the presence of fluorophores originating from HA content as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in "HA and WM" solution during SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.69.



Figure 4.69. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities displayed a slight decrease throughout irradiation period of 60 min however they displayed more intense fluorescence in comparison to *E. coli* inactivation in HA solution (Figure 4.64). Similar finfings were also obtained from solar photolytic treatment of *E. coli* both in sole HA solution (Figure 4.16) and "HA and WM" solution (Figure 4.21). It could be indicated that the presence of WM in the solution matrix caused inhibition of HA degradation. Different from the TiO₂ solar photocatalytic treatment (Figure 4.45), SynTiO₂ expressed lower degradation of fulvic-like and humic-like substances. Moreover, Regions I, II and IV were quenched by fulvic-like and humic-like substances originating from HA as expected.

<u>4.2.2.9. Mechanistic evaluation of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in "HA and WM" solution. Upon irradiation, as bacteria inactivation products of SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.28.</u>

SynTiO ₂ solar photocatalytic	<i>E. coli</i> in "HA and WM"					
treatment	Initial	t=0	60 min			
Total K, mg/L	4.48	4.38	4.47			
Protein, mg/L	6.65	5.95	13.3			
Carbohydrate, mg/L	2.02	2.05	28.4			

Table 4.28. Total K, protein, and carbohydrate content of time dependent *E. coli* inactivation in "HA and WM" solution upon SynTiO₂ solar photocatalytic treatment.

Total K content remained nearly constant under all conditions. The reason could be attributed to various attractive and repulsive electrostatic interactions as well as passive transport mechanism of *E. coli* cells regulating K^+ ion uptake from and/or release to reaction medium containing WM solution.

Upon solar photocatalytic treatment, substantial increase in both protein (6.65-13.3 mg/L) and carbohydrate (2.02-28.4 mg/L) concentrations were detected. Carbohydrate concentration showed higher increase than protein concentration similar to *E. coli* inactivation conducted in sole HA solution (Table 4.26). Protein contents and $FI_{sync280}$ displayed a strong correlation as R²=0.93. Both protein and carbohydrate contents were plotted against NPOC separately and showed moderate correlations with each other as R²=0.51 and R²=0.60. Moreover, it could be concluded that increased
NPOC (Table 4.27) and intense EEM contour plots (Figure 4.69) were roughly correlated with detected high carbohydrate content.

4.2.2.10. Comparative evaluation of different experimental matrixes on SynTiO₂ solar photocatalytic inactivation mechanism of *E. coli*. Upon solar inactivation process via SynTiO₂ photocatalysis, total K, protein, and carbohydrate contents and bacteria enumerations attained in comparison to various conditions were given in Figure 4.70. Total K contents varied in between 0.61-4.48 mg/L being the lowest for the initial condition of *E. coli* in sole HA solution and the highest for the initial sample of *E. coli* in "HA and WM" solution. Total K content attained in WM containing treatments showed increased presence due to high total K concentration of sole WM solution (3.84 mg/L, Section 4.1.5). Considering total K concentrations of different matrix conditions individually, no significant difference was detected between initial, t=0 and t_{irr}=60 min conditions. Similar to previous treatments carried out in HA solution (Table 4.8 and 4.17), lower total K contents were obtained among other solution matrixes upon use of SynTiO₂ specimen. Except the presence of sole WM solution, total K and *E. coli* counts displayed a rough correlation as R² > 0.71.



Figure 4.70. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in various aqueous matrixes.

Covering all conditions, protein contents varied in between 2.35-14.8 mg/L being the lowest for initial and the highest for $t_{irr}=60$ min conditions of *E. coli* in WM solution. Similarly, carbohydrate contents varied in between 1.83-45.0 mg/L being the lowest upon treatment for $t_{irr}=60$ min of *E. coli* in WM solution whereas the highest for $t_{irr}=60$ min sample of *E. coli* in IsoT solution. Protein contents

under initial and t=0 conditions in HA solution containing reaction matrixes were higher than the case attained in IsoT and WM solutions. *E. coli* in IsoT solution for t_{irr} =60 min displayed a significant increase in carbohydrate content which could be resulting from an experimental error as explained in Section 4.2.2.3. Except inactivation profiles attained in WM solution, under all conditions an increasing trend for protein and carbohydrate contents were observed. *E. coli* inactivation upon t_{irr} =60 min showed a decrease in "HA and WM" solution (4.67E+02 CFU/mL) however the presence of sole HA did not cause a significant retardation effect on *E. coli* inactivation (3.35E+01 CFU/mL).

Released protein and carbohydrate contents, calculating by subtracting initial contents from the final contents attained upon $t_{irr}=60$ min, and bacteria LRV of SynTiO₂ solar photocatalytic treatment of *E. coli* in various solution matrixes were given in Figure 4.71. Calculation for WM solution revealed negative values indicating degradation of carbohydrate content. Released protein content was higher for WM solution (12.5 mg/L). It could be concluded that the presence of WM solution displayed selectivity towards degradation of carbohydrates rather than protein. Released carbohydrate content (42.7 mg/L) and LRV of *E. coli* (5.64) in IsoT solution were detected as the highest ones. *E. coli* LRV was inversely proportional to protein release for IsoT and WM solutions. Moreover, in the presence of HA and "HA and WM" solutions, *E. coli* LRV were plotted against released protein concentration and no correlation was detected, whereas *E. coli* LRV and released carbohydrate contents displayed a strong correlation as R²=0.84 excluding the presence of sole WM solution.



Figure 4.71. Released protein and carbohydrate content, and bacteria LRV of *E. coli* inactivation upon SynTiO₂ solar photocatalytic treatment in various aqueous matrixes.

Upon SynTiO₂ solar photocatalytic treatment of *E. coli* in various matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV₂₈₀, and NPOC data under initial, t=0, and t_{irr}=60 min

conditions were given in Figure 4.72. Similar UV₂₈₀ values were obtained for initial, t=0, and t_{irr}=60 min samples of *E. coli* in IsoT solution. Both protein concentrations and UV₂₈₀ showed an increasing trend for HA solution containing experimental conditions. All conditions, except HA initial solution, expressed higher FI_{sync280} than protein contents. Moreover, a rough correlation as $R^2 > 0.67$ was detected between protein contents and FI_{sync280} in the absence of WM solution.



Figure 4.72. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and bacteria enumeration results of SynTiO₂ solar photocatalytic treatment of *E. coli* in various aqueous matrixes.

Carbohydrate concentrations were higher than NPOC contents in t_{irr} =60 min samples of IsoT, HA and "HA and WM" solutions. NPOC measurements displayed an increasing trend under all experimental conditions. No correlation was detected between carbohydrate concentrations and NPOC in initial samples; however, t=0 and t_{irr} =60 min data displayed a rough correlation as R² > 0.72 except the presence of sole HA solution.

4.2.3. 0.25% Fe-TiO₂ Solar Photocatalytic Inactivation of E. coli

4.2.3.1. 0.25% Fe-TiO₂ solar photocatalytic *E. coli* inactivation under specified reaction conditions, reductions, and kinetics. E. coli suspension (No=1.18E+06 CFU/mL) was prepared in IsoT solution and subjected to solar photocatalytic treatment using 0.25% Fe-TiO₂ specimen. Upon irradiation for 30 min, E. coli cell count decreased to 3.45E+02 CFU/mL. Following prolonged light exposure of t_{irr}=60 min, E. coli cell count further decreased to 6.13E+01 CFU/mL. Under these conditions LRV was 4.28 and reduction was 99.99 %. Upon preparation of E. coli (N_o=9.17E+05 CFU/mL) suspension in WM solution, for an irradiation period of 30 min, E. coli count decreased to 2.16E+04 CFU/mL. Following light exposure of tirr=60 min, E. coli count reached to 2.95E+03 CFU/mL revealing LRV as 2.49 and reduction as 99 %. In case that E. coli (N₀=6.13E+05 CFU/mL) suspension was prepared in HA solution, for an irradiation period of 30 min, E. coli count decreased to 9.79E+03 CFU/mL. Upon light exposure of tirr=60 min, E. coli count diminished to 5.00E+03 CFU/mL expressing LRV as 2.09 and reduction as 99 %. E. coli (N₀=9.26E+05 CFU/mL) suspension was also prepared in a solution of "HA and WM" components and was subjected to an irradiation period of 30 min in which E. coli count decreased to 1.17E+05 CFU/mL. Following light exposure of t_{irr}=60 min, E. coli count declined to 6.73E+03 CFU/mL revealing LRV as 2.14 and reduction as 99 %. Based on these results, the effect of solution matrix could be displayed in terms of LRV of E. coli in an increasing order as: HA < "HA and WM" < WM < IsoT. E. coli inactivation profiles and first order kinetic parameters were presented in Figure 4.73 and Table 4.29.



Figure 4.73. Irradiation time dependent *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in various solution matrixes.

0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution resulted in a rather fast reduction with k=0.174 min⁻¹ and half-life as 3.98 min. In the presence of HA, a reaction rate constant of k=0.0774 min⁻¹ and half-life as 8.95 min were attained. The effect of reaction medium could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; IsoT > WM > "HA and WM" > HA. Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) also followed the same trend as; 2.05E+05 > 8.16E+04 > 7.71E+04 > 4.74E+04 for IsoT, WM, "HA and WM", and HA, respectively.

Motriy	First	IDV	Daduction 0/			
Maurx	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min		Keuuciioii, %	
<i>E. coli</i> in IsoT	0.174	3.98	2.05E+05	4.28	99.99	
<i>E. coli</i> in WM	0.0890	7.79	8.16E+04	2.49	99	
<i>E. coli</i> and HA	0.0774	8.95	4.74E+04	2.09	99	
<i>E. coli</i> in "HA and WM"	0.0833	8.32	7.71E+04	2.14	99	

Table 4.29. 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

Solar photocatalytic treatment of *E. coli* in IsoT solution with 0.25% Fe-TiO₂ showed the highest bacteria reduction percentage among other experiment media, as 99.99 %. However, its LRV, which was detected as 4.28, was lower than the results of TiO₂ (Table 4.11) and SynTiO₂ (Table 4.20). Birben and colleagues studied with several doped TiO₂ types as; N-doped, Se-doped and Se-N co-doped under similar experimental conditions, except the use of deionized water as working solution, and they achieved rate constants (k, min⁻¹) as 0.153, 0.155 and 0.317 for N-doped, Se-N co-doped and Se-doped, respectively (Birben et al., 2017a). It could be indicated that 0.25% Fe-TiO₂ expressed a slightly higher reduction rate constant than N-doped and Se-N co-doped TiO₂ species. On the other hand, rate constant of Se-doped TiO₂ was higher than 0.25% Fe-TiO₂.

0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM solution showed lower % reduction (99 %) than *E. coli* in IsoT solution (99.99 %) due to the inhibition effect of WM components (Section 2.5). LRV for *E. coli* inactivation in WM solution was the lowest in comparison to either solar photolytic (Table 4.2) or TiO₂/SynTiO₂ photocatalytic treatments carried out so far (Tables 4.11 and 4.20).

The role of HA presence in *E. coli* inactivation using 0.25% Fe-TiO₂ solar photocatalysis could be explained as resulting in the lowest *E. coli* LRV as 2.09. It could be indicated that 0.25% Fe-TiO₂ had selectivity towards *E. coli* rather than organic matter like SynTiO₂ due to detected higher *E. coli* reduction than sole TiO₂. Birben and colleagues, having worked with the same 0.25% Fe-TiO₂, indicated that doped TiO₂ showed a retardation effect on HA removal rates compared to undoped TiO₂ (Birben et al., 2017b). This information could support *E. coli* selectivity suggestion about 0.25% Fe-TiO₂. Moreover, in another study of Birben and colleagues, in which under similar experimental conditions other doped TiO₂ species were employed, reaction rate constants were 0.163, 0.110 and 0.096 min⁻¹ for N-doped, Se-doped and Se-N co-doped, respectively (Birben et al., 2017a). Bacteria reduction rate constant of 0.25% Fe-TiO₂ as 0.0774 min⁻¹ was considerably lower in comparison to other doped TiO₂ species. It could be concluded that in the presence of HA, N-doped TiO₂ has the fastest bacteria reduction among other doped TiO₂ species.

E. coli in "HA and WM" solution solar photocatalytic treatment with 0.25% Fe-TiO₂ showed higher bacteria LRV than HA, as 2.14. It could be indicated that, the addition WM solution displayed no significant retardation effect as different than the presence of only HA solution. In the view of these very close *E. coli* LRV of WM, HA and "HA and WM" solutions, it could be concluded that the presence of HA more negatively affects *E. coli* inactivation than WM solution. Irrespective of the minor differences in LRV of these three solution matrixes, similar bacteria reduction percentages were detected as 99 %.

4.2.3.2. Organic matter formation and removal upon 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.74 and 4.75, respectively. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-250 nm displaying the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) that should be recorded in longer wavelengths as $\lambda \ge 280$ nm and extending to visible region. Detected absorbances at $\lambda < 210$ nm were characteristic to present IsoT solution (Tong et al., 2020). UV spectral features displayed similarities to the ones attained upon solar photolytic (Figure 4.2) and photocatalytic treatments with the use of bare TiO₂ specimens as TiO₂ (Figure 4.26) and SynTiO₂ (Figure 4.50).

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =280 nm (Figure 4.75). Moreover, the presence of fluorophoric groups expressing very minor intensities at λ_{emis} ≈470 nm could also be encountered similar to findings attained with the solar photolytic treatment (Figure 4.3) and use of TiO₂ (Figure 4.27) and SynTiO₂ (Figure 4.51) specimens in solar photocatalytic treatments. An irradiation time dependent decreasing order in FI_{sync280} was observed throughout the experiment. On the other hand, contrary to very low intensities of FI_{sync470} in the range of 0.0948-0.752, dominant FI_{sync280} displayed changes in between 24.1-51.0. In the absence of HA as solution matrix, the key fluorophoric group was recognized as FI_{sync280}.



Figure 4.74. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.



Figure 4.75. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

From a general perspective, NPOC data expressed variations in the range of 0.847-4.07 mg/L (Figure 4.76). NPOC content increased up to $t_{irr}=20 \text{ min } (0.847-4.01 \text{ mg/L})$ due to the release of organics from the damaged and destructed *E. coli* cells, followed by a minor fluctuation trend up to $t_{irr}=40 \text{ min } (3.69-4.07 \text{ mg/L})$ and then decreased to 2.91 mg/L during solar photocatalytic treatment. As the most significant specified UV-vis parameter, UV₂₅₄ displayed variations in the range of 0.0337-0.0422. The highest UV₂₅₄ was attained for initial condition (0.0422) as could also be considered as similar to t=0 (0.0418) and $t_{irr}=10 \text{ min } (0.0417)$ conditions although variations were insignificant.



Figure 4.76. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Based on NPOC contents, all UV-vis and fluorescence specific parameters as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated (Figure 4.77). Most importantly, SUVA₂₅₄ displayed variations (0.828-4.982 L/m mg) during 0.25% Fe-TiO₂ solar photocatalytic treatment indicating concomitant differences in both UV₂₅₄ and NPOC removals.



Figure 4.77. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Although SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀ were also presented, respective specified parameters were very low in comparison to total NPOC contents of the organic matter. Similar findings were

In a similar manner, SFI_{sync470} expressed variations in the range of 0.0326-0.829 expressing insignificant contribution of fluorophores detected at λ_{emis} =470 nm. As indicated above, SFI_{sync280} expressed considerable variations in the range of 7.86-60.3. The main fluorophoric group was accepted as SFI_{sync280} similar to previous solar photocatalytic treatments conducted with the use of TiO₂ (Figure 4.29) and SynTiO₂ (Figure 4.53) in IsoT solution.

For simplicity purposes, based on the irradiation time dependent variations in UV-vis and fluorescence spectroscopic properties along with NPOC data, characteristic properties of organic matter under initial condition, upon introduction of 0.25% Fe-TiO₂ (t=0) and at the end of 0.25% Fe-TiO₂ solar photocatalytic treatment for irradiation period of 60 min were presented in Table 4.30.

Table 4.30. Characteristic properties of organic matter present in 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	FI _{sync280}	SFI _{sync280}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg			
Initial	0.0059	0.0094	0.0316	0.0422	0.847	4.98	51.1	60.3	1.33
t=0	0.0070	0.0115	0.0321	0.0418	0.908	4.60	49.6	54.6	1.68
60 min	0.0024	0.0038	0.0203	0.0356	2.91	1.22	24.1	8.27	1.84

Upon irradiation for 60 min in 0.25% Fe-TiO₂ solar photocatalytic treatment at which 4.28 log reduction of *E. coli* in IsoT was attained (Table 4.29), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0024, UV₃₆₅: 0.0038, UV₂₈₀: 0.0203, UV₂₅₄: 0.0356, and NPOC as 2.91 mg/L. After t_{irr}=60 min, calculated SUVA₂₅₄ as 1.22 L/m mg represents that the remaining organic matter could be considered having hydrophilic and aliphatic character (Edzwald et al., 1985).

Due to the degradation of protein content, $FI_{sync280}$ displayed a decrease at the end of 60 irradiation period. $SFI_{sync280}$ expressed a decrease after t_{irr} =60 min in accordance with the increased NPOC content due to released organic material from bacteria. Moreover, $FI_{sync470}$ of initial, t=0, and t_{irr} =60 min conditions were detected as 0.624, 0.753, and 0.0948, respectively. Almost insignificant $SFI_{sync470}$ of initial, t=0, and t_{irr} =60 min conditions were calculated as 0.737, 0.829, and 0.0326, respectively. At t_{irr} =60 min, FI was calculated as 1.84 that could mostly be related to microbially derived organic matter. Similar finding was also reported by Birben and colleagues for N-doped TiO₂ (Birben et al., 2017a).

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in IsoT solution during 0.25% Fe-TiO₂ solar photocatalytic treatment were displayed in Figure 4.78.



Figure 4.78. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation periods, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli*. However, fluorescence intensities displayed quite low changes after 30 min of irradiation period in accordance with relatively close *E. coli* enumeration

results obtained between t_{irr} =30 min (3.45E+02 CFU/mL) and 60 min (6.13E+01 CFU/mL) (Figure 4.73). Moreover, EEM contour plots of 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution have more intense results than TiO₂ solar photocatalytic treatment in IsoT solution (Figure 4.30) which could be resulting from the lower bacteria reduction percentage of 0.25% Fe-TiO₂ (LRV as 5.90 for TiO₂; LRV as 4.28 for 0.25% Fe-TiO₂ solar photocatalytic inactivation of *E. coli* in IsoT solution) (Tables 4.11 and 4.29).

<u>4.2.3.3.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in IsoT solution. Upon irradiation, as bacteria inactivation products of 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.31.

Table 4.31. Total K, protein, and carbohydrate contents of time dependent E. coli inactivation inIsoT solution upon 0.25% Fe-TiO2 solar photocatalytic treatment.

0.25% Fe-TiO ₂ solar	E. coli in IsoT solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	1.35	1.12	1.15			
Protein, mg/L	1.20	1.07	6.44			
Carbohydrate, mg/L	1.58	2.18	6.00			

Total K content displayed a slight fluctuation as 0.23 mg/L similar to SynTiO₂ (Table 4.22). At t=0, total K showed an immediate adsorption removal as 17.0 % and decreased from 1.35 mg/L (initial) to 1.12 mg/L. After irradiation period of 60 min, almost the same total K concentration was detected with t=0 condition. The reason could be attributed to adsorption of released K⁺ ions from damaged bacteria onto negatively charged *E. coli* cell wall particles and/or 0.25% Fe-TiO₂ particles having dominantly negatively charges sites at experimental pH (\approx 7) due to its pH_{zpc} as 4.80 (Birben et al., 2017).

E. coli in IsoT solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment showed increase in both protein and carbohydrate content at the end of 60 min treatment period. Similar increase was detected for protein and carbohydrate contents after 60 min of irradiation period. Protein concentrations were plotted against $FI_{sync280}$ in order to detect if there is any correlation between these parameters; and a strong correlation as R²=1.00 was detected. Furthermore, NPOC data were plotted against protein and carbohydrate concentrations separately, both parameters displayed strong correlations with NPOC as R²=1.00 and R²=0.99, respectively.

4.2.3.4. Organic matter formation and removal upon 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM solution. UV-vis absorption spectra of organic matter in solution were presented in Figure 4.79. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-320 nm in the slight presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀) originating from the components of WM solution. UVvis absorption spectra obtained from organic matter in WM solution displayed a declining trend without any absorption peaks complying UV-vis absorption spectra of humic substance which was represented in Section 2.4. (Figure 2.16a). However, the use of NO₃⁻ containing WM solution resulted in high light absorption at λ < 210 nm originating from the presence of NO₃⁻ having its λ_{max} at approximately 200 nm (Krishnan and Guha, 1934; Edwards at el., 2001).



Figure 4.79. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm (Figure 4.80). FI_{sync} data displayed a significant increase at t_{irr}=40 min followed by a declining trend during solar photocatalytic treatment. It could be indicated that fluorophoric groups could be reemerged and degraded again throughout specified irradiation period. The presence of fluorophores was evident at λ_{emis} > 300 nm displaying a tailing effect rather than a well-defined peak. Moreover, the presence of fluorophoric groups expressing very minor intensities at λ_{emis} ≈470 nm could also be encountered similar to findings attained in IsoT solution (Figure 4.75). Furthermore, in the absence of humic content, FI_{sync470} displayed almost featureless intensities in the range of 0.293-1.47 however dominant FI_{sync280} expressed variations in between 34.6-52.6. The main fluorophoric groups were accepted as FI_{sync280}.



Figure 4.80. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

NPOC expressed variations in the range of 0.755-2.46 mg/L (Figure 4.81). NPOC content displayed a minor decrease between initial (0.957 mg/L) and t=0 (0.755 mg/L) due to adsorptive removal of organic content after introduction of photocatalyst into the system, and then followed by an increasing trend up to t_{irr}=30 min (2.46 mg/L) originating from released organics from the damaged and destructed *E. coli* cells and decreased during solar photocatalytic treatment (1.88 mg/L).



Figure 4.81. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Similar to previous solar photolytic (Figure 4.9) and photocatalytic treatments of *E. coli* in WM solution (Figures 4.33 and 4.57), all UV-vis displayed quite low presence. On the other hand, Color₄₃₆

and UV₃₆₅ expressed lower presence than UV₂₈₀ and UV₂₅₄ due to non-existence and subsequent degradation of color forming moieties of the released organic matrix upon irradiation, and the use of WM solution, which was a colorless solution as well. As the most significant UV-vis parameter, UV₂₅₄ displayed variations in between 0.0424 and 0.0752 being the lowest for t_{irr}=60 min and the highest for t=0 conditions. The role of Fe doping of TiO₂ could not be visualized under the specified experimental conditions.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.82. SUVA₂₅₄ displayed variations in the range of 1.87-9.96 L/m mg during 0.25% Fe-TiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals. Due to minor presence of Color₄₃₆ and UV₃₆₅, calculated SCoA₄₃₆ and SUVA₃₆₅ was almost insignificant. On the other hand, calculated specific UV-vis and fluroscence parameters presented higher presence from prior solar photolytic (Figure 4.10) and photocatalytic treatments conducted in WM solution (Figures 4.34 and 4.58).

Moreover, due to the absence of humic matter in the solution matrix, $SFI_{sync470}$ displayed almost insignificant presence in between 0.119-1.38 however owing to released protein content from damaged bacteria cells, $SFI_{sync280}$ expressed variations in the range of and 18.4-66.8. The highest $SFI_{sync470}$ and $SFI_{sync280}$ was calculated for t=0 condition in accordance with the lowest NPOC content of the sample. The predominant fluorophoric group was recognized as $SFI_{sync280}$ in the absence of HA solution as solution media.



Figure 4.82. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter present under initial condition, upon introduction of 0.25% Fe-TiO₂ (t=0) and at the end of 0.25% Fe-TiO₂ solar photocatalytic treatment for irradiation period of 60 min were presented in Table 4.32.

Table 4.32. Characteristic properties of organic matter present in 0.25% Fe-TiO₂ solar photolytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis S	pectroscop	pic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	FI _{sync280}	SFI _{sync280}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg			
Initial	0.0072	0.0111	0.0343	0.0433	0.957	4.53	52.6	54.9	1.32
t=0	0.0149	0.0289	0.0691	0.0752	0.755	9.96	50.5	66.8	1.23
60 min	0.0039	0.0077	0.0293	0.0424	1.88	2.26	34.6	18.4	1.74

Upon solar photocatalytic treatment using 0.25% Fe-TiO₂ specimen for irradiation period of 60 min, where 2.49 log reduction of *E. coli* in WM was attained (Table 4.29), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0039, UV₃₆₅: 0.0077, UV₂₈₀: 0.0293, UV₂₅₄: 0.0424, and NPOC as 1.88 mg/L.

SUVA₂₅₄ was calculated as 4.53, 9.95, and 2.26 L/m mg at for initial, t=0, and t_{irr}=60 min conditions, respectively. According to findings of Edzwald and colleagues, SUVA₂₅₄ < 3 indicates the presence of hydrophilic and aliphatic content and SUVA₂₅₄ > 4 reveals hydrophobic and aromatic substances (Edzwald et al., 1985). It could be indicated that initial and t=0 solutions had hydrophobic and aromatic characteristics however t_{irr} =60 min showed hydrophilic and aliphatic features. Decreased SUVA₂₅₄ could be due to degradation of aromatic organics after 60 min of irradiation period.

 $FI_{sync280}$ decreased at t_{irr} =60 min due to degradation of protein content released from damaged *E*. *coli* cells. At the same time, calculated $SFI_{sync280}$ displayed a decrease at t_{irr} =60 min in accordance with the increased NPOC due to released proteins from bacteria. On the other hand, $FI_{sync470}$ were detected as 1.08, 1.04, and 0.352 for initial, t=0, and t_{irr} =60 min conditions, respectively. Owing to the degradation of released organic content from damaged *E*. *coli* cells, $FI_{sync470}$ at t_{irr} =60 min expressed a decline. $SFI_{sync470}$ of initial, t=0, and t_{irr} =60 min conditions were calculated as 1.13, 1.38, and 0.187, respectively.

FI of initial, t=0, and t_{irr}=60 min conditions were calculated as 1.32, 1.23, and 1.74, respectively. FI \leq 1.4 represents the presence of humic fluorophores originating from the presence of humic matter and FI \geq 1.9 indicates the existence of microbially derived organic material (Sen-Kavurmaci and Bekbolet, 2014). Besides the absence of HA as a solution matrix, calculated FI values of initial and t=0 samples was as less than 1.4, the reason could be attributed to the formation of "in-situ generated humic-like matter" originating from extracellular and/or intracellular organic matter founding in the structure of *E. coli* (Uyguner-Demirel et al., 2018). After 60 min of irradiation period, calculated FI was 1.74 which could mostly be related to microbially derived organic content, was the same with Se-N co-doped TiO₂ presented by Birben and colleagues (Birben et al., 2017a).

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution during 0.25% Fe-TiO₂ solar photocatalytic treatment were displayed in Figure 4.83.



Figure 4.83. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation times released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli*. However, at 40 min of irradiation period, increased intensities could be visualized at Region I. It could be indicated that, enhanced protein release from bacteria occurred at t_{irr}=40 min, increased protein content could also be followed via FI_{sync280} data at t_{irr}=40 min (Figure 4.81). Moreover, different from TiO₂ solar photocatalytic treatment of *E. coli* in WM solution (Figure 4.35), higher fluorescence intensities were detected from EEM fluorescence contour plots with the use of 0.25% Fe-TiO₂.

<u>4.2.3.5.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in WM solution. Upon irradiation, as bacteria inactivation products of 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.33.

0.25% Fe-TiO ₂ solar	E. coli in WM solution				
photocatalytic treatment	Initial	t=0	60 min		
Total K, mg/L	4.11	4.16	3.98		
Protein, mg/L	1.79	1.29	9.15		
Carbohydrate, mg/L	2.81	0.911	4.44		

Table 4.33. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Total K content displayed a slight increase at t=0 which could be resulting from K⁺ ion release from *E. coli* cells after adsorption of photocatalyst particles onto the cell surface. After an irradiation period of 60 min, total K content showed a decrease (3.98 mg/L), that could be due to K⁺ ion intake via passive transport by bacteria. Moreover, the reason of high bacteria count result at t_{irr}=60 min as 2.95E+03 CFU/mL (Figure 4.73) could be attributed to the use of excessive K⁺ ions in solution by bacteria to maintain inner osmatic pressure of cells and increase the possibility of survival.

After solar photocatalytic treatment of *E. coli* in WM solution in the presence of 0.25% Fe-TiO₂, increased protein and carbohydrate concentrations were detected. However, increase in protein concentration (1.29 mg/L for t=0; 9.15 mg/L for t_{irr}=60 min) was more evident that than the increase in carbohydrate content (0.911 mg/L for t=0; 4.44 mg/L) after 60 min of solar irradiation period.

Protein contents were plotted against $FI_{sync280}$ and NPOC separately in order to detect if there is any correlation between these parameters; and protein concentrations displayed a strong correlation with both $FI_{sync280}$ and NPOC parameters as R^2 =1.00 and R^2 =0.99, respectively. Furthermore, carbohydrate and NPOC contents showed a strong correlation with each other as R^2 =0.85, as well.

4.2.3.6. Organic matter formation and removal upon 0.25% Fe-TiO₂ solar photocatalytic treatment of E. coli in HA solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.84 and 4.85. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains, UV absorbing centers as well as color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀). On the other hand, HA_i displayed distinctly different trend in wavelength region below λ =230 as expected. Upon introduction of 0.25% Fe-TiO₂ (t=0) into the reaction medium, increased absorbances were recorded similar to TiO₂ solar photocatalytic treatment of *E. coli* in HA solution (Figure 4.36). It could be attributed to mechanistic effect of photocatalyst on E. coli cell wall, which could cause cell wall damage and release of organic matter into the solution. UV-vis absorption spectra did not follow an irradiation time dependent decreasing order in contradistinction to previous solar photocatalytic treatments carried out with undoped TiO2 specimens (Figures 4.36 and 4.60). It could be indicating that emergence and removal of light absorbing centers could be pronounced throughout 60 min of irradiation duration. Moreover, higher absorbances were detected in accordance with bare TiO₂ specimens as TiO₂ (Figure 4.36) and SynTiO₂ (Figure 4.60).



Figure 4.84. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =370 nm and two minor peaks at λ_{emis} =440 nm and 480 nm representing the presence of aromatic hydrocarbons in the solution (Dujmov and Sučević, 1990) (Figure 4.85). After 30 min of irradiation period, FI_{sync} data displayed a declining trend. Moreover, in comparison to previous solar photolytic and photocatalytic treatments of *E. coli* in HA solution (Figures 4.13, 4.37, and 4.61), the highest synchronous scan fluorescence intensities were measured in the use of 0.25% Fe-TiO₂.

On the other hand, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 60.7-198 and 9.96-114, respectively (Figure 4.86). $FI_{sync280}$ of sole HA_i could not be detected however $FI_{sync280}$ of initial solution increased to 9.96 indicating the presence of protein content in the solution. The main increase in $FI_{sync280}$ was detected between initial and t=0 conditions, therefore; it could be indicated that, detach of protein containing structures from cell wall and increased protein release due to cell wall damage could be occurred right after the photocatalyst and *E. coli* contact.



Figure 4.85. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Along with the presence of HA solution, NPOC expressed variations in the range of 4.21-6.67 mg/L (Figure 4.86). NPOC content increased up to t=0 condition (4.42-6.22 mg/L) and followed by a decreased trend in t_{irr} =10 min (4.21 mg/L) and again followed by fluctuations in a range of 4.21-6.67 mg/L during solar photocatalytic treatment. The highest NPOC was detected at t_{irr} =60 min due to released organic content from bacteria. Specified UV-vis parameters expressed higher presence relative to previous solar photolytic (Figure 4.14) and photocatalytic treatment sets of *E. coli* in HA solution (Figures 4.38 and 4.62). However, variations in UV-vis parameters did not followed an

irradiation time dependent order. As the most significant parameter, UV_{254} showed changes in the range of 0.4851-0.7778, being the highest for t_{irr}=30 min and the lowest for irradiation period of 60 min conditions.



Figure 4.86. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.87. SUVA₂₅₄ displayed variations in between 7.27-14.3 L/m mg during 0.25% Fe-TiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals.



Figure 4.87. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Furthermore, $SFI_{sync470}$ and $SFI_{sync280}$ expressed variations in the range of 10.2-34.6 and 1.67-22.2, respectively. In the presence of HA solution as a reaction matrix, the key fluorophoric group was recognized as $FI_{sync470}$. According to undetected $FI_{sync280}$ of HA_i, $SFI_{sync280}$ could not be calculated as well.

Characteristic properties of organic matter present under initial and t=0 conditions, and at the end of solar photocatalytic inactivation of *E. coli* in HA solution for 60 min of irradiation period using 0.25% Fe-TiO₂ were presented in Table 4.34.

Table 4.34. Characteristic properties of organic matter present in 0.25% Fe-TiO2 solar photolytictreatment of *E. coli* in HA solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	БТ	SEI	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	$_{280}$ UV ₂₅₄ mg/L L/m mg $^{\text{Fl}_{\text{sync}4}}$	Γ1 _{sync470}	SF1 _{sync470}	1.1		
Initial	0.0767	0.1663	0.4270	0.5064	5.95	8.51	60.7	10.2	1.05
t=0	0.1106	0.2573	0.6197	0.6858	6.22	11.0	161	25.9	1.08
60 min	0.0674	0.1595	0.4114	0.4851	6.67	7.27	70.6	10.6	1.16

Upon 60 min solar photocatalytic treatment of *E coli* in the presence of HA solution at which 2.09 log *E. coli* reduction was achieved (Table 4.29), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0674, UV₃₆₅: 0.1595, UV₂₈₀: 0.4114, UV₂₅₄: 0.4851, and NPOC as 6.67 mg/L. SUVA₂₅₄ of initial, t=0, and t_{irr}=60 min solutions were calculated as 8.51, 11.0, and 7.27 L/m mg, respectively. The use of HA solution as reaction medium caused relatively high initial NPOC and SUVA₂₅₄ as expected. All calculated SUVA₂₅₄ values were higher than 4 (Figure 4.86), indicating that throughout the 60 min light exposure, samples solutions displayed hydrophobic and aromatic characteristics (Edzwald et al., 1985). Moreover, synchronous scan fluorescence spectra revealing the presence of aromatic hydrocarbons in the solutions (Figure 4.85) could be correlated with high SUVA₂₅₄ data.

 $FI_{sync470}$ increased at t_{irr} =60 min due to both released organic content from damaged *E. coli* cells and present HA solution. At the end of the irradiation period of 60 min SFI_{sync470} almost remained the same as the initial in accordance with slightly increased NPOC. On the other hand, $FI_{sync280}$ measurements of initial, t=0 and t_{irr} =60 min conditions were detected as 9.96, 86.5, and 80.0, respectively. According to NPOC and $FI_{sync280}$, $SFI_{sync280}$ of initial, t=0, and t_{irr} =60 min conditions were calculated as 1.67, 13.9, and 12.0, respectively. FI at t_{irr} =60 min was calculated as 1.16 indicating the presence of fluorophores coming from humic matter containing solution matrix as expected. Similar FI values were also presented by Birben and colleagues (Birben et al., 2017a). EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in HA solution during 0.25% Fe-TiO₂ solar photocatalytic treatment were displayed in Figure 4.88.



Figure 4.88. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Humic-like fluorophore intensities displayed a decreasing trend throughout the treatment period however at t_{irr} =60 min fulvic-like fluorescence intensities expressed an increase, which could be attributed to transformation from humic-like region to fulvic-like with increased humic matter degradation. Moreover, visualized

decreased fluorescence intensities were very low with respect to TiO_2 solar photocatalytic treatment (Figure 4.40). It could be indicated that 0.25% Fe-TiO₂ had a selectivity towards bacteria on contrary to TiO_2 displaying lower *E. coli* reduction (*E. coli* LRV as 2.09 for 0.25% Fe-TiO₂; *E. coli* LRV as 1.45 for TiO_2) (Figures 4.11 and 4.29) and higher organic content degradation (Figures 4.38 and 4.86). Moreover, besides EEM contour plot at t_{irr}=60 min showing similarities with SynTiO₂ (Figure 4.64), with the use of 0.25% Fe-TiO₂ lower intensities were attained, therefore; it could be concluded that 0.25% Fe-TiO₂ had lower bacteria selectivity than SynTiO₂.

4.2.3.7. Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in HA solution. Upon destruction of bacteria cells in HA solution via 0.25% Fe-TiO₂ solar photocatalytic treatment, released total K, protein, and carbohydrate contents of in initial, t=0, and t_{irr}=60 min solutions were presented in Table 4.35.

Table 4.35. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

0.25% Fe-TiO ₂ solar	E. coli and HA				
photocatalytic treatment	Initial	t=0	60 min		
Total K, mg/L	0.59	0.44	0.45		
Protein, mg/L	10.4	9.52	14.3		
Carbohydrate, mg/L	2.50	3.09	3.61		

Total K content showed an immediate adsorptive removal as 25.0 % at t=0 and remained stable at t_{irr} =60 min. Low total K content detected at t_{irr} =60 min could be resulting from the adsorption of K⁺ ions onto oppositely charged *E. coli*, photocatalyst, and/or HA surfaces and removal via membrane filtration applied prior to total K analysis. Similar low total K contents were also obtained from previous solar photolytic and photocatalytic experiment sets carried out in HA solution (Tables 4.8, 4.17, and 4.26) complying with the suggestion that HA has an inhibitory effect on the detection of total K concentration.

E. coli in HA solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment displayed increase in both protein and carbohydrate contents at the end of 60 min irradiation period. Protein concentration showed a higher increase (10.4-14.3 mg/L) than carbohydrate concentration (2.50-3.61 mg/L). In comparison with treatments conducted in the presence of IsoT and WM solutions, higher initial protein (10.4 mg/L) and carbohydrate (2.50 mg/L) contents were detected due to the presence of HA solution. No correlation was detected between $FI_{sync280}$ and protein content however NPOC

measurements displayed strong correlations with both protein and carbohydrate concentrations as $R^2=0.72$ and $R^2=0.97$, respectively.

4.2.3.8. Organic matter formation and removal upon 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.89 and 4.90. Irradiation time dependent absorbance changes were different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅ and UV₂₈₀). UV-vis absorption spectra did not follow an irradiation time dependent decreasing order similar to previous SynTiO₂ solar photocatalytic treatment of *E. coli* in sole HA solution (Figures 4.84). On the other hand, the use of "HA and WM" solution as a reaction medium, caused distinctly different trend in UV absorption at λ < 240 nm as expected.



Figure 4.89. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =370 nm and two minor peaks at λ_{emis} =440 nm and 470 nm (Figure 4.90). On the other hand, data at λ_{emis} =370 nm displayed the highest presence in comparison with previous treatment sets conducted in the presence of "HA and WM" solution (Figures 4.18, 4.42, and 4.66) and expressed variations in the range of 13.0-593. From a general perspective, FI_{sync} data displayed an increase between t_{irr}=30 min and 40 min and followed by a drastic decrease at t_{irr}=60 min. Furthermore, similar to TiO₂ and SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution (Figures 4.42 and 4.66), t=0 condition expressed a significant increase in FI_{sync} data. The reason could be

attributed to the possible mechanistic effect of photocatalyst on cell wall structure, causing release of cytoplasmic organic content from *E. coli* cells.

Moreover, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 52.1-130 and 43.1-103, respectively (Figure 4.91). In the presence of HA solution as a constituent of aqueous reaction matrix, $FI_{sync470}$ expressed higher intensities than $FI_{sync280}$ as expected and recognized as the main fluorophoric group. On the other hand, $FI_{sync280}$ "HA and WM" solution and initial condition could not be detected however t=0 displayed an $FI_{sync280}$ as 63.5.



Figure 4.90. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

From a general perspective, NPOC expressed variations in the range of 4.24-5.97 mg/L (Figure 4.91). NPOC content increased after the introduction of bacteria into "HA and WM" solution (4.24-5.97 mg/L) and slightly decreased after the addition of 0.25% Fe-TiO₂ (4.54 mg/L) due to immediate adsorptive removal or organic content as 24.0 %. An increasing trend in NPOC content was recorded up to t_{irr} =20 min (5.46 mg/L) due to the release of organics from damaged and destructed *E. coli* cells and followed by a decrease until t_{irr} =40 min (4.74 mg/L) then again expressed a slight increase during solar photocatalytic treatment (4.88 mg/L).

Specified UV-vis parameters did not follow a continuous irradiation time dependent order similar to SynTiO₂ solar photocatalytic *E. coli* inactivation conducted in the presence of sole HA solution (Figure 4.86). As the most significant parameter, UV_{254} expressed variations in between 0.3395-0.4589, being the highest for t=0 and the lowest for t_{irr}=60 min conditions. A decreasing trend

was observed in all specified UV-vis parameters between t=0 and t_{irr} =30 min conditions, however a significant increase and a drastic decrease followed each other at t_{irr} =40 min and 60 min conditions, respectively.



Figure 4.91. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.92.



Figure 4.92. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

As the most evident specific UV-vis parameter, $SUVA_{254}$ displayed variations in the range of 6.96-100 L/m mg during 0.25% Fe-TiO₂ solar photocatalytic treatment indicating differences in both UV_{254} and NPOC removals. In accordance with obtained specified UV-vis parameters and NPOC contents (4.24-5.97 mg/L), calculated specific UV-vis parameter displayed similar emergence and removal pattern with specified UV-vis parameters (Figure 4.91).

On the other hand, $SFI_{sync470}$ and $SFI_{sync280}$ expressed variations in the range of 9.13-28.7 and 8.84-21.7, respectively. The dominant fluorophoric group was accepted as $SFI_{sync470}$ due to its remarkable presence in comparison to $SFI_{sync280}$. According to undetected $FI_{sync280}$ of "HA and WM" solution and initial condition, their $SFI_{sync280}$ could not be calculated either.

Characteristic properties of organic matter upon initial and t=0 conditions, and at the end of 0.25% Fe-TiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution for 60 min were presented in Table 4.36.

Table 4.36. Characteristic properties of organic matter present in 0.25% Fe-TiO₂ solar photolytic treatment of *E. coli* in "HA and WM" solution.

Irradiation	UV-vis S	Spectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	EI	SEI	ы
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	F1 _{sync470}	SF1sync470	ГІ
Initial	0.0683	0.1461	0.3732	0.4494	5.97	7.53	54.5	9.13	1.16
t=0	0.0763	0.1737	0.4060	0.4589	4.54	10.1	130	28.7	1.21
60 min	0.0377	0.0948	0.2736	0.3395	4.88	6.96	54.8	11.2	1.27

Upon t_{irr}=60 min 0.25% Fe-TiO₂ solar photocatalytic treatment of *E coli* in the presence of "HA and WM" at which 2.14 log *E. coli* reduction was achieved (Table 4.29), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0377, UV₃₆₅: 0.0948, UV₂₈₀: 0.2736, UV₂₅₄: 0.3395, and NPOC as 4.88 mg/L. The use of HA solution as a matrix component caused relatively high initial NPOC and SUVA₂₅₄ as expected. NPOC expressed a decrease after 60 min of irradiation time indicating degradation of organics. SUVA₂₅₄ at t_{irr}=60 min was calculated as 6.96 L/m mg indicating the remaining organic content in solution displayed aromatic and hydrophobic characteristics (Edzwald et al., 1985).

FI_{sync470} decreased at t_{irr}=60 min due to degradation of organic content released from damaged *E. coli* cells and organic matrix content. At the end of 60 min treatment period SFI_{sync470} increased in accordance with decreased NPOC. Moreover, FI_{sync280} of initial, t=0 and t_{irr}=60 min conditions were determined as \leq 0.01, 63.5, and 43.1 respectively. According to NPOC and FI_{sync280} measurements,

SFI_{sync280} values of initial, t=0, and t_{irr}=60 min conditions were calculated as ≤ 0.01 , 14.0, and 8.84 respectively. FI of t_{irr}=60 min was calculated as 1.27 indicating the presence of fluorophores originating from humic content as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in "HA and WM" solution by 0.25% Fe-TiO₂ solar photocatalytic treatment were displayed in Figure 4.93.



Figure 4.93. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores

were mainly originated from fulvic-like and humic-like substances. Humic-like fluorophore intensities displayed a slight decrease throughout the treatment however they had more intense fluorescence than *E. coli* in HA solution set (Figure 4.88) which is a similar pattern that was detected from previous "HA and WM" solar photolytic and photocatalytic treatments (Figures 4.21, 4.45, and 4.69). After t_{irr} =30 min, intensity of fulvic-like fluorophore displayed an increase while humic-like fluorophore intensities decreased slightly, which could be attributed to a shift of treated high molecular size HA towards lower molecular size fractions (Birben et al., 2017b).

4.2.3.9. Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in "HA and WM" solution. As inactivation products of 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min solutions were presented in Table 4.37.

Table 4.37. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-TiO₂ solar photocatalytic treatment.

0.25% Fe-TiO ₂ solar	E. coli in "HA and WM"					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.36	4.29	4.23			
Protein, mg/L	12.6	4.35	16.6			
Carbohydrate, mg/L	4.66	2.65	5.49			

Total K content showed a decrease at t_{irr} =60 min which could be resulting from the adsorption of K⁺ ions onto oppositely charged *E. coli*, photocatalyst, and HA surfaces. Moreover, at t=0 condition total K displayed 1.61 % of immediate adsorptive removal however it was lower than calculated adsorptive total K content removals of IsoT and HA solutions (17.0 % for *E. coli* in IsoT and 25.0 % for *E. coli* and HA). Due to the use of WM solution as an aqueous matrix component, higher total K concentrations were obtained in comparison with IsoT (Table 4.31) and sole HA solutions (Table 4.35).

0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution expressed an increase in both protein and carbohydrate concentration after t_{irr} =60 min owing to release of cell components from *E. coli*. While protein concentration showed a significant increase (12.6-16.6 mg/L), carbohydrate concentration increased slightly (4.66-5.49 mg/L). Protein concentrations displayed no correlation with either FI_{sync280} or NPOC data. Furthermore, no correlation was detected between NPOC and carbohydrate contents.

4.2.3.10. Comparative evaluation of different experimental matrixes on 0.25% Fe-TiO₂ solar photocatalytic inactivation mechanism of *E. coli*. Total K, protein, and carbohydrate concentrations and bacteria enumerations attained upon 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes in comparison with initial, t=0 and t_{irr}=60 min samples were given in Figure 4.94. Total K contents varied in between 0.44-4.36 mg/L being the lowest for t=0 condition of *E. coli* in HA solution and the highest for initial sample of *E. coli* in "HA and WM" solution. Total K content of WM solution containing conditions displayed higher concentrations as expected. Except the presence of sole HA solution, a rough correlation as $R^2 > 0.75$, was detected between total K contents and bacteria counts of initial and t_{irr}=60 solutions, however t=0 data displayed no correlation.



Figure 4.94. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in various aqueous matrixes.

Covering all conditions protein contents varied in between 1.07-16.6 mg/L being the lowest for t=0 sample of *E. coli* in IsoT solution and the highest for t_{irr}=60 min sample of *E. coli* in "HA and WM" solution. Similarly, carbohydrate contents varied in between 0.91-6.00 mg/L being the lowest for t=0 condition of *E. coli* in WM solution and the highest for t_{irr}=60 min sample of *E. coli* in IsoT solution. At t_{irr}=60 min, all solar photocatalytic treatment conditions displayed increased protein contents, with the highest being detected from "HA and WM" solution (16.6 mg/L). All carbohydrate contents at t_{irr}=60 min displayed increasing trends with respect to their initial conditions. Although similar bacteria counts were obtained from WM, HA and "HA and WM" solutions, there were no resemblance detected between protein and carbohydrate contents. Only total K content of WM and "HA and WM" solution as reaction medium.

Released protein and carbohydrate contents were calculated by subtracting initial data from the final contents attained upon irradiation period of 60 min and represented with *E. coli* LRV in Figure 4.95. Despite the higher LRV of IsoT solution as 4.28, released protein content was detected as the highest in *E. coli* in WM solution (7.36 mg/L) which could be attributed to the degradation of released proteins in IsoT solution. In the presence of HA and "HA and WM" solution) and in accordance with that, their released protein and carbohydrate concentrations were also detected as close to each other. It could be indicated that released carbohydrate contents of WM, HA and "HA and WM" solutions were directly proportional to decreased LRV of *E. coli* individually to detect if there was any correlation between these parameters. In the absence of IsoT solution, released protein concentrations and bacteria LRV showed a strong correlation as $R^2=1.0$. Moreover, released carbohydrate concentrations for all solutions as $R^2=0.99$.



Figure 4.95. Released protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon 0.25% Fe-TiO₂ solar photocatalytic treatment in various aqueous matrixes.

Upon 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV₂₈₀ and NPOC data under initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.96. In the presence of HA solution, protein contents and UV₂₈₀ showed increased presence similar to the finding obtained from solar photolytic (Figure 2.24) and SynTiO₂ photocatalytic treatments (Figure 4.72). Similar protein concentrations and FI_{sync280} were detected between initial and t=0 conditions in both IsoT and WM solutions. FI_{sync280} data showed a significant increase after addition of photocatalyst to the solution matrix in only sole HA solution.



 $FI_{sync280}$ of "HA and WM" initial sample was detected as ≤ 0.01 . Excluding the presence of sole HA matrix, $FI_{sync280}$ and protein data expressed a strong correlation as $R^2 > 0.90$.

Figure 4.96. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and bacteria enumeration results of 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in various aqueous matrixes.

HA containing conditions displayed higher NPOC than carbohydrate contents, except $t_{irr}=60$ min condition of "HA and WM" solution. *E. coli* in IsoT and WM solutions showed significant increase in carbohydrate content at $t_{irr}=60$ min, due to released organics from damaged cells. It could be indicated that the presence of HA solution was directly proportional to NPOC content which also complied with the data of previous solar photolytic (Figure 4.24) and photocatalytic treatments (Figures 4.48 and 4.72). In the absence of sole HA solution, NPOC and carbohydrate contents displayed a strong correlation as $R^2=0.86$ for initial conditions. However, t=0 and $t_{irr}=60$ min samples showed a strong correlation as $R^2 > 0.88$ except in the use of sole WM solution.

4.2.4. 0.50% Fe-TiO₂ Solar Photocatalytic Inactivation of E. coli

4.2.4.1. 0.50% Fe-TiO₂ solar photocatalytic *E. coli* inactivation under specified reaction conditions, reductions, and kinetics. E. coli suspension (No=6.18E+05 CFU/mL) was prepared in IsoT solution and subjected to solar photocatalytic treatment using 0.50% Fe-TiO₂ specimen. Upon an irradiation period of 40 min, E. coli count decreased to 3.34E+03 CFU/mL. Following further light exposure of tirr=60 min, E. coli count decreased to 2.23E+03 CFU/mL. Under these conditions LRV was 2.44 and reduction was 99 %. Upon preparation of E. coli (No=6.71E+05 CFU/mL) suspension in WM solution, for an irradiation period of 40 min, E. coli count decreased to 2.04E+03 CFU/mL. Following prolonged light exposure of tirr=60 min, E. coli count reached to 2.18E+02 CFU/mL revealing LRV as 3.49 and reduction as 99.9 %. E. coli (No=6.25E+05 CFU/mL) suspension was prepared in HA solution and for an irradiation period of 40 min, E. coli count decreased to 1.29E+03 CFU/mL. Following further light exposure of tirr=60 min, E. coli count declined to 2.71E+02 CFU/mL expressing LRV as 3.36 and reduction as 99.9 %. E. coli (No=5.34E+05 CFU/mL) suspension was prepared in a solution comprising of "HA and WM" components and was subjected to an irradiation period of 40 min in which E. coli count decreased to 3.54E+04 CFU/mL. Upon light exposure of tirr=60 min, E. coli count further diminished to 7.74E+03 CFU/mL revealing LRV as 1.84 and reduction as 96.8 %. Based on these results, the effect of solution matrix could be displayed in terms of LRV of E. coli in an increasing order as: "HA and WM" < IsoT < HA < WM. Irradiation time dependent E. coli inactivation profiles and first order kinetic parameters were presented in Figure 4.97 and Table 4.38.



Figure 4.97. Irradiation time dependent *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in various solution matrixes.

0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM solution resulted in a rather fast reduction with k=0.135 min⁻¹ and half-life as 5.13 min. In the presence of "HA and WM" solution, a reaction rate constant of k=0.0734 min⁻¹ and half-life as 9.44 min was attained. The effect of reaction medium could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; WM > HA > IsoT > "HA and WM". Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) also followed the same trend as; 9.06E+04 > 7.81E+04 > 5.19E+04 > 3.92E+04 for WM, HA, IsoT, "HA and WM", respectively.

Motrix	First	IDV	Deduction 0/			
Mauix	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min		Reduction, %	
<i>E. coli</i> in IsoT	0.0839	8.26	5.19E+04	2.44	99	
<i>E. coli</i> in WM	0.135	5.13	9.06E+04	3.49	99.9	
<i>E. coli</i> and HA	0.125	5.54	7.81E+04	3.36	99.9	
<i>E. coli</i> in "HA and WM"	0.0734	9.44	3.92E+04	1.84	96.8	

Table 4.38. 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

E. coli in IsoT solution solar photocatalytic treatment with 0.50% Fe-TiO₂ experiment showed lower bacteria reduction than WM and HA aqueous media, as 99 %. *E. coli* LRV was obtained as 2.44. Different from the previous sets of solar photolytic (Table 4.2) and photocatalytic experiments (Tables 4.11, 4.20, and 4.29) *E. coli* in IsoT solution displayed lower LRV from WM and HA solutions. It could be resulting from the presence of optimal osmotic conditions provided by 0.85 % IsoT solution. Moreover, detected reaction rate constant as 0.0839 was lower than N-doped, Sedoped, and Se-N co-doped TiO₂ results presented by Birben and colleagues (Birben et al., 2017a).

0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM solution displayed the highest *E. coli* LRV among other experiment media, as 3.49. Higher bacteria reduction results could be explained with the presence of common anions and cations in the WM solution. The presence of anions and cations could cause production of increased amount of ROS thus enhance the inactivation rate of bacteria.

In the presence of HA solution, similar reduction of bacteria was detected with *E. coli* in WM condition, as 99.9 % however LRV showed a slight decrease as 0.13. According to the study of Birben and colleagues, it could be concluded that 0.50% Fe-TiO₂ showed a faster reduction rate than Se-doped and Se-N co-doped but slower than N-doped TiO₂ (Birben et al., 2017a). Also, LRV of *E. coli* in the presence of 0.50% Fe-TiO₂ was detected as higher than that in the presence of 0.25% Fe-TiO₂ (Table 4.29) therefore it could be concluded that 0.50% Fe-TiO₂ displayed higher selectivity against *E. coli*, rather than HA.

E. coli in "HA and WM" solution upon solar photocatalytic treatment with 0.50% Fe-TiO₂ showed the lowest bacteria reduction, as 96.8 % which could be due to retardation effect of the presence of both WM and HA solutions. It could be resulting from the beneficial role of "HA and WM" solution on *E. coli* survival by acting as a nutrient source and giving resistance to bacteria.

4.2.4.2. Organic matter formation and removal upon 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.98 and 4.99. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-280 nm in the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀) that should be measured in longer wavelengths. The highest absorption was detected at t_{irr}=60 min between λ =205 nm and 320 nm resembling with the UV-vis absorption spectra feature of t_{irr}=60 min sample of TiO₂ solar photocatalytic treatment (Figure 4.26).



Figure 4.98. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at $\lambda_{emis}=280$ nm, especially at t_{irr}=20 and 40 min (Figure 4.99). The highest intensity at $\lambda_{emis}=280$ nm was detected at t_{irr}=20 min (32.8) due to released protein content from damaged bacteria and following t_{irr}=40 min, intensity at $\lambda_{emis}=280$ nm displayed a slight decrease (27.4) however it also expressed a minor peak at $\lambda_{emis}=320$ nm representing the presence of aromatic hydrocarbons originating from released organic content from bacteria (Dujmov and Sučević, 1990). Moreover, the presence of fluorophoric groups expressing very minor intensities as FI_{sync}≈6.5 could also be

encountered at $\lambda_{emis} > 320$ nm. From a general perspective, after $t_{irr}=20$ min, an irradiation time dependent decreasing order in $FI_{sync280}$ was observed. On the other hand, in the absence of humic matter $FI_{sync470}$ expressed very low intensities as in between 0.486-2.54 however dominant $FI_{sync280}$ displayed changes in the range of and 8.97-32.8.



Figure 4.99. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

NPOC expressed variations in the range of 3.47-6.09 mg/L (Figure 4.100). NPOC content increased between initial (3.85 mg/L) and t=0 (6.09 mg/L) and decreased up to t_{irr}=20 min (3.47 mg/L) and again increased to t_{irr}=40 min (5.35 mg/L) due to the release of organics from damaged and destructed *E. coli* cells and followed by a decreasing trend during solar photocatalytic treatment. Specified UV-vis parameters displayed higher values from previous solar photolytic (Figure 4.4) and photocatalytic treatment sets (Figures 4.28, 4.52, and 4.76) carried out in the presence of IsoT solution. The highest data was detected at initial condition as 0.0131 for Color₄₃₆, 0.0213 for UV₃₆₅, 0.0551 for UV₂₈₀, and 0.0670 for UV₂₅₄. All specified UV-vis parameters expressed variations throughout the irradiation duration however a decrease was detected after 60 min of irradiation period.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.101. SUVA₂₅₄ displayed variations in the range of 0.626-1.74 L/m mg during solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ expressing differences in both UV₂₅₄ and NPOC removals. Despite quite low presence of specific UV-vis parameters, they expressed a significant decrease at t=0 condition and followed by an increasing trend until the end of 60 min irradiation period. Furthermore, besides almost insignificant
SFI_{sync470} varied in between 0.137-0.597, SFI_{sync280} expressed changes in the range of and 1.47-9.45. In accordance with the highest FI_{sync280} of irradiation period of 20 min as 32.8 (Figures 4.99 and 4.100), SFI_{sync280} of the same condition (9.45) was calculated as the highest among initial and other irradiation period conditions. In the absence of HA as a solution matrix component, the main fluorophoric group was accepted as SFI_{sync280} similar to previous solar photolytic (Figure 4.5) and photocatalytic *E. coli* inactivation treatments carried out in the presence of IsoT solution (Figures 4.29, 4.53, and 4.77). The role of Fe doping dose of TiO₂ could not be visualized under the specified experimental conditions.



Figure 4.100. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.



Figure 4.101. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

For simplicity purposes, based on the irradiation time dependent variations in UV-vis and fluorescence spectroscopic properties along with NPOC data, characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-TiO₂ (t=0) and at the end of 0.50% Fe-TiO₂ solar photocatalytic for irradiation period of 60 min were presented in Table 4.39.

Table 4.39. Characteristic properties of organic matter present in 0.50% Fe-TiO₂ solar photolytic treatment of *E. coli* in IsoT solution.

Irradiation time, min	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	EI	SEI	EI
	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1 _{sync280}	SI Isync280	ГІ
Initial	0.0131	0.0213	0.0551	0.0670	3.85	1.74	10.9	2.84	1.48
t=0	0.0043	0.0070	0.0249	0.0381	6.09	0.626	8.97	1.47	2.05
60 min	0.0054	0.0086	0.0292	0.0370	3.63	1.02	9.92	2.73	1.49

Upon irradiation of 60 min at which 2.44 log reduction of *E. coli* in IsoT solution was attained (Table 4.38), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0054, UV₃₆₅: 0.0086, UV₂₈₀: 0.0292, UV₂₅₄: 0.0370, and NPOC as 3.63 mg/L. At the end of irradiation period of 60 min, SUVA₂₅₄ was calculated as 1.02 L/m mg represents that the remaining organic matter could be considered as having hydrophilic and aliphatic character (Edzwald et al., 1985).

Due to release and degradation of proteins, $FI_{sync280}$ showed some fluctuation throughout the irradiation period. According to $FI_{sync280}$ and NPOC contents, calculated $SFI_{sync280}$ also exhibited similar variations. On the other hand, initial, t=0, and t_{irr}=60 min $FI_{sync470}$ data were detected as 2.30, 0.837, and 2.54, respectively. Owing to the release of organics, $FI_{sync470}$ increased at the end of 60 min of irradiation period. $SFI_{sync470}$ values of initial, t=0, and t_{irr}=60 min conditions were calculated as 0.597, 0.137, and 0.701, respectively. FI was calculated as 1.49, which was in between the region of humic fluorophores and microbially derived fluorophores, after irradiation period of 60 min (Sen-Kavurmaci and Bekbolet, 2014). The reason could be attributed to degradation of released organic content from damaged and/or destructed bacteria cells.

In accordance with the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in IsoT solution during solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ were also followed and illustrated in Figure 4.102. EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions.



Figure 4.102. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Decreased fluorescence intensities were detected at t=0 which could be attributed to the immediate adsorptive removal of organics in the presence of photocatalyst. At t_{irr}=20 min, increased intensities were detected at Region I due to the increased protein release from bacteria. The presence of improved protein release could also be followed by increase in FI_{sync280} (Figures 4.99 and 4.100) and SFI_{sync280} (Figure 4.101). After 20 min of irradiation, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of released organic matter and reduction of *E. coli*. Fluorescence intensities were displayed quite low decrease after 20 min of irradiation period in accordance with relatively close *E. coli* enumeration results obtained from t_{irr}=20 min (3.87E+03 CFU/mL), 40 min (3.34E+03 CFU/mL), and 60 min (2.23E+03 CFU/mL) (Figure 4.97). EEM contour plots of 0.50% Fe-TiO₂ solar photocatalytic treatment in IsoT solution (Figure 4.78). However, both solar photocatalytic treatment sets expressed quite low changes in fluorescence intensities and bacteria counts (Figures 4.73 and 4.97) after t_{irr}=20 min.

Besides increased NPOC contents of t=0 and t_{irr} =40 min (Figure 4.100), there was no increased fluorescence could be visualized by EEM contour plot of these conditions. The reason could be

attributed to either the masking effect of released polysaccharides on protein contents of the samples or the presence of organic content other than proteins which could contribute the enhanced NPOC detection.

<u>4.2.4.3.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in IsoT solution. During irradiation, as bacteria inactivation products of 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.40.

Table 4.40. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

0.50% Fe-TiO ₂ solar	E. coli in IsoT solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	1.08	1.00	0.83			
Protein, mg/L	1.28	3.64	5.52			
Carbohydrate, mg/L	2.12	2.20	25.9			

Total K content displayed a decrease in both t=0 and irradiation period of 60 min conditions. 7.4 % of total K showed an immediate adsorptive removal at t=0. Moreover, further decrease of total K content at t_{irr} =60 min could be due to adsorption of released K⁺ ions onto oppositely charged *E. coli* cell wall and/or surface of photocatalyst particles and removal by membrane filtration prior total K analysis.

E. coli in IsoT solution upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ displayed increase in both protein and carbohydrate contents at the end of 60 min treatment period. Carbohydrate concentration showed a significant increase (25.9 mg/L) while protein concentration displayed a slight increase (5.52 mg/L). Glycoproteins that are present in the structure of *E. coli*, and Fe³⁺ released from doped-TiO₂ could interfere with the carbohydrate determination process and result in high carbohydrate concentration at the end of irradiation period of 60 min (Martens and Frankenburger, 1990, Rao and Pattabiraman, 1989). Birben and colleagues reported that when sole 0.25% Fe-TiO₂ was irradiated for 60 min in deionized water, 0.013 mg/L iron release was detected (Birben et al., 2017b). However, released Fe³⁺ analysis of both t=0 and t_{irr}=60 min solutions of *E. coli* in IsoT solution upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ were detected as below detection limit (BDL). No correlations were detected neither between NPOC and protein or carbohydrate concentrations, nor between FI_{sync280} and protein contents.

4.2.4.4. Organic matter formation and removal upon 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in WM solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.103 and 4.104. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-280 nm in the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀) that should be measured in longer wavelengths. Only t=0 condition displayed higher absorption than following irradiation period conditions at λ > 220 nm. Similar absorption feature of t=0 was also detected with the use of 0.25% Fe-TiO₂ upon solar photocatalytic treatment of *E. coli* in WM solution (Figure 4.79). On the other hand, absorbance detected at λ < 210 nm were characteristic to NO₃⁻ which was a component of WM solution (Krishnan and Guha, 1934; Edwards at el., 2001).



Figure 4.103. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =280 and a minor peak at λ_{emis} =420 nm revealing the presence of protein and aromatic hydrocarbon content, respectively (Figure 4.104). At λ_{emis} =280 nm, FI_{sync} data showed a significant increase at t_{irr}=20 min and followed by a decreasing trend. However, decreased fluorescence intensities at λ_{emis} =420 nm until 40 min of irradiation time, displayed a drastic increase at t_{irr}=60 min that could be originating from released aromatic organic content from damaged and destructed *E. coli* cells. Obtained changes in FI_{sync} data was not follow an irradiation time dependent decreasing or increasing order which could be indicating that the presence of both degradation and reemergence of fluorophoric groups could be visualized during specified irradiation period. Furthermore, contrary to minor presence of FI_{sync470} in the range of 0.570-2.74, FI_{sync280} displayed variations significant

presence in between 8.71-30.7. The key fluorophoric groups were accepted as $FI_{sync280}$, in the absence of humic matter in solution.



Figure 4.104. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

From a general perspective, NPOC expressed variations in the range of 1.40-6.12 mg/L (Figure 4.105). NPOC content displayed an insignificant decrease between initial (1.49 mg/L) and t=0 (1.40 mg/L) and increased up to t_{irr} =20 min (3.33 mg/L) due to release of organics from damaged and destructed *E. coli* cells, then decreased up to t_{irr} =40 min (3.08 mg/L) and followed by a sharp increase during solar photocatalytic treatment (6.12 mg/L).



Figure 4.105. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

All UV-vis parameters displayed quite low presence however as the most significant parameter, UV_{254} expressed changes in between 0.0340-0.0712 being the lowest for t_{irr}=20 min which could also be considered as similar to t_{irr}= 40 min (0.0354) and 60 min (0.0345), and the highest for t=0 condition.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.106. SUVA₂₅₄ displayed variations in between 0.563-5.09 L/m mg during 0.50% Fe-TiO₂ solar photocatalytic treatment indicating the differences in both UV₂₅₄ and NPOC removals. Furthermore, SFI_{sync470} displayed almost featureless presence in the range of 0.0931-1.28 however dominant SFI_{sync280} expressed variations in between 1.42-15.4.



Figure 4.106. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter present under initial condition, upon introduction of 0.25% Fe-TiO₂ (t=0) and at the end of 0.50% Fe-TiO₂ solar photocatalytic treatment for irradiation period of 60 min were presented in Table 4.41.

Table 4.41. Characteristic properties of organic matter present in 0.50% Fe-TiO₂ solar photolytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis S	pectroscop	pic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ , L/m mg	FI 200	SFI _{sync280}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L		Γ1 _{sync280}		
Initial	0.0090	0.0127	0.0377	0.0454	1.49	3.04	8.88	5.95	1.35
t=0	0.0156	0.0301	0.0676	0.0712	1.40	5.09	21.7	15.5	1.21
60 min	0.0016	0.0037	0.0211	0.0345	6.12	0.563	8.71	1.42	1.92

Upon an irradiation period of 60 min, 3.49 log reduction of *E. coli* in WM solution was attained in the presence of 0.50% Fe-TiO₂ specimen (Table 3.38). Released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0016, UV₃₆₅: 0.0037, UV₂₈₀: 0.0211, UV₂₅₄: 0.0345, and NPOC as 6.12 mg/L. SUVA₂₅₄ was calculated as 0.563 L/m mg at t_{irr}=60 min representing that the remaining organic matter could still be considered as having hydrophilic and aliphatic character (Edzwald et al., 1985). The decrease of SUVA₂₅₄ between t=0 (5.09 L/m mg) and t_{irr}=60 min conditions could be attributed to the degradation of present organic content in the solution during irradiation period. Owing to the degradation of released protein content from damaged *E. coli* cells, FI_{sync280} showed a decrease at t_{irr}=60 min. SFI_{sync280} decreased at t_{irr}=60 min in accordance with the increased NPOC due to released organic material from bacteria. Furthermore, FI_{sync470} of initial, t=0, and t_{irr}=60 min runs were detected as 0.803, 1.79, and 0.570, respectively. SFI_{sync470} values of initial, t=0, and t_{irr}=60 min conditions were calculated as 0.538, 1.28, and 0.0931, respectively. Due to the presence of microbially derived organic material, FI at t_{irr}=60 min was calculated as 1.92, as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution during solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ were displayed in Figure 4.107.



Figure 4.107. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II, and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Obtained increased intensities of Regions I, II, and IV at t=0 condition could be attributed to cell wall damage occurred right after the introduction of photocatalyst into the solution, similar increases could also be followed by synchronous scan fluorescence spectra (Figure 4.104), specified UV-vis parameters (Figure 4.105), and specific UV-vis and fluroscence parameters (Figure 4.106). Following further irradiation period released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli* in WM solution which could be resulting from the lower bacteria LRV of *E. coli* in IsoT solution condition (LRV as 2.44 for *E. coli* in IsoT solution; LRV as 3.49 for *E. coli* in WM solution) (Figure 4.38), also released protein content of *E. coli* in WM solution could possibly be removed throughout the irradiation and result in lower intensities as well.

4.2.4.5. Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in WM solution. As *E. coli* inactivation products of solar photocatalytic treatment of *E. coli* in WM solution in the presence of 0.50% Fe-TiO₂, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.42.

0.50% Fe-TiO ₂ solar	E. coli in WM solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.12	3.86	3.91			
Protein, mg/L	1.08	1.10	14.7			
Carbohydrate, mg/L	0.618	1.25	3.21			

Table 4.42. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Total K content showed a slight fluctuation like TiO_2 solar photocatalytic treatment (Table 4.15). At t=0, total K content displayed an immediate adsorptive removal as 6.3 % and after irradiation period of 60 min, a slight increase in total K was obtained owing to release of K⁺ ions from damaged bacteria cells.

Upon introduction of 0.25% Fe-TiO₂ into the solution (t=0), protein content expressed almost insignificant increase at t=0 however carbohydrate content increased higher than the half of initial condition. Increased carbohydrate content at t=0 could be correlated with increased data obtained

from synchronous scan fluorescence spectra (Figure 4.104), specified UV-vis parameters (Figure 4.105), specific UV-vis and fluroscence parameters (Figure 4.106), and increased EEM intensities at Region IV representing microbial by-products (Figure 4.107). Both protein and carbohydrate contents increased at t_{irr} =60 min. Besides minor increase in carbohydrate content at t_{irr} =60 min (0.618-3.21 mg/L), protein concentration displayed a significant increase (1.08-14.7 mg/L), similar increase in protein content was also detected from SynTiO₂ solar photocatalytic treatment in WM solution (Table 4.24). Protein and FI_{sync280} showed no correlation with each other however NPOC displayed strong correlations with both protein and carbohydrate contents as R²=1.00 and R²=0.94, respectively.

4.2.4.6. Organic matter formation and removal upon 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in HA solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.108 and 4.109. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀). The absorbance of HA_i expressed a rather different trend in λ < 235 nm region as expected. Upon addition of bacteria into the solution (initial), due to organic matter exudates from *E. coli* following 0.45 µm filtration increased absorbances were detected. Moreover, further increase of absorbances at UV region were recorded after the introduction of 0.50% Fe-TiO2 into the reaction medium due to released cytoplasmic organic content from bacteria via mechanistic effect of photocatalyst adsorbed onto the cell wall. On the other hand, after t=0 condition, UV-vis absorption spectra followed an irradiation time dependent decreasing order during 60 min of irradiation period.



Figure 4.108. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed intense peaks at λ_{emis} =370 nm and two minor peaks at λ_{emis} =440 nm and 480 nm that represent the presence of aromatic hydrocarbons in the solution as expected (Dujmov and Sučević, 1990) (Figure 4.109). Significant increase in FI_{sync} data, especially for FI_{sync370}, were observed at t_{irr}= 20 min and 40 min conditions, which could be indicating that the increased released of organic content from damaged bacteria cells. Furthermore, in the presence of HA solution, evidently present FI_{sync470} expressed changes in the range of 67.8-106 however FI_{sync280} displayed lower presence varied in between 4.76-83.7 as expected (Figure 4.110). The main fluorophoric group referred to FI_{sync370}.



Figure 4.109. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

In the presence of HA solution, NPOC contents expressed variations in the range of 5.27-13.7 mg/L (Figure 4.110). NPOC content increased between HA_i (5.52 mg/L) and initial (7.57 mg/L) due to introduction of bacteria into the solution but decreased up to t=0 (5.27 mg/L) owing to immediate adsorptive removal of organic content as 30.4 %. Afterwards, NPOC displayed fluctuations up to 0.44 mg/L until t_{irr}=40 min (5.31 mg/L) and a sharp increase was observed as 13.7 mg/L due to release of organics from damaged and destructed *E. coli* cells during solar photocatalytic treatment. The highest NPOC release was attained upon irradiation of 60 min. All UV-vis parameters expressed high presence due to the use of HA solution. After t=0 condition, all specified UV-vis parameters followed a sequential decreasing trend similar to TiO₂ and SynTiO₂ solar photocatalytic treatment sets (Figures 4.38 and 4.62). However, demonstrated decreasing trend was not evident as in TiO₂ solar

photocatalytic treatment. As the most significant parameter UV_{254} expressed variations in the range of 0.4688-0.5911.



Figure 4.110. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Based on NPOC contents attained under all conditions, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.111.



Figure 4.111. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Most importantly, SUVA₂₅₄ displayed variations in the range of 3.43-11.2 L/m mg during solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ expressing differences in both UV₂₅₄ and

NPOC removals. Furthermore, $SFI_{sync470}$ and $SFI_{sync280}$ expressed variations in the range of 5.68-20.1 and 0.728-12.9, respectively. In presence of HA solution as a reaction medium, the predominant fluorophoric group was accepted as $SFI_{sync470}$.

For comparative evaluation purposes, characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-TiO₂ (t=0) and at the end of 0.50% Fe-TiO₂ solar photocatalytic inactivation of *E. coli* in HA solution for 60 min of irradiation period were presented in Table 4.43.

Table 4.43. Characteristic properties of organic matter present in 0.50% Fe-TiO₂ solar photolytic treatment of *E. coli* in HA solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ , L/m mg	FI _{sync470}	SFI _{sync470}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L				
Initial	0.0909	0.1837	0.4552	0.5390	7.57	7.12	81.3	10.7	1.04
t=0	0.0891	0.1967	0.5119	0.5911	5.27	11.2	106	20.1	1.05
60 min	0.0641	0.1486	0.3942	0.4688	13.7	3.43	77.6	5.68	1.17

Upon 60 min of irradition period, upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂, 3.36 log *E. coli* reduction was achieved in HA solution (Table 4.38). Released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0641, UV₃₆₅: 0.1486, UV₂₈₀: 0.3942, UV₂₅₄: 0.4688, and NPOC as 13.7 mg/L.

SUVA₂₅₄ of initial, t=0, and following irradiation period of 60 min condition were calculated as 7.12, 11.2, and 3.43 L/m mg, respectively. At the beginning of the treatment, sample solutions (initial and t=0) displayed hydrophobic and aromatic features however after t_{irr} =60 min aromaticity of the organic content showed a significant decrease and expressed almost completely aliphatic characteristics (Edzwald et al., 1985). The presence of HA as solution matrix caused relatively high initial NPOC and SUVA₂₅₄ as expected.

 $FI_{sync470}$ displayed a decrease at $t_{irr}=60$ min due to degradation of organic matter in solution however detected $FI_{sync280}$ followed a different pattern. $FI_{sync280}$ measurements of initial, t=0 and $t_{irr}=60$ min conditions were detected as 5.51, 56.1, and 83.7, respectively. Released proteins from damaged bacteria caused an increase in $FI_{sync280}$ data at the end of 60 min of irradiation time. In accordance with increased NPOC at $t_{irr}=60$ min, $SFI_{sync470}$ decreased by almost half with respect to initial condition. On the other hand, $SFI_{sync280}$ values of initial, t=0, and $t_{irr}=60$ min samples were calculated as 0.728, 10.6, and 6.12, respectively. After solar light exposure of 60 min, FI was calculated as 1.17 indicating the presence of fluorophores emerging from present humic matter in the solution as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in HA solution during 0.50% Fe-TiO₂ solar photocatalytic treatment were displayed in Figure 4.112.



Figure 4.112. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in HA solution upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Due to the quenching effect of humic-like components on microbial by-products and protein-like fluorophores, and encapsulation of proteins by humic substances, the presence of Region I, II and IV could not significantly be identified as expected (Tomaszewski et al., 2011; Wang et al., 2015). Both humiclike and fulvic-like fluorophore intensities displayed a decreasing trend throughout the 60 min irradiation period however the decrease was very low with respect to TiO_2 solar photocatalytic treatment (Figure 4.40). It could be attributed that 0.50% Fe-TiO₂ has a selectivity towards bacteria rather than organic matrix similar to the behavior of SynTiO₂ and 0.25% Fe-TiO₂ specimens. Moreover, EEM contour plots of 0.50% Fe-TiO₂ have similarities with 0.25% Fe-TiO₂ (Figure 4.88) but displayed higher intensities than 0.25% Fe-TiO₂; therefore, it could be concluded that 0.50% Fe-TiO₂ has higher affinity for bacteria than 0.25% Fe-TiO₂.

<u>4.2.4.7. Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in HA solution. As *E. coli* inactivation products of 0.50% Fe-TiO₂ solar photocatalytic treatment in HA solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.44.</u>

0.50% Fe-TiO ₂ solar	<i>E. coli</i> and HA					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	0.54	0.57	0.57			
Protein, mg/L	8.55	8.75	11.3			
Carbohydrate, mg/L	3.02	3.99	77.9			

Table 4.44. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Low total K concentrations were obtained from *E. coli* in the presence of HA solution as expected. Total K content showed a slight increase after initial condition and remained stable between t=0 and $t_{irr}=60$ min conditions. Same total K content of t=0 and $t_{irr}=60$ min solutions could be due to the elimination of released K⁺ ions, which adsorbed onto bacteria, HA and photocatalyst surfaces, by membrane filtration taking place prior to total K analysis.

E. coli in HA solution upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ displayed increase in both protein (8.55-11.3 mg/L) and carbohydrate (3.02-77.9 mg/L) content at t_{irr} =60 min. Protein concentrations were plotted against FI_{sync280} and a moderate correlation was detected as R²=0.66. Carbohydrate concentration showed an unexpected significant increase after 60 min of irradiation period (77.9 mg/L) which could be due to an experimental error or by reason of the presence of some interfering compounds like cellulosic fibers and other insoluble carbohydrates, or NO₃⁻ and Fe³⁺ (Martens and Frankenberger, 1990). Moreover, investigators reported that when humic material is treated with H₂SO₄, which was used in phenol-sulfuric acid method, it could give absorbance in the range of λ =350-500 nm and may interfere with the total carbohydrate determination (Sieburth and Jensen, 1969; Meadows and Campbell, 1978). Leached Fe³⁺ content was measured in both t=0 and t_{irr}=60 min samples as 0.032 mg/L and 0.051 mg/L, respectively. Therefore, the reason of high carbohydrate content of t_{irr}=60 min could be attributed to presence of HA, Fe³⁺, and

glycoproteins originating from bacteria. On the other hand, NPOC displayed strong correlations with either protein and carbohydrate contents as $R^2=0.89$ and $R^2=0.92$, respectively.

4.2.4.8. Organic matter formation and removal upon 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution. UV-vis absorption spectra of organic matter in solution were recorded and presented in Figure 4.113. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀).

Different from HA_i, the use of "HA and WM" solution matrix, absorbance characteristics displayed a different trend at $\lambda < 260$ nm. Especially between $\lambda=200-210$ nm a shoulder was detected due to the strong absorption spectrum of NO₃⁻ having its λ_{max} at approximately 200 nm (Krishnan and Guha, 1934; Edwards at el., 2001). Similar absorption behaviors could also be visualized with previous solar photolytic (Figure 4.17) and photocatalytic treatment sets carried out in the presence of "HA and WM" solution (Figures 4.41, 4.65, and 4.89). On the other hand, UV-vis absorption spectra displayed an irradiation time dependent decreasing order during 60 min solar light exposure duration.



Figure 4.113. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =370 nm and a minor peak at λ_{emis} =480 nm (Figure 4.114). Similar to TiO₂ solar photocatalytic treatment of *E. coli* conducted in "HA and WM" solution, with the addition of 0.50% Fe-TiO₂ into

the reaction matrix (t=0), a drastic increase in FI_{sync} data was recorded. It could be indicated that, due to the mechanistic effect of photocatalyst on the cell wall causing release of cytoplasmic material of bacteria, FI_{sync} could be increase with increased organic matter content in solution. From a general view, FI_{sync} data displayed an increase between $t_{irr}=20$ min and 40 min and followed by a significant decrease at $t_{irr}=60$ min.

Furthermore, despite predominant $FI_{sync470}$ expressing variations in between 28.1-95.0, quite low $FI_{sync280}$ displayed changes in the range of 4.07-70.8 (Figure 4.115). The key fluorophoric group was recognized as $FI_{sync470}$ due to the presence of HA in the solution matrix.



Figure 4.114. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Along with the presence of "HA and WM" solution, NPOC expressed variations in the range of 4.76-12.9 mg/L (Figure 4.115). NPOC content slightly decrease between HA_i (4.83 mg/L) and "HA and WM" solutions (4.76 mg/L). After the addition of *E. coli* into the reaction solution (initial condition), a significant increase was recorded in NPOC content (7.07 mg/L). However, at t=0 condition due to an immediate adsorptive removal of organics as 25.9 %, NPOC content displayed a drastic decrease (5.24 mg/L). Afterwards, NPOC expressed fluctuations in a range of 0.38 mg/L up to t_{irr}=40 min (5.62 mg/L) and followed by a sharp increase (12.9 mg/L) due to the release of organics from damaged and destructed bacterial cells during solar photocatalytic treatment. Detected NPOC increase in between t_{irr}=40 min (3.54E+04 CFU/mL for t_{irr}=40 min; 7.74E+03 CFU/mL for t_{irr}=60 min) (Figure 4.97). On the other hand, all specified UV-vis parameters displayed an irradiation time

dependent decreasing order, except slightly increased UV_{280} and UV_{254} at t_{irr}=40 min condition. As the most evident parameter, UV_{254} expressed changes in the range of 0.3151-0.5020, being the highest for HA_i solution and the lowest for t_{irr}=60 min condition.



Figure 4.115. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

Based on NPOC contents, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.116.



Figure 4.116. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

SUVA₂₅₄, as the most significant specific UV-vis parameter, displayed changes in between 2.45-10.4 L/m mg during 0.50% Fe-TiO₂ solar photocatalytic treatment expressing the differences in both UV₂₅₄ and NPOC removals. On the other hand, SFI_{sync470} and SFI_{sync280} expressed variations in the range of 2.18-18.1 and 0.316-7.89, respectively. Due to remarkable presence of SFI_{sync470} in comparison to SFI_{sync280}, it was accepted as the dominant fluorophoric group.

For comparative evaluation purposes, characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-TiO₂ (t=0) and at the end of 0.50% Fe-TiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution for 60 min of solar light exposure were presented in Table 4.45.

Table 4.45. Characteristic properties of organic matter present in 0.50% Fe-TiO₂ solar photolytic treatment of *E. coli* in "HA and WM" solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ , L/m mg	FI 170	SFI _{sync470}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L		Γlsync470		
Initial	0.0724	0.1520	0.3935	0.4727	7.07	6.69	72.0	10.2	1.14
t=0	0.0544	0.1286	0.3567	0.4182	5.24	7.98	95.0	18.1	1.19
60 min	0.0234	0.0661	0.2395	0.3151	12.9	2.45	28.1	2.18	1.27

Upon solar photocatalytic treatment in the presence of 0.50% Fe-TiO₂ for t_{irr} =60 min, at which 1.84 log *E. coli* reduction was achieved in "HA and WM" solution (Table 4.38), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0234, UV₃₆₅: 0.0661, UV₂₈₀: 0.2395, UV₂₅₄: 0.3151, and NPOC as 12.9 mg/L.

SUVA₂₅₄ of initial, t=0, and t_{irr}=60 min conditions were calculated as 6.69, 7.98, and 2.45 L/m mg, respectively. According to findings of Edzwald and colleagues, SUVA₂₅₄ < 3 indicates that sample display hydrophilic and aliphatic features however SUVA₂₅₄ > 4 reveals hydrophobic and aromatic characteristics of solution (Edzwald et al., 1985). Therefore, it could be indicated that whereas initial and t=0 conditions showed hydrophobic and aromatic features, t_{irr}=60 min condition expressed hydrophilic and aliphatic characteristics due to degradation of aromatic organics throughout 60 min of solar light exposure duration. On the other hand, the presence of HA solution as a rection matrix constituent caused relatively high initial NPOC and SUVA₂₅₄ as expected.

 $FI_{sync470}$ decreased at t_{irr}=60 min due to degradation of organic content arising from damaged *E*. *coli* cells and organic matrix. At the end of 60 min irradiation period, calculated $SFI_{sync470}$ decreased in accordance with increased NPOC content. On the other hand, $FI_{sync280}$ measurements of initial, t=0

and t_{irr} =60 min conditions were detected as 15.8, 70.8, and 4.07, respectively. Furthermore, SFI_{sync280} values of initial, t=0, and t_{irr} =60 min conditions were calculated as 2.23, 13.5, and 0.316, respectively. Due to degradation of both humic content and released cytoplasmic organics containing proteins, FI_{sync280} and consequently FI_{sync280} showed a decrease after 60 min of solar irradiation period. FI was calculated as 1.27 for t_{irr} =60 min conditions indicating the presence of fluorophores emerging from present humic matter in experimental solution matrix as expected (Sen-Kavurmaci and Bekbolet, 2014).

In accordance with the UV-vis absorbance and synchronous scan fluorescence spectroscopic analyses, EEM fluorescence contour plots of organic matter during irradiation time dependent *E. coli* inactivation in "HA and WM" solution under 0.50% Fe-TiO₂ solar photocatalytic treatment conditions were also followed and illustrated in Figure 4.117. It should be indicated that both emergence and removal mechanisms were prevailing under the specified conditions.



Figure 4.117. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in the presence of "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like

and fulvic-like fluorophore intensities displayed a decreasing trend throughout 60 min solar light exposure period. However, solar photocatalytic treatment conducted in "HA and WM" solution expressed more intense fluorescence than the ones attained for *E. coli* in HA solution (Figure 4.112) most probably due to retardation effect of WM components on organic matter degradation and bacteria cell destruction. Intensity differences between EEM contour plots of treatments carried out with sole HA and "HA and WM" solution could also be correlated with *E. coli* count obtained at t_{irr} =60 min (2.71E+02 CFU/mL for *E. coli* in sole HA solution and 7.74E+03 CFU/mL for *E. coli* in "HA and WM" solution) (Figure 4.97).

<u>4.2.4.9.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in "HA and WM" solution. Upon solar irradiation, as inactivation products of 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min solutions were presented in Table 4.46.

Table 4.46. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment.

0.50% Fe-TiO ₂ solar	<i>E. coli</i> in "HA and WM"					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.61	4.42	4.35			
Protein, mg/L	6.47	4.66	10.0			
Carbohydrate, mg/L	2.73	1.95	4.25			

Total K content showed a decreasing trend, which could be resulting from the adsorption of released K⁺ ions onto negatively charged *E. coli*, photocatalyst and/or HA surfaces. On the other hand, 4.1 % of immediate adsorptive total K removal was calculated for t=0 condition. A similar pattern was also observed upon solar photocatalytic treatment of *E. coli* in "HA and WM" solution in the presence of 0.25% Fe-TiO₂ (Table 4.37).

E. coli in the presence of "HA and WM" solution upon solar photocatalytic treatment using 0.50% Fe-TiO₂ displayed increase in both protein (6.47-10.0 mg/L) and carbohydrate content (2.37-4.25 mg/L) at the end of 60 min irradiation period due to release of cell components from damaged *E. coli* and the presence of humic matter in solution matrix. Increase in both protein and carbohydrate concentrations could be expressed with strong correlations with increased NPOC content, where R² values were calculated as 0.99 for both relations. On the other hand, protein contents were plotted against $FI_{sync280}$ data and a strong correlation was obtained between these parameters as R²=0.73.

4.2.4.10. Comparative evaluation of different experimental matrixes on 0.50% Fe-TiO₂ solar photocatalytic inactivation mechanism of *E. coli*. Total K, protein, and carbohydrate concentrations and bacteria enumerations of 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes in comparison with initial, t=0, and t_{in}=60 min conditions were given in Figure 4.118. Total K contents varied in between 0.54-4.61 mg/L being the lowest for initial sample of *E. coli* in HA solution and the highest for initial condition of *E. coli* in "HA and WM" solution. Total K content of WM containing reaction media displayed higher presence than IsoT and HA solutions as expected. Except *E. coli* in HA solution, other matrixes showed a decreasing trend in total K concentrations during t_{in} =60 min. The lowest total K contents were obtained from *E. coli* in HA solution, also previous solar photolytic (Figure 4.22) and photocatalytic treatment runs (Figures 4.46, 4.70, and 4.94) displayed similar patterns. Total K contents were plotted against *E. coli* counts and except the presence of sole WM solution a strong correlation was detected as $R^2 > 0.92$.



Figure 4.118. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in various aqueous matrixes.

Covering all conditions protein contents varied in between 1.08-14.7 mg/L being the lowest for initial sample of *E. coli* in WM solution and the highest for t_{irr} =60 min sample of the same reaction matrix. Similarly, carbohydrate concentrations varied in between 0.62-77.9 mg/L being the lowest for initial sample of *E. coli* in WM solution and the highest for t_{irr} =60 min sample of *E. coli* in HA solution. At the end of t_{irr} =60 min, all treatment sets displayed increased protein contents. In the presence of sole HA solution, initial (8.55 mg/L) and t=0 (8.75 mg/L) samples showed higher protein

concentrations than other initial and t=0 condition. Moreover, all carbohydrate concentrations at $t_{irr}=60$ min displayed an increasing trend with respect to their initial contents.

Released protein and carbohydrate concentrations and bacteria LRV of 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes were given in Figure 4.119. Released protein and carbohydrate concentrations were calculated by subtracting initial data from final contents attained upon t_{irr}=60 min. In the presence of sole WM and HA solutions, *E. coli* showed similar LRV however their released protein and carbohydrate contents were detected as significantly different from each other. The highest released protein content was obtained from WM solution (13.6 mg/L) which complied with the findings of previous solar photocatalytic treatment sets (Figures 4.47, 4.71, and 4.95). In the presence of sole HA solution, the highest carbohydrate release (74.9 mg/L) was detected which could be due to the released carbohydrate content from bacteria and present organic matrix, and also possible experimental error and/or interferences which were indicated at Section 4.2.4.7. Between *E. coli* LRV and released protein contents a strong correlation was detected as R²=0.91 except the presence of sole HA solution matrix. Furthermore, released carbohydrate content and LRV of *E. coli* showed a strong correlation in the absence of sole WM solution as R²=0.99.



Figure 4.119. Released protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon 0.50% Fe-TiO₂ solar photocatalytic treatment in various aqueous matrixes.

Upon 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV₂₈₀ and NPOC data under initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.120. UV₂₈₀ values increased in HA solution containing reaction matrixes, like previous solar photolytic (Figure 4.24) and photocatalytic treatment sets (Figures 4.48, 4.72, and 4.96). The presence of HA solution caused the detection of higher protein concentrations for initial and t=0 conditions. In the presence of "HA and WM" solution, $FI_{sync280}$ data

showed a significant increase after addition of photocatalyst into the solution. Except the presence of sole HA solution, initial protein concentration and $FI_{sync280}$ displayed a strong correlation as R²=0.94, however no correlation was detected for t=0 and t_{irr}=60 min conditions.



Figure 4.120. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and bacteria enumeration results of 0.50% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in various aqueous matrixes.

E. coli in IsoT solution showed a significant increase in carbohydrate content at t_{irr} =60 min due to organics released from damaged bacteria. In the presence of sole HA solution, higher NPOC contents were obtained for initial and t_{irr} =60 min samples. A strong correlation as R² = 0.93 was detected for initial carbohydrate and NPOC concentrations. Moreover, for t=0 condition a strong correlation was detected as R²=0.99 in the absence of sole HA solution. Different from initial and t=0 conditions, carbohydrate and NPOC contents of t_{irr} =60 min displayed a rough correlation as R²=0.74 in the absence of "HA and WM" solution.

0.25% Fe-SynTiO₂ solar photocatalytic E. coli inactivation under specified reaction 4.2.5.1. conditions, reductions, and kinetics. E. coli suspension (No=4.03E+05 CFU/mL) was prepared in IsoT solution and subjected to solar photocatalytic treatment using 0.25% Fe-SynTiO₂ as a Fe-doped photocatalyst specimen. Upon irradiation period of 30 min, E. coli count decreased to 2.20E+03 CFU/mL. Following further light exposure of tirr=60 min, E. coli count decreased to 5.60E+02 CFU/mL. Under these conditions LRV was 2.86 and reduction was 99 %. Upon preparation of E. coli suspension (N_o=2.23E+05 CFU/mL) in WM solution, for an irradiation period of 30 min, E. coli count decreased to 1.18E+02 CFU/mL. Upon prolonged light exposure of t_{irr}=60 min, *E. coli* count decreased to 4.00E+01 CFU/mL revealing LRV as 3.75 and reduction as 99.9 %. E. coli $(N_0=2.62E+05 \text{ CFU/mL})$ suspension was prepared in HA solution and exposed to an irradiation period of 30 min, E. coli count decreased to 6.71E+02 CFU/mL. Following further light exposure of tirr=60 min, E. coli count decreased to 4.00E+01 CFU/mL revealing LRV as 3.82 and reduction as 99.9 %. E. coli (N₀=7.83E+05 CFU/mL) suspension was prepared in a solution composed of "HA and WM" components and was subjected to an irradiation period of 30 min in which E. coli count decreased to 8.51E+03 CFU/mL. Following light exposure of tirr=60 min, E. coli count decreased to 2.49E+03 CFU/mL revealing LRV as 2.50 and reduction as 99 %. Based on these results, the effect of solution matrix could be displayed in terms of LRV of E. coli in an increasing order as: "HA and WM" < IsoT < WM < HA. E. coli inactivation profiles and first order kinetic parameters were presented in Figure 4.121 and Table 4.47.



Figure 4.121. Irradiation time dependent *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in various solution matrixes.

0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in the presence of HA solution resulted in a rather fast removal with k=0.152 min⁻¹ and half-life as 4.56 min. In the presence of "HA and WM" solution, a reaction rate constant of k=0.0943 min⁻¹ and half-life as 7.35 min were attained. The effect of reaction media on bacteria inactivation could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; HA > WM > IsoT > "HA and WM". Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) followed as; 7.38E+04 > 4.67E+04 > 3.98E+04 > 3.26E+04 for "HA and WM", IsoT, HA and WM solutions, respectively.

Moteix	First	IDV	Domoval 04		
Mauix	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min		Kellioval, %
<i>E. coli</i> in IsoT	0.116	5.97	4.67E+04	2.86	99.7
<i>E. coli</i> in WM	0.146	4.75	3.26E+04	3.75	99.9
<i>E. coli</i> and HA	0.152	4.56	3.98E+04	3.82	99.9
<i>E. coli</i> in "HA and WM"	0.0943	7.35	7.38E+04	2.50	99.7

Table 4.47. 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

E. coli in IsoT solution upon solar photocatalytic treatment with 0.25% Fe-SynTiO₂ experiment showed lower bacteria reduction than WM and HA media, as 99.7 %. The reason could be attributed the use of IsoT solution providing higher resistance to bacteria against solar photocatalytic process due to optimal osmotic pressure. However, obtained *E. coli* LRV as 2.86 was lower than LRV of 0.25% Fe-TiO₂ (4.28) (Table 4.29) could be indicating no relevance between same doping concentration among bare TiO₂ species as TiO₂ and SynTiO₂. Moreover, detected reaction rate constant as 0.116 was lower than N-doped, Se-doped, and Se-N co-doped TiO₂ results presented by Birben and colleagues (Birben et al., 2017a).

0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution showed higher bacteria reduction than IsoT, as 99.9 %. Common anions and cations present in the solution could cause production of extra ROSs and enhanced bacteria inactivation rate.

In the presence of HA solution, 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* displayed the highest *E. coli* LRV among other experiment media, as 3.82. Due to this high LRV it could be suggested that 0.25% Fe-SynTiO₂ has selectivity against bacteria rather than organic matrix. Moreover, Birben and colleagues, which they work with other doped TiO₂ species on the degradation of HA and reduction of *E. coli*, achieved higher reaction rate with N-doped and lower reaction rates with Se-doped and Se-N co-doped TiO₂ photocatalyst specimens than 0.25% Fe-SynTiO₂ (Birben et al., 2017a).

0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution displayed the lowest bacteria LRV, as 2.50. It could be owing to two different effects of solution matrix as inhibition on the photocatalytic removal process of bacteria and providing resistance to bacteria by playing a role as a nutrient source.

4.2.5.2. Organic matter formation and removal upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.122 and 4.123. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-290 nm excluding the presence of aromatic domains and color forming moieties that should be measured in longer wavelengths. UV spectral features expressed similarities to the ones attained upon solar photolytic (Figure 4.2) and photocatalytic treatments (Figures 4.26, 4.50, 4.74 and 4.98).



Figure 4.122. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =280 nm for all irradiation periods and also a minor peak was detected at λ_{emis} =440 nm of t_{irr}=10 min datum representing the presence of aromatic hydrocarbons originating from released organic content from bacteria (Dujmov and Sučević, 1990) (Figure 4.123). Furthermore, the presence of fluorophoric groups expressing very minor intensities as FI_{sync} < 5.5 could also be encountered at λ_{emis} > 300 nm. Similar low peaks around FI_{sync}=2-10 were also detected from solar photolytic (Figure 4.3) and 0.50% Fe-TiO₂ solar photocatalytic treatment (Figure 4.99) of *E. coli* in IsoT solution conditions. An irradiation time dependent decreasing order in FI_{sync280} data was observed. On the



Figure 4.123. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

NPOC contents slightly increased between initial (3.82 mg/L) and t=0 min (4.16 mg/L) and besides slight decrease at t_{irr} =10 min (4.09 mg/L), followed by an increasing trend until t_{irr} =40 min (5.65 mg/L) due to the release of organics from damaged and destructed *E. coli* cells, then displayed a sharp decrease at t_{irr} =60 min (4.25 mg/L) (Figure 4.124).



Figure 4.124. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

All UV-vis parameters displayed quite low presence (Figure 4.124). The highest UV₂₅₄ was attained upon t_{irr} =60 min (0.0378) as could also be considered as similar to t_{irr} =40 min (0.0369) although variations were insignificant. UV₂₈₀ displayed almost insignificant changes varied in between 0.0201-0.0249. Due to non-existence and subsequent degradation of color forming moieties of the released organic matrix upon irradiation and the usage of IsoT solution, Color₄₃₆ and UV₃₆₅ expressed minor presence similar to previous solar photolytic (Figure 4.4) and photocatalytic treatment sets (Figures 4.28, 4.52, 4.76, and 4.100). The role of both Fe doping of TiO₂ and 0.25% Fe doping on different TiO₂ specimens as TiO₂ and SynTiO₂ could not be visualized under the specified experimental conditions.

Based on NPOC contents expressing variations in the range of 3.82-5.65 mg/L, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.125. As the most significant parameter, SUVA₂₅₄ displayed changes in the range of 0.653-0.911 L/m mg during 0.25% Fe-SynTiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals. The highest SUVA₂₅₄ was calculated for initial (0.911 L/m mg) condition as could also be considered as similar to t_{irr}=60 min (0.889 L/m mg). Moreover, SUVA₂₈₀ displayed noticeable presence varied in between 0.412-0.652 L/m mg being the highest for initial and the lowest for irradiation period of 20 min conditions. Although SCoA₄₃₆ and SUVA₃₆₅ were also presented, respective specified parameters were very low in comparison to total NPOC contents of the organic matter present in solution.



Figure 4.125. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Furthermore, SFI_{sync470} displayed very low chances in between 0.117-0.712 expressing insignificant contribution of fluorophores detected at λ_{emis} =470 nm. However, SFI_{sync280} expressed considerable variations in the range of 2.38-6.30. In the absence of HA as a solution matrix component, the predominant fluorophoric group was accepted as SFI_{sync280}.

For simplicity purposes, based on the irradiation time dependent variations in UV-vis and fluorescence spectroscopic properties along with NPOC data, characteristic properties of organic matter under initial condition, upon introduction of 0.25% Fe-SynTiO₂ (t=0) and at the end of solar photocatalytic for irradiation period of 60 min were presented in Table 4.48.

Table 4.48. Characteristic properties of organic matter present in 0.25% Fe-SynTiO₂ solar photolytic treatment of *E. coli* in IsoT solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ , L/m mg	FI 200	SFI _{sync280}	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L		1 ⁻¹ sync280		
Initial	0.0028	0.0049	0.0249	0.0348	3.82	0.911	18.1	4.75	1.33
t=0	0.0023	0.0037	0.0208	0.0303	4.16	0.728	22.8	5.49	1.65
60 min	0.0032	0.0052	0.0235	0.0378	4.25	0.117	12.9	3.03	1.77

There was no significant difference detected between the initial, t=0 and t_{irr}=60 min UV-vis spectroscopic parameter results of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. Upon irradiation period of 60 min at which 2.86 log reduction of *E. coli* in IsoT solution was attained (Table 4.47), released organic matter expressed the following specified UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0032, UV₃₆₅: 0.0052, UV₂₈₀: 0.0235, UV₂₅₄: 0.0378, and NPOC as 4.25 mg/L. SUVA₂₅₄ value at t_{irr}=60 min was calculated as 0.117 L/m mg, which could be considered as insignificant, represents that the remaining organic matter could be having hydrophilic and aliphatic character (Edzwald et al., 1985).

Due to degradation of protein content, $FI_{sync280}$ displayed a decrease at the end of $t_{irr}=60$ min. Moreover, $SFI_{sync280}$ expressed a decrease at $t_{irr}=60$ min (4.75 for initial; 3.03 for $t_{irr}=60$ min) in accordance with the increased NPOC. Furthermore, $FI_{sync470}$ of initial, t=0, and $t_{irr}=60$ min conditions were detected as 2.72, 0.623, and 0.499, respectively. According to $FI_{sync470}$ and NPOC, calculated $SFI_{sync470}$ of initial, t=0, and $t_{irr}=60$ min samples, as 0.712, 0.150, and 0.117, respectively, were almost insignificant. FI at $t_{irr}=60$ min was calculated as 1.77 which was between indication value of organic matter, originating from humic matter or microbial content (Sen-Kavurmaci and Bekbolet, 2014). Calculated FI could mostly be related to microbially derived organic matter. Similar finding was also reported by Birben and colleagues for N-doped TiO₂ (Birben et al., 2017a). EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in IsoT solution during 0.25% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.126.



Figure 4.126. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation periods released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli*. Moreover, it could be indicated that due to lower bacteria reduction of 0.25% Fe-SynTiO₂ than SynTiO₂ (*E. coli* LRV as 2.86 for 0.25% Fe-SynTiO₂; *E. coli* LRV as 5.64 for SynTiO₂) (Tables 4.20 and 4.47), EEM contour plots of 0.25% Fe-SynTiO₂ have more intense results than SynTiO₂ solar photocatalytic treatment (Figure 4.54).

<u>4.2.5.3.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in IsoT solution. As bacteria inactivation products of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr} =60 min solutions were presented in Table 4.49.

E. coli in IsoT solution 0.25% Fe-SynTiO₂ solar photocatalytic treatment Initial t=060 min Total K, mg/L 1.94 0.85 1.05 Protein, mg/L 3.83 4.33 4.94 Carbohydrate, mg/L 1.52 1.27 78.0

Table 4.49. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content showed an immediate adsorptive removal as 56.2 % at t=0, and slightly increased at t_{irr} =60 min due to K⁺ ion release from *E. coli* cells having damaged cell walls. Similar high immediate adsorptive removal of total K was also detected from TiO₂ solar photocatalytic treatment carried out with IsoT solution (48.0 %) (Section 4.2.1.3).

E. coli in IsoT solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment displayed increase in both protein and carbohydrate content at the end of 60 min treatment period. No correlation was detected between protein concentrations and FI_{sync280} results however a strong correlation was obtained from protein and NPOC contents as R^2 =0.82. Carbohydrate content significantly increased at t_{irr}=60 min (78.0 mg/L) and similar finding was also detected from SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution (45.0 mg/L) (Table 4.22). The usage of H₂SO₄ at the carbohydrate concentration determination process, which could cause the formation of degradation products absorbing light in the same spectral band with carbohydrate s, and the presence of glycoproteins and Fe³⁺ could be the cause of unexpected high carbohydrate results (Dawson and Liebezeit, 1981; Martens and Frankenburger, 1990; Josefsson et al. 1972; Rao and Pattabiraman, 1989). However, for both t=0 and irradiation period of 60 min conditions of *E. coli* in IsoT solution upon solar photocatalytic treatment in the presence of 0.25% Fe-SynTiO₂ Fe³⁺ release was detected as BDL. On the other hand, carbohydrate and NPOC concentrations displayed a strong correlation as R^2 =1.00.

4.2.5.4. Organic matter formation and removal upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in WM solution were presented in Figures 4.127 and 4.128. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-280 nm in the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀) that should be measured in longer wavelengths. A slightly increasing trend in between λ =240 nm and 280 nm could be visualized under all irradiation period conditions, which could be due to released organic aromatic content from lysed *E. coli* cells and EPS surrounding cell wall (Birben et al., 2021a). Similar absorption pattern was also detected from previous solar photolytic (Figure 4.7) and photocatalytic treatments carried out in the presence of WM solution (Figures 4.31, 4.55, 4.79, and 4.103).



Figure 4.127. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed minor peaks at λ_{emis} =280 nm as FI_{sync} in the range of 14.6-23.5. However, initial, t=0, t_{irr}=10 min and 30 min samples displayed intense peaks at λ_{emis} =420 nm. The highest FI_{sync} as 213 was detected from t_{irr}=10 min at λ_{emis} =420 nm, representing the presence of aromatic hydrocarbons (Dujmov and Sučević, 1990), could be attributed to carbohydrate release from bacteria. Intense peaks at λ_{emis} =420 nm were also attained in previous solar photolytic (Figure 4.8), SynTiO₂ (Figure 4.56) and 0.50% Fe-TiO₂ (Figure 4.104) photocatalytic treatment sets carried out in WM solution however the highest FI_{sync} (213) was detected upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution. On the



Figure 4.128. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

All UV-vis parameters displayed very low presence however as the most significant parameter, UV_{254} expressed variations in between 0.0459-0.0798 (Figure 4.129). UV_{280} , as the second evident UV-vis parameter, displayed changed in the range of 0.0357-0.0665. The highest UV_{254} and UV_{280} values were recorded upon $t_{irr}=20$ min condition.



Figure 4.129. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Furthermore, due to absence and subsequent degradation of color forming moieties of the released extracellular/intracellular organic matrix upon irradiation and the usage of WM solution, $Color_{436}$ and UV_{365} expressed quite low presence similar to previous sets of *E. coli* inactivation in WM solution upon solar photolytic (Figure 4.9) and photocatalytic treatments (Figures 4.33, 4.57, 4.81, and 4.105).

From a general perspective, NPOC expressed changes in the range of 1.09-2.17 mg/L (Figure 4.129). NPOC content displayed a minor decrease between initial (1.62 mg/L) and t=0 (1.47 mg/L) conditions, followed by an increase at t_{irr}=10 min (1.89 mg/L) and decreased at t_{irr}=20 min (1.09 mg/L). an increasing trend was visualized up to following further irradiation period of 40 min (2.17 mg/L) and NPOC displayed a decrease at t_{irr}=60 min (1.64 mg/L). The role of 0.25% Fe doping on different TiO₂ specimens as TiO₂ and SynTiO₂ could not be visualized under the specified experimental conditions.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.130. SUVA₂₅₄ displayed variations in the range of 2.54-7.32 L/m mg during 0.25% Fe-SynTiO₂ solar photocatalytic treatment indicating differences in both UV₂₅₄ and NPOC removals. On the other hand, besides quite low SFI_{sync470} displaying changes in between 0.601-6.05, SFI_{sync280} expressed variations in the range of 6.41-27.2. The main fluorophoric group was recognized as SFI_{sync280} in the absence of HA solution as reaction medium.



Figure 4.130. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial condition, upon introduction of 0.25% Fe-SynTiO₂ as photocatalyst specimen (t=0), and at the end of the 0.25% Fe-SynTiO₂ solar photocatalytic inactivation of *E. coli* in WM solution for irradiation period of 60 min were presented in Table 4.50.

Table 4.50. Characteristic properties of organic matter present in 0.25% Fe-SynTiO₂ solar photolytic treatment of *E. coli* in WM solution.

Irradiation time, min	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ , L/m mg	FI and	SFI _{sync280}	FI
	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L		Γ1 _{sync280}		
Initial	0.0078	0.0114	0.0357	0.0459	1.62	2.83	21.8	13.4	1.39
t=0	0.0132	0.0211	0.0441	0.0544	1.47	3.70	20.5	14.0	1.30
60 min	0.0088	0.0153	0.0474	0.0631	1.64	3.85	17.0	10.4	1.49

Upon irradiation period of 60 min 0.25% Fe-SynTiO₂ solar photocatalytic treatment at which 3.75 log reduction of *E. coli* in WM solution was attained (Table 4.47), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0088, UV₃₆₅: 0.0153, UV₂₈₀: 0.0474, UV₂₅₄: 0.0631, and NPOC as 1.64 mg/L.

After irradiation period of 60 min, SUVA₂₅₄ was calculated as 3.85 L/m mg. Edzwald and colleagues reported that, SUVA₂₅₄ < 3 indicates the presence of hydrophilic and aliphatic content and SUVA₂₅₄ > 4 reveals hydrophobic and aromatic substances (Edzwald et al., 1985). It could be indicated that calculated SUVA₂₅₄ as 3.85 L/m mg would be denoted the higher presence of hydrophobic and aromatic groups in the solution.

 $FI_{sync280}$ showed a decrease at $t_{irr}=60$ min due to degradation of released protein content from damaged bacteria. $SFI_{sync280}$ displayed a decrease at $t_{irr}=60$ min according to increased NPOC. On the other hand, in the absence of humic matter as a component of reaction solution quite low $FI_{sync470}$ were detected as 1.19, 4.56, and 2.37 for initial, t=0, and $t_{irr}=60$ min conditions, respectively. Based on $FI_{sync470}$ and NPOC data, $SFI_{sync470}$ of initial, t=0, and $t_{irr}=60$ min samples were calculated as 0.733, 3.12, and 1.44, respectively.

After irradiation period of 60 min, FI was calculated as 1.49, which was in between the region indicating the presence of humic fluorophores and microbially derived fluorophores (Sen-Kavurmaci and Bekbolet, 2014). The reason of closeness the FI value to the fluorophores originating humic matter could be attributed to degradation of released organic content from damaged *E. coli* cells.
EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution during 0.25% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.131.



Figure 4.131. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation times released microbial by-products and proteins from *E. coli* cells showed a decreasing trend however at t_{irr} =60 min intensities of Region I, II and IV showed an increase. Moreover, at the end of 60 min of irradiation period, Region III representing the presence of fulvic-like fluorophores displayed increased intensities. Also, bacteria count showed a slight

increase between t_{irr} =40 min (3.70E+01 CFU/mL) and 60 min (4.00E+01 CFU/mL) indicating that the presence of bacteria propagation. It could be attributed that bacteria used anions and cations not only to resist the solar photocatalytic process but to reproduce and this propagation caused an increase in the intensities of EEM fluorescence contour plots.

<u>4.2.5.5. Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in WM solution. Upon irradiation total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution were presented in Table 4.51.</u>

0.25% Fe-SynTiO ₂ solar	E. coli in WM solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.11	4.04	3.97			
Protein, mg/L	1.80	2.18	6.95			
Carbohydrate, mg/L	1.95	3.16	74.7			

Table 4.51. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Immediate adsorptive removal of total K content at t=0 was calculated as 1.7 %. Total K content showed a decreasing trend which could be due to released K⁺ ion adsorption onto negatively charged ruptured *E. coli* cell wall and photocatalyst surfaces. Similar decreasing pattern was also obtained from SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution (Table 4.24).

E. coli in WM solution upon solar photocatalytic treatment in the presence of 0.25% Fe-SynTiO₂ specimen displayed increase in both protein and carbohydrate content at the end of 60 min treatment period however carbohydrate concentration showed a significant increase (1.95-74.7 mg/L) like *E. coli* in IsoT solution condition (Table 4.49). Although detection of slight increase in bacteria count, measured carbohydrate concentration was fairly high than expected, therefore; it could be mentioned that the presence of some experimental errors or interferences. The presence of glycoproteins, Fe³⁺, and NO₃⁻ which was a component of WM solution, could be the cause of interferences affecting carbohydrate determination process (Martens and Frankenberger, 1990; Rao and Pattabiraman, 1989). However, Fe³⁺ release was detected as BDL for both t=0 and t_{irr}=60 min conditions. On the other hand, protein concentrations were plotted against FI_{sync280} and a strong correlation was obtained as R²=0.96. Moreover, NPOC content displayed a strong correlation with both protein and carbohydrate contents as R²=0.95 and R²=0.97, respectively.

4.2.5.6. Organic matter formation and removal upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.132 and 4.133. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀). Irradiation time dependent sequential degradation of organic matter was not detected in contradiction to previous sets of solar photocatalytic treatment of *E. coli* in HA solution carried out with TiO₂, SynTiO₂, and 0.50% Fe-TiO₂ (Figures 4.36, 4.60, and 4.108). It should be indicated that, the absorbance of HA_i could be visualized a rather different trend at λ < 235 nm, as expected.



Figure 4.132. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of the emerged organic matter displayed an intense peak at λ_{emis} =380 nm and two minor peaks at λ_{emis} =440 and 480 nm representing the presence of aromatic hydrocarbons (Dujmov and Sučević, 1990) (Figure 4.133). Due to masking effect of humic fluorophores on protein fluorophores, measured intensities at λ_{emis} =280 nm was quite low as expected. After t_{irr}=30 min, an irradiation time dependent decreasing order in FI_{sync} data was observed with a significant decrease after irradiation period of 40 min.

On the other hand, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 55.3-119 and 8.71-27.4, respectively (Figure 4.134). In the presence of HA as solution matrix, the predominant fluorophoric groups were recognized as $FI_{sync470}$. $FI_{sync280}$ of HA_i could not be detected however after



Figure 4.133. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

All UV-vis parameters expressed high presence due to the use of HA solution as expected (Figure 4.134). Specified UV-vis parameters displayed a decrease at $t_{irr}=60$ min. Most importantly, UV₂₅₄ expressed variations in the range of 0.4777-0.5446 being the highest for HA_i solution and the lowest for $t_{irr}=60$ min condition.



Figure 4.134. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

However, obtained differences in specified UV-vis parameters were quite low and similar to $SynTiO_2$ solar photocatalytic treatment of *E. coli* in HA solution (Figure 4.62). Moreover, obtained data could be supported that the interpretation which was made about 0.25% Fe-SynTiO₂ having higher selectivity against *E. coli* rather than HA in Section 4.2.5.1.

Along with the presence of HA solution, NPOC expressed considerable variations in the range of 4.46-14.4 mg/L (Figure 4.134). NPOC content increased between HA_i (4.76 mg/L) and initial (8.48 mg/L) due to possibly EPS content of introduced bacteria, then followed a decreasing trend up to $t_{irr}=20 \text{ min}$ (4.46 mg/L) due to degradation of organics both originating from HA and *E. coli* content. Afterward, followed by an increase to $t_{irr}=30 \text{ min}$ (5.17 mg/L) and again followed by a decrease to $t_{irr}=40 \text{ min}$ (4.75 mg/L) and displayed a sharp increase at $t_{irr}=60 \text{ min}$ (14.4 mg/L) due to further release of organics from damaged and destructed *E. coli* cells.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.135. SUVA₂₅₄ displayed variations in between 3.32-11.5 L/m mg during 0.25% Fe-SynTiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals.

Furthermore, $SFI_{sync470}$ and $SFI_{sync280}$ expressed variations in the range of 3.84-22.4 and 1.03-6.14, respectively. The key fluorophoric group was accepted as $SFI_{sync470}$ in the presence of HA solution. According to undetected $FI_{sync280}$ of HA_i solution, $SFI_{sync280}$ could not be calculated as well.



Figure 4.135. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial condition, upon introduction of 0.25%Fe-SynTiO₂ (t=0) and at the end of 0.50% Fe-TiO₂ solar photocatalytic inactivation of *E. coli* in HA solution for 60 min of irradiation period were presented in Table 4.52.

Table 4.52. Characteristic properties of organic matter present in 0.25% Fe-SynTiO₂ solar photolytic treatment of *E. coli* in HA solution.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹ NPOC, SUVA ₂₅₄ ,		UV-vis Spectroscopic Parameters, cm ⁻¹ NPOC, SUVA ₂₅₄ , FI		rs, cm ⁻¹ NPOC, SUVA ₂₅ .		SEI	БI	
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1 _{sync470}	SF1 _{sync470}	ГІ
Initial	0.0736	0.1622	0.4320	0.5154	8.48	6.08	75.3	8.88	1.05
t=0	0.0840	0.1789	0.4408	0.5208	5.35	9.73	119	22.2	1.06
60 min	0.0661	0.1501	0.3931	0.4777	14.4	3.32	55.3	3.84	1.10

Upon t_{irr}=60 min of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E coli* in HA solution, at which 3.82 log *E. coli* reduction was achieved (Table 4.47), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0661, UV₃₆₅: 0.1501, UV₂₈₀: 0.3931, UV₂₅₄: 0.4777, and NPOC as 14.4 mg/L.

SUVA₂₅₄ of initial, t=0, and t_{irr} =60 min conditions were calculated as 6.08, 9.73, and 3.32 L/m mg, respectively. Differences in SUVA₂₅₄ could be indicate that initial and t=0 solutions expressed hydrophobic and aromatic features however after t_{irr} =60 min with the degradation of organic matter, aromaticity of the organic content displayed a significant decrease and expressed almost completely aliphatic characteristics (Edzwald et al., 1985). On the other hand, the use of HA solution as aqueous medium caused relatively high initial NPOC and SUVA₂₅₄ as expected.

 $FI_{sync470}$ expressed a decrease at $t_{irr}=60$ min due to degradation of organic content. Calculated $SFI_{sync470}$ displayed a decrease at $t_{irr}=60$ min in accordance with the increased NPOC. On the other hand, $FI_{sync280}$ measurements of initial, t=0 and $t_{irr}=60$ min conditions were detected as 8.71, 23.9, and 23.4, respectively. $FI_{sync280}$ demonstrated an increase at $t_{irr}=60$ min due to released protein content from bacteria. $SFI_{sync280}$ of initial, t=0, and $t_{irr}=60$ min runs were calculated based on NPOC and $FI_{sync280}$ data, as 1.03, 4.47, and 1.62, respectively. Moreover, FI at $t_{irr}=60$ min was calculated as 1.10 representing the presence of fluorophores originating from humic content present in reaction solution as expected.

In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent

E. coli inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.136.



Figure 4.136. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities displayed a decreasing trend throughout solar irradiation duration however obtained intensities at t_{irr} =60 min were quite high with respect to TiO₂ solar photocatalytic treatment (Figure 4.40). It could be concluded that 0.25% Fe-SynTiO₂ has a selectivity towards bacteria

irrespective to TiO₂ due to higher bacteria LRV of 0.25% Fe-SynTiO₂ (*E. coli* LRV as 1.45 for TiO₂; *E. coli* LRV as 3.82 for 0.25% Fe-SynTiO₂) (Tables 4.11 and 4.47).

<u>4.2.5.7.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in HA solution. As bacteria inactivation products of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr} =60 min solutions were presented in Table 4.53.

0.25% Fe-SynTiO ₂ solar	E. coli and HA				
photocatalytic treatment	Initial	t=0	60 min		
Total K, mg/L	0.55	0.55	0.63		
Protein, mg/L	7.05	8.78	13.5		
Carbohydrate, mg/L	2.80	3.12	25.7		

Table 4.53. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content displayed no immediate adsorptive removal at t=0, following t_{irr} =60 min showed a slight increase expressing the presence of K⁺ ion release from *E. coli* cells after 60 min of irradiation. Similar inhibited total K contents were also obtained from previous solar photolytic (Table 4.8) and photocatalytic treatment sets (Tables 4.17, 4.26, 4.35, and 4.44) carried out in the presence of HA solution.

Upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution, both protein and carbohydrate contents displayed increase in at the end of t_{irr} =60 min. No correlation was detected when protein concentrations were plotted against FI_{sync280} data. Carbohydrate concentration showed higher increase (2.80-25.7 mg/L) than protein concentration (7.05-13.5 mg/L) after 60 min of solar irradiation period. Detected significantly increased carbohydrate concentration as 25.7 mg/L could be due to remaining humic content and released carbohydrate content from bacteria. Increased NPOC content (Figure 4.134) and the presence of relatively intense EEM contour plots at t_{irr}=60 min. (Figure 4.136) could support increased protein and carbohydrate contents at t_{irr}=60 min. Protein concentrations and NPOC displayed a moderate correlation with each other as R²=0.67 however carbohydrate and NPOC contents showed a strong correlation as R²=0.88. Moreover, 0.029 mg/L of Fe³⁺ release was detected at t_{irr}=60 min, which could be result in increased carbohydrate concentration due to the interfering presence of Fe³⁺ to total carbohydrate determination process (Martens and Frankenberger, 1990). 4.2.5.8. Organic matter formation and removal upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.137 and 4.138. Irradiation time dependent absorbance changes were significantly different from each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀).

Different from previous sets of solar photolytic (Figure 4.17) and photocatalytic treatments conducted in "HA and WM" solution (Figures 4.41, 4.65, 4.89 and 4.113), except "HA and WM" solution and initial condition, quite low absorbances were detected at $\lambda < 250$ nm. On the other hand, similar to TiO₂ solar photocatalytic treatment (Figure 4.41), after the introduction of photocatalyst into the solution (t=0), a drastic decrease was attained at $\lambda < 230$ nm, which revealing significant surface interactions.

Moreover, UV-vis absorption spectra did not follow an irradiation time dependent decreasing order similar to 0.25% Fe-TiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution (Figures 4.89). The reason could be attributed that the re-emergence of removed light absorbing centers following further irradiation periods.



Figure 4.137. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed two intense peaks at λ_{emis} =390 nm and 470 nm, and a minor peak at λ_{emis} =280 nm, due to presence of HA solution as a

matrix component, as expected (Figure 4.138). After the introduction of photocatalyst into the aqueous reaction solution (t=0) an increase in FI_{sync} at $\lambda_{emis} < 410$ nm, especially at $\lambda_{emis} < 300$ nm was detected. It could be indicated that released cytoplasmic content from *E. coli* with the mechanistic effect of present photocatalyst contained higher proteinaceous content than aromatic hydrocarbons. The most significant decrease in FI_{sync} data was observed after t_{irr}=40 min.

Furthermore, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 29.3-68.7 and 1.14-31.9, respectively (Figure 4.139). The predominant fluorophoric group was determined as $FI_{sync470}$ due to the use of HA solution as a reaction medium constituent.



Figure 4.138. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

NPOC contents expressed variations in the range of 4.56-5.52 mg/L (Figure 4.139). NPOC content displayed a decreasing trend up to t=0 condition (5.52-4.63 mg/L), revealing an immediate adsorptive removal as 16.1 %, and slightly increased up to t_{irr} =10 min (5.02 mg/L) due to release of cytoplasmic organic content from damaged bacteria, and followed by a decreasing trend until t_{irr} =30 min (4.56 mg/L). Afterwards NPOC showed an increasing trend during prolonged solar light irradiation period of 60 min (5.37 mg/L) due to release of organics from damaged and destructed *E. coli* cells. Moreover, obtained increase in NPOC content could also be roughly correlated with continuous but slowed down bacteria decrease between irradiation period of 30 min and 60 min conditions (8.51E+03 CFU/mL for t_{irr} =30 min; 4.24E+03 CFU/mL for t_{irr} =40 min; 2.49E+03 CFU/mL for t_{irr} =60 min) (Figure 4.121).

All specified UV-vis parameters displayed a significant decrease at t=0 condition due to adsorptive removal or organic content via membrane filtration prosess applied prior to spectroscopic analysis (Figure 4.139). Besides an increase recorded at t_{irr} =20 min, specified UV-vis parameters expressed a decrease at t_{irr} =60 min. On the other hand, between t_{irr} =20 min and 40 min quite low differences were measured in specified UV-vis and NPOC parameters however, almost 1 log bacteria reduction was achieved among indicated irradiation periods (4.15E+04 CFU/mL for t_{irr} =20 min; 4.24E+03 CFU/mL for t_{irr} =40 min) (Figure 4.121). Therefore, it could be indicated that between these irradiation periods solar photocatalytic system worked against the destruction of *E. coli* rather than the degradation of already present and released organic content in the solutions. On the other hand, as the most significant parameter, UV₂₅₄ displayed variations in the range of 0.2521-0.5166, being the highest for initial solution and the lowest for t_{irr} =60 min condition.



Figure 4.139. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Based on NPOC contents of the samples, all specific parameters as SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} were calculated and presented in Figure 4.140. SUVA₂₅₄ displayed variations in between 4.70-9.46 L/m mg during 0.25% Fe-SynTiO₂ solar photocatalytic treatment indicating the differences in both UV₂₅₄ and NPOC removals. Similar to behavior of specified UV-vis parameters (Figure 4.139), all calculated specific UV-vis and fluorescence parameters expressed insignificant changes between t_{irr} =20 min and 40 min conditions. On the other hand, beside predominant SFI_{sync470} displaying variations in the range of 5.45-14.4, quite low SFI_{sync280} expressed changes in between 0.213-6.83.



Figure 4.140. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial condition, at t=0 condition, and at the end of the 0.25% Fe-SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution for 60 min were presented in Table 4.54.

Table 4.54. Characteristic properties of organic matter present in 0.25% Fe-SynTiO₂ solar photolytic treatment of *E. coli* in "HA and WM" solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	FLaura 470	SEI	EI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	ГІ
Initial	0.0904	0.1726	0.4317	0.5166	5.46	9.46	66.8	12.2	1.18
t=0	0.0512	0.1122	0.2970	0.3577	4.63	7.73	66.9	14.4	1.19
60 min	0.0267	0.0663	0.1965	0.2521	5.37	4.70	29.3	5.45	1.26

Upon t_{irr}=60 min solar photocatalytic treatment of *E coli* in "HA and WM" solution in the presence of 0.25% Fe-SynTiO₂, at which 2.50 log *E. coli* reduction was achieved (Table 4.47), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0267, UV₃₆₅: 0.0663, UV₂₈₀: 0.1965, UV₂₅₄: 0.2521, and NPOC as 5.37 mg/L. After irradiation period of 60 min, SUVA₂₅₄ was calculated as 4.69, indicating that the presence of hydrophobic and aromatic substances in solution (Edzwald et al., 1985). The use of HA solution caused relatively high initial NPOC and SUVA₂₅₄ values as expected. FI_{sync470} showed a decrease at t_{irr}=60 min due to degradation of organic content released from damaged *E. coli* cells and already present organic matrix content. After solar light exposure period of 60 min SFI_{sync470} decreased while NPOC result displayed a slight decrease

based on initial content. Moreover, $FI_{sync280}$ of initial, t=0 and irradiation period of 60 min conditions were recorded as 8.57, 20.1, and 1.14, respectively. $SFI_{sync280}$ of initial, t=0, and t_{irr}=60 min samples were calculated as 1.57, 4.34, and 0.213, respectively. Due to degradation of released cytoplasmic protein content, both $FI_{sync280}$ and $SFI_{sync280}$ displayed a decrease at t_{irr}=60 min condition. FI at t_{irr}=60 min was calculated as 1.26 expressing the presence of fluorophores coming from present HA solution as expected.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in "HA and WM" solution during 0.25% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.141.



Figure 4.141. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities displayed a decrease throughout irradiation period of 60 min however, more intense fluorescence could be visualized relative to *E. coli* in HA solution (Figure 4.136). The reason could be attributed to the presence of WM solution which acting a retardation effect on both reduction of *E. coli* and degradation of humic content.

<u>4.2.5.9. Mechanistic evaluation of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in "HA and WM" solution.</u> Total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples, as bacteria inactivation products of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution were presented in Table 4.55.

Table 4.55. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment.

0.25% Fe-SynTiO ₂ solar	E. coli in "HA and WM"				
photocatalytic treatment	Initial	t=0	60 min		
Total K, mg/L	4.59	4.47	4.41		
Protein, mg/L	5.10	4.79	10.2		
Carbohydrate, mg/L	5.01	4.52	36.2		

Total K content displayed an immediate adsorptive removal at t=0 as 2.61 % and followed by a further decrease at t_{irr} =60 min condition. Decreased total K content at 60 min of irradiation period could be due to the adsorption of K⁺ ions onto negatively charged *E. coli*, photocatalyst and HA surfaces, and elimination with membrane filtration prior to total K analysis. Similar trends were also obtained from previous Fe-doped solar photocatalytic treatment sets carried out in the presence of "HA and WM" solution (Tables 4.37 and 4.46).

E. coli in "HA and WM" solution upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment displayed increased protein (5.10-10.2 mg/L) and carbohydrate concentration (5.01-36.2 mg/L) results at the end of t_{irr}=60 min treatment period. A moderate correlation was obtained between protein and FI_{sync280} data as R²=0.68. Carbohydrate content showed higher increase than protein. This significantly increased carbohydrate concentration could be due to possible extra absorbance at λ =350-500 nm, which includes the measured absorbance value of carbohydrate determination as λ =490 nm, originating from H₂SO₄ treated remaining humic content (Sieburth and Jensen, 1969;

Meadows and Campbell, 1978). Furthermore, released Fe^{3+} concentration at $t_{irr}=60$ min was detected as 0.008 mg/L, which could also cause an interference in carbohydrate determination process (Martens and Frankenberger, 1990). There were no correlations detected between NPOC contents and neither protein nor carbohydrate concentrations.

4.2.5.10. Comparative evaluation of different experimental matrixes on 0.25% Fe-SynTiO₂ solar photocatalytic inactivation mechanism of *E. coli*. Total K, protein, and carbohydrate concentrations and bacteria count results of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes in comparison with initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.142. Total K contents varied in between 0.55-4.59 mg/L being the lowest for initial and t=0 samples of *E. coli* in HA solution and the highest for initial sample of *E. coli* in "HA and WM" solution. WM solution containing samples displayed higher total K contents than IsoT and HA solutions. Except *E. coli* in HA solution solar photocatalytic treatment set, total K concentrations expressed a decreasing trend. In the absence of sole WM solution, total K contents and *E. coli* counts displayed a strong correlation as $R^2 > 0.89$.



Figure 4.142. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in various aqueous matrixes.

Covering all conditions protein contents varied in between 1.80-13.5 mg/L being the lowest for initial sample of *E. coli* in WM solution and the highest for t_{irr} =60 min sample of *E. coli* in HA solution. Similarly, carbohydrate contents varied in between 1.27-78.0 mg/L being the lowest for t=0 condition and the highest for t_{irr} =60 min sample of *E. coli* in IsoT solution. At t_{irr} =60 min, all matrix

conditions displayed increased protein and carbohydrate concentrations. The highest protein concentrations for initial (7.05 mg/L), t=0 (8.78 mg/L), and t_{irr}=60 min (13.5 mg/L) samples were detected in only HA solution matrix. On the other hand, *E. coli* in IsoT, WM and "HA and WM" solution conditions showed significantly increased t_{irr}=60 min carbohydrate concentrations, which could be resulting from an experimental error as explained in Sections 4.2.5.3, 4.2.5.5, and 4.2.5.9.

Released protein and carbohydrate concentrations and bacteria LRV of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in various solution matrixes were given in Figure 4.143. Released protein and carbohydrate concentrations were calculated by subtracting initial from t_{irr} =60 min contents. While WM solution displayed the highest released protein in previous solar photocatalytic treatment sets (Figures 4.47, 4.71, 4.95, and 4.119), upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment the highest released protein was obtained from HA solution as 6.49 mg/L. Moreover, LRV of *E. coli* in HA solution was detected as the highest as 3.82. It could be concluded that high bacteria LRV caused the highest released protein concentration. The highest released carbohydrate content was calculated for IsoT solution as 76.5 mg/L complying with its highest t_{irr} =60 min carbohydrate content. A strong correlation was detected between released protein contents and *E. coli* LRV as R² = 0.97 except the presence of "HA and WM" solution; however, there was no correlation detected between released carbohydrate concentrations and bacteria LRV.



Figure 4.143. Released protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment in various aqueous matrixes.

Upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, $FI_{sync280}$, UV₂₈₀ and NPOC data under initial, t=0, and irradiation period of 60 min conditions were given in Figure 4.144. In the presence of HA solution, higher UV₂₈₀ values were recorded similar to previous solar photolytic (Figure 4.24) and

photocatalytic treatment sets (Figures 4.48, 4.72, 4.96, and 4.120). The use of HA solution as a matrix component resulted in the detection of higher protein contents for initial and t=0 conditions. In the presence of sole HA solution, $FI_{sync280}$ data showed a significant increase at t=0 condition as 23.9. Excluding the presence of "HA and WM" solution, protein concentrations and $FI_{sync280}$ showed a strong correlation as $R^2 > 0.83$.



Figure 4.144. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and *E. coli* enumeration results of 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in various solution matrixes.

Higher NPOC contents were obtained in conditions containing HA solution, besides increased NPOC of initial and t=0 samples, increased carbohydrate concentrations were detected as well. IsoT and WM solutions expressed significantly high carbohydrate concentrations at t_{irr}=60 min, however NPOC content did not show any substantial increase, which supporting the assumption of the presence of an experimental error in carbohydrate determination process. Carbohydrate and NPOC contents displayed a rough correlation as $R^2 > 0.63$, except the presence of sole HA solution.

4.2.6. 0.50% Fe-SynTiO₂ Solar Photocatalytic Inactivation of E. coli

0.50% Fe-SynTiO₂ solar photocatalytic E. coli inactivation under specified reaction 4.2.6.1. conditions, reductions, and kinetics. E. coli suspension ($N_0=5.42E+05$ CFU/mL) was prepared in IsoT solution and subjected to solar photocatalytic treatment using 0.50% Fe-SynTiO₂. Upon irradiation period of 40 min, E. coli count decreased to 1.81E+03 CFU/mL. Following prolonged light exposure of t_{irr}=60 min, E. coli count declined to 3.80E+01 CFU/mL. Under these conditions LRV was 4.15 and reduction was 99.99 %. Upon preparation of E. coli (N₀=5.95E+05 CFU/mL) suspension in WM solution, for an irradiation period of 40 min, E. coli count decreased to 2.79E+02 CFU/mL. Following further light exposure of tirr=60 min, E. coli count reached to 1.33E+02 CFU/mL revealing LRV as 3.65 and reduction as 99.9 %. E. coli (No=6.44E+05 CFU/mL) suspension was prepared in HA solution and for an irradiation period of 40 min, E. coli count decreased to 7.21E+02 CFU/mL. Upon light exposure of tirr=60 min, E. coli count diminished to 1.27E+02 CFU/mL revealing LRV as 3.71 and reduction as 99.9 %. E. coli (No=5.74E+05CFU/mL) suspension was prepared in "HA and WM" components and was subjected to an irradiation period of 40 min in which *E. coli* count decreased to 1.01E+04 CFU/mL. Following prolonged light exposure of t_{irr}=60 min, *E.* coli count decreased to 3.38E+03 CFU/mL revealing LRV as 2.23 and reduction as 99 %.

Based on these results, the effect of solution matrix could be displayed in terms of LRV of *E*. *coli* in an increasing order as: "HA and WM" < WM < HA < IsoT. *E. coli* inactivation profiles and first order kinetic parameters were presented in Figure 4.145 and Table 4.65.



Figure 4.145. Irradiation time dependent *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in various solution matrixes.

Solar photocatalytic treatment of *E. coli* in the presence of 0.50% Fe-SynTiO₂ in IsoT solution resulted in a rather fast removal with k=0.158 min⁻¹ and half-life as 4.39 min. In the presence of "HA and WM" solution, a reaction rate constant of k=0.0850 min⁻¹ and half-life as 8.15 min were attained. The effect of reaction medium could also be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in a decreasing order as; IsoT > HA > WM > "HA and WM". Irrespective of the minor differences in initial bacteria counts, inactivation rates (CFU/mL min) were ordered as; 9.53E+04 > 8.69E+04 > 8.56E+04 > 4.88E+04 for HA, WM, IsoT, and "HA and WM", respectively.

Motrix	First	IDV	Paduation %			
Maurx	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min		Reduction, 70	
<i>E. coli</i> in IsoT	0.158	4.39	8.56E+04	4.15	99.99	
<i>E. coli</i> in WM	0.146	4.75	8.69E+04	3.65	99.9	
<i>E. coli</i> in HA	0.148	4.68	9.53E+04	3.71	99.9	
<i>E. coli</i> in "HA and WM"	0.0850	8.15	4.88E+04	2.23	99	

Table 4.56. 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli*: Inactivation parameters.

Solar photocatalytic treatment of *E. coli* in IsoT solution in the presence of 0.50% Fe-SynTiO₂ displayed the highest bacteria reduction among other experimental media, as 99.99 %. According to a recent study of Birben and colleagues, reaction rate constant of 0.50% Fe-SynTiO₂, which was detected as 0.158 min⁻¹, was higher than the rate constants of N-doped and Se-N co-doped TiO₂ and lower than Se-doped TiO₂ (Birben et al., 2017a).

E. coli in WM solution upon solar photocatalytic treatment with 0.50% Fe-SynTiO₂ displayed lower reduction than *E. coli* in IsoT solution, as 99.9 %. The presence of WM solution produced an inhibitory effect on the inactivation of bacteria.

E. coli inactivation in the presence of HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment displayed the same *E. coli* reduction with WM solution as, 99.9 %. However, LRV of *E. coli* was slightly higher than that in WM solution, as 3.71. Considering high *E. coli* LRV, it could be suggested that 0.50% Fe-SynTiO₂ has selectivity towards bacteria rather than organic matrix. A reaction rate constant of 0.148 min⁻¹ was obtained for HA solution condition. Birben and colleagues achieved a higher reaction rate constant with N-doped TiO₂ and lower reaction rate constants with Se-doped and Se-N co-doped TiO₂ species on the degradation of HA and *E. coli* (Birben et al., 2017a).

Solar photocatalytic treatment of *E. coli* in "HA and WM" solution showed the lowest bacteria reduction as 99 %. The presence of both HA and WM could retard the bacteria inactivation by inhibiting the process or giving resistance to bacteria.

4.2.6.2. Organic matter formation and removal upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.146 and 4.147. Irradiation time dependent absorbance changes were significantly similar to each other in UV wavelength range of λ =200-240 nm in the absence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀) that should be measured in longer wavelengths. Moreover, resembling with TiO₂ (Figure 4.26) and 0.50% Fe-TiO₂ (Figure 4.98) solar photocatalytic treatments, the highest absorption was detected at t_{irr}=60 min in the wavelength range of λ =205-320 nm.



Figure 4.146. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed an intense peak at λ_{emis} =280 nm (Figure 4.147). An irradiation time dependent decreasing order in FI_{sync280} data was observed up to t_{irr}=40 min and no significant difference was detected between t_{irr}=40 min and 60 min. On the other hand, the presence of fluorophoric groups expressing very minor intensities as FI_{sync} < 5.5 could also be encountered at λ_{emis} > 300 nm. Similar minor peaks around FI_{sync}=2-10 were also detected from solar photolytic (Figure 4.3), 0.50% Fe-TiO₂ (Figure 4.99), and 0.25% Fe-SynTiO₂ (Figure 4.123) solar photocatalytic treatment of *E. coli* in IsoT solution conditions. Moreover, beside very low FI_{sync470} displaying changes in between 0.587-2.27, predominant FI_{sync280} expressed variations in the range of 7.12-29.0 in the absence of humic matter in solution matrix.

NPOC expressed variations in the range of 2.38-6.76 mg/L during 60 min of irradiation period (Figure 4.148). NPOC content increased between initial (2.38 mg/L) and t=0 (3.00 mg/L), remained

nearly stable up to t_{irr}=20 min (2.95 mg/L) and followed by a sharp increase at t_{irr}=40 min (6.76 mg/L) due to release of organics from damaged and destructed *E. coli* cells. Subsequently, a decreasing trend was observed at 60 min of solar photocatalytic treatment period (5.23 mg/L). Despite, all UV-vis parameters displayed quite low values, due to absence and adsorptive removal of color forming structures of released organic matter in the solution, Color₄₃₆ and UV₃₆₅ values of t=0 condition could be considered as negligible (<0.0005). On the other hand, as the most significant parameter, UV₂₅₄ displayed variations in the range of 0.0228-0.0612 being the highest for initial condition.



Figure 4.147. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.



Figure 4.148. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.149. SUVA₂₅₄ displayed variations in the range of 0.704-2.57 L/m mg during 0.50% Fe-SynTiO₂ solar photocatalytic treatment showing differences in both UV₂₅₄ and NPOC removals. In accordance with negligible Color₄₃₆ and UV₃₆₅ values of t=0 condition (Figure 4.148), SCoA₄₃₆ and SUVA₃₆₅ of t=0 solution were calculated as 0.000 and 0.017 L/m mg, respectively. Furthermore, SFI_{sync470} displayed almost insignificant presence in the range of 0.132-0.954 however dominant SFI_{sync280} expressed changes in between 1.09-12.2. The role of both Fe doping of TiO₂ and 0.50% Fe doping on different TiO₂ specimens as TiO₂ and SynTiO₂ could not be visualized under the specified experimental conditions.



Figure 4.149. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial condition, upon t=0 condition, and at the end of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution for irradiation period of 60 min were presented in Table 4.57.

Table 4.57. Characteristic properties of organic matter present in 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution.

Irradiation	UV-vis S	Spectroscop	pic Parame	ters, cm ⁻¹	NPOC, SUVA ₂₅₄ ,	EI	SEI	БI	
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γ1 _{sync280}	SF1 _{sync280}	ГІ
Initial	0.0135	0.0215	0.0516	0.0612	2.38	2.57	29.0	12.2	1.18
t=0	0.0000	0.0005	0.0158	0.0228	3.00	0.760	23.2	7.73	1.53
60 min	0.0097	0.0153	0.0398	0.0524	5.23	1.00	7.12	1.36	1.72

Upon irradiation period of 60 min of 0.50% Fe-SynTiO₂ solar photolytic treatment at which 4.15 log reduction of *E. coli* in IsoT solution was attained (Table 4.56), released organic matter expressed following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0097, UV₃₆₅: 0.0153, UV₂₈₀: 0.0398, UV₂₅₄: 0.0524, and NPOC as 5.23 mg/L. SUVA₂₅₄ at t_{irr}=60 min was calculated as 1.00 L/m mg represents that remaining organic matter could be considered having hydrophilic and aliphatic character (Edzwald et al., 1985). Due to degradation of released protein content, FI_{sync280} displayed a decrease at the end of 60 min of irradiation period. SFI_{sync280} also expressed a decrease at t_{irr}=60 min, in accordance with the increased NPOC result. On the other hand, FI_{sync470} of initial, t=0, and t_{irr}=60 min conditions were detected as 2.27, 0.978, and 0.721, respectively. Based on FI_{sync470} and NPOC data, SFI_{sync470} values of initial, t=0, and t_{irr}=60 min samples were calculated as 0.954, 0.326, and 0.138, respectively. After 60 min of irradiation period, FI was calculated as 1.72 which could mostly be associated with the presence of microbially derived organic matter. Similar finding was also reported by Birben and colleagues for N-doped TiO₂ (Birben et al., 2017a).

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in IsoT solution during 0.50% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.150.

Figure 4.150. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation periods, released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of organic matter and reduction of *E. coli*. EEM contour plots displayed similarities with 0.25% Fe-SynTiO₂ (Figure 4.126) but displayed more intense results at 60 min of irradiation period. Significant irradiation time dependent decreasing trend of fluorescence intensities until t_{irr} =40 min especially at Region I and II could be roughly correlated with decreasing FI_{sync280} data (Figures 4.147 and 4.148). Similar to minor decrease of FI_{sync280} between t_{irr} =40 min and 60 min, EEM fluorescence contour plots displayed quite low decrease in intensities as well.

<u>4.2.6.3.</u> Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in IsoT solution. From a general perspective, as inactivation products of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in IsoT solution total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr}=60 min samples were presented in Table 4.58.

0.50% Fe-SynTiO ₂ solar	E. coli in IsoT solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	0.96	0.79	0.81			
Protein, mg/L	0.683	1.95	5.85			
Carbohydrate, mg/L	2.43	2.24	23.9			

Table 4.58. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in IsoT solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content showed a similar trend with SynTiO₂ (Table 4.22) and 0.25% Fe-TiO₂ (Table 4.49) solar photocatalytic treatments. At t=0, total K expressed an immediate adsorptive removal as 17.7 %, and slightly increased at t_{irr} =60 min due to K⁺ ion release from damaged bacteria cells.

E. coli in IsoT solution upon 0.50% Fe-TiO₂ solar photocatalytic treatment displayed increased protein and carbohydrate concentrations at the end of t_{irr} =60 min due to the release of organics from damaged bacteria cells. Carbohydrate content showed significant increase (23.9 mg/L) while protein concentration displayed a slight increase (5.85 mg/L). Protein concentrations were plotted against FI_{sync280} and NPOC data separately and displayed strong correlations as R²=1.00 with both parameters. Moreover, carbohydrate and NPOC concentrations displayed a strong correlation as R²=0.95.

4.2.6.4. Organic matter formation and removal upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.151 and 4.152. Irradiation time dependent absorbance changes displayed minor differences from each other in UV wavelength range of λ =200-500 nm in the slight presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀) that should be measured in longer wavelengths. Irradiation period of 60 min condition expressed higher absorption at λ > 210 nm which could be due to released organic content from lysed *E. coli* cells and EPS surrounding cell wall (Birben et al., 2021a). However, all irradiation period conditions expressed higher absorption than previous solar photolytic and photocatalytic treatments carried out in the presence of WM solution (Figures 4.7, 4.31, 4.55, 4.79, 4.103, and 4.127).

Figure 4.151. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of all irradiation periods displayed a minor peak at λ_{emis} =280 nm however initial and t_{irr}=20 min samples expressed intense peaks at λ_{emis} =420 nm which could represent the release of organics from damaged bacteria (Figure 4.152). Despite lower intensities attained in 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution, similar intense peaks at λ_{emis} =420 nm were also detected upon 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution (Figure 4.128). An irradiation time dependent decreasing order in FI_{sync} data at λ_{emis} =280 nm was noticeable. On the other hand, FI_{sync470} expressed almost featureless presence in the range of 0.943-1.85 however due to existence of proteinaceous content originating from bacteria FI_{sync280} expressed variations in between 14.6-22.1. In the absence of HA as a solution matrix component, predominant fluorophoric group was recognized as FI_{sync280}.

Figure 4.152. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

All UV-vis parameters expressed low presence however obtained values were higher than prior solar photolytic (Figure 4.9) and photocatalytic treatments of *E. coli* in WM solution (Figures 4.33, 4.57, 4.81, 4.105 and 4.129). UV₂₅₄ which was the most significant parameter, displayed changes in the range of 0.1146-0.2748 being the highest for t_{irr} =60 min condition.

Figure 4.153. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

NPOC content increased up to $t_{irr}=20 \text{ min } (4.07-7.97 \text{ mg/L})$ due to release of organics from damaged and destructed *E. coli* cells, then decreased up to $t_{irr}=40 \text{ min } (4.89 \text{ mg/L})$ and followed by a

sharp increase (13.9 mg/L) during the solar photocatalytic treatment. Significantly increased NPOC at $t_{irr}=60$ min could also be correlated with high UV₂₅₄, which is also a surrogate parameter of DOC.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.154. SUVA₂₅₄ displayed changes in between 1.55-3.66 L/m mg during 0.50% Fe-SynTiO₂ solar photocatalytic treatment indicating differences in both UV₂₅₄ and NPOC removals. Furthermore, SFI_{sync470} displayed almost insignificant presence varied in between 0.112-0.436 however SFI_{sync280} expressed variations in the range of 1.05-5.22. The key fluorophoric group was accepted as SFI_{sync280}.

Figure 4.154. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-SynTiO₂ (t=0) and at the end of 0.50% Fe-SynTiO₂ solar photocatalytic treatment for irradiation period of 60 min were presented in Table 4.59.

Table 4.59. Characteristic properties of organic matter present in 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution.

Irradiation	UV-vis S	UV-vis Spectroscopic Parameters, cm ⁻¹				C, SUVA $_{254}$,	FI	SEI	БI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	F1 _{sync280}	SI Isync280	ГІ
Initial	0.0348	0.0526	0.1230	0.1489	4.07	3.66	21.3	5.22	1.60
t=0	0.0271	0.0415	0.0960	0.1146	4.24	2.70	22.1	5.21	1.49
60 min	0.0699	0.1052	0.2341	0.2748	13.9	1.98	14.6	1.05	1.45

Upon 60 min of solar photocatalytic treatment in the presence of 0.50% Fe-SynTiO₂ at which 3.65 log reduction of *E. coli* in WM solution was attained (Table 4.56), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0699, UV₃₆₅: 0.1052, UV₂₈₀: 0.2341, UV₂₅₄: 0.2748, and NPOC as 13.9 mg/L. SUVA₂₅₄ was calculated as 1.98 L/m mg at t_{irr}=60 min, indicating that samples solution displayed the presence of hydrophilic and aliphatic content (Edzwald et al., 1985). Owing to the degradation of released protein content from *E. coli* cells, FI_{sync280} decreased at t_{irr}=60 min. SFI_{sync280} displayed a decrease at t_{irr}=60 min in accordance with the increased NPOC result. On the other hand, FI_{sync470} were detected as 0.943, 1.85, and 1.56 for initial, t=0, and t_{irr}=60 min conditions, respectively. According to FI_{sync470} and NPOC results, SFI_{sync470} of initial, t=0, and t_{irr}=60 min samples were calculated as 1.45, which was in between the region of humic fluorophores and microbially derived fluorophores (Sen-Kavurmaci and Bekbolet, 2014). The reason could be attributed to degradation of released organic content from damaged bacteria.

EEM fluorescence contour plots of organic matter emerged and removed during irradiation time dependent *E. coli* inactivation in WM solution during 0.50% Fe-SynTiO₂ solar photocatalytic treatment were displayed in Figure 4.155.

Figure 4.155. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots displayed the presence of Regions I, II and IV expressing that the fluorophores were mainly originated from microbial by-products and proteins under all irradiation conditions. Following further irradiation times released microbial by-products and proteins from *E. coli* cells showed a decreasing trend in accordance with the solar photocatalytic degradation of released organic matter and reduction of *E. coli*. However, fluorescence intensities of Region I which indicating the existence of aromatic proteins showed an increase at $t_{irr}=60$ min, similar to increase in UV₂₈₀ which also represent the presence of aromatic proteins especially tyrosine and tryptophane containing ones (Edelhoch, 1948). The reason could be attributed to the release of proteins from damaged *E. coli* cells after 60 min of irradiation period. Moreover, EEM contour plots of 0.50% Fe-SynTiO₂ have lower intensities than 0.25% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution (Figure 4.131).

<u>4.2.6.5. Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in WM solution. As *E. coli* inactivation products of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in WM solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr} =60 min samples were presented in Table 4.60.</u>

0.50% Fe-SynTiO ₂ solar	E. coli in WM solution					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.32	3.91	3.89			
Protein, mg/L	1.03	1.15	10.7			
Carbohydrate, mg/L	1.80	2.06	47.0			

Table 4.60. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content showed a declining trend that is similar to measurements detected from SynTiO₂ (Table 4.24) and 0.25% Fe-SynTiO₂ (Table 4.51) solar photocatalytic treatment of *E. coli* in WM solution. An immediate adsorptive removal of total K as 9.5 % was calculated at t=0. Further decrease of total K concentration could be resulting from adsorption of released K⁺ ions onto oppositely charged *E. coli* cell wall particles and photocatalyst surfaces.

E. coli in WM solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment displayed increase in both protein and carbohydrate content at t_{irr} =60 min. Carbohydrate concentration showed a significant increase (1.08-47.0 mg/L) after 60 min of irradiation period, like *E. coli* in IsoT solution upon SynTiO₂ solar photocatalytic treatment (Table 4.22). Released Fe³⁺ concentration was detected

as 0.006 mg/L after an irradiation time of 60 min. The presence of NO_3^- in WM solution, use of Fe³⁺ doped photocatalyst, and the presence of glycoproteins founding in *E. coli* structure could be interference factors for carbohydrate determination process and the cause of detected high carbohydrate content (Martens and Frankenberger, 1990; Rao and Pattabiraman, 1989). FI_{sync280} and protein contents displayed a strong correlation with each other as R²=0.99. Moreover, both protein and carbohydrate concentrations showed a strong correlation with NPOC data as R²=1.00.

4.2.6.6. Organic matter formation and removal upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.156 and 4.157. Irradiation time dependent absorbance changes were similar to each other in UV wavelength range of λ =200-400 nm due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀). Upon introduction of bacteria into the solution (initial), due to exudating organic matter from *E. coli* following 0.45 µm filtration process increased absorbances were detected. UV-vis absorption spectra did not follow an irradiation time dependent decreasing order similar to previous sets of solar photocatalytic treatment of *E. coli* in HA solution in the presence of 0.25% Fe-TiO₂ (Figure 4.84) and 0.25% Fe-SynTiO₂ (Figure 4.132). It could be indicated that emergence and removal of light absorbing centers could be pronounced throughout 60 min of irradiation duration.

Figure 4.156. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed intense peaks at λ_{emis} =390 nm and 470 nm indicating the presence of aromatic hydrocarbons in the solution as

expected (Dujmov and Sučević, 1990) (Figure 4.157). From a general perspective, after the fluorescence intensity increase at t=0 condition, an irradiation time dependent sequential decreasing order in FI_{sync} data was observed. The reason of increased FI_{sync} at t=0 could be attributed to immediate interaction of 0.50% Fe-SynTiO₂ with HA and *E. coli* cell surface causing mechanistic damage to bacteria cell wall and cytoplasmic organic matter release.

On the other hand, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 41.8-81.9 and 1.02-13.5, respectively (Figure 4.158). Due to the presence of HA solution as reaction medium, $FI_{sync470}$ were recognized as the predominant fluorophoric group, as expected. $FI_{sync280}$ of HA_i and initial conditions could not be detected however at t=0 condition, $FI_{sync280}$ increased to 11.8 representing the presence of proteinaceous content in the sample, which also support the occurrence of mechanistic bacteria cell wall damage of in the presence of photocatalyst specimen.

Figure 4.157. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Along with the presence of HA solution, NPOC expressed variations in the range of 5.76-14.9 mg/L (Figure 4.158). NPOC content showed an increasing trend (5.76-14.9) throughout the solar photocatalytic treatment due to release of organics from damaged and destructed *E. coli* cells, that consistent with gradually decreased *E. coli* counts (Figure 4.145).

Specified UV-vis parameters expressed high presence due to use of HA solution (Figure 4.158). As the most significant parameter, UV_{254} showed changes in between 0.5406-0.6281, being the highest for irradiation period of 20 min and the lowest for 60 min irradiation period conditions. All

other specified UV-vis parameters followed a similar trend with UV_{254} which could be due to release of cytoplasmic organic content at $t_{irr}=20$ min and its degradation occurred during further irradiation periods.

Figure 4.158. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Based on NPOC contents, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.159. SUVA₂₅₄ displayed a decreasing trend (9.50-3.62 L/m mg) during solar photocatalytic treatment indicating differences in both UV₂₅₄ and NPOC removals.

Figure 4.159. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Furthermore, despite dominant $SFI_{sync470}$ displaying changes in between 2.80-14.2, almost insignificant $SFI_{sync280}$ expressed variations in the range of 0.0684-1.09. $SFI_{sync470}$ was accepted as the main fluorophoric group due to its remarkable presence in comparison to $SFI_{sync280}$. Moreover, according to undetected $FI_{sync280}$ of HA_i and initial conditions, $SFI_{sync280}$ could not be detected either.

For simplicity purposes, based on the irradiation time dependent variations in UV-vis and fluorescence spectroscopic properties along with NPOC data, characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-SynTiO₂ (t=0) and at the end of 0.50% Fe-SynTiO₂ solar photocatalytic for irradiation period of 60 min were presented in Table 4.61.

Table 4.61. Characteristic properties of organic matter present in 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution.

Irradiation	UV-vis S	pectroscop	oic Parame	ters, cm ⁻¹	NPOC,	SUVA ₂₅₄ ,	FI	SFI.	БI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	Γlsync470	SF1sync470	ГІ
Initial	0.0970	0.2027	0.5154	0.6099	9.15	6.67	72.3	7.91	1.03
t=0	0.0974	0.2074	0.5102	0.5998	11.1	5.38	80.4	7.21	1.04
60 min	0.0787	0.1777	0.4479	0.5406	14.9	3.62	41.8	2.80	1.08

Upon t_{irr}=60 min of solar photocatalytic treatment of *E coli* in HA solution at which 3.71 log *E*. *coli* reduction was achieved (Table 4.56), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.0787, UV₃₆₅: 0.1777, UV₂₈₀: 0.4479, UV₂₅₄: 0.5406, and NPOC as 14.9 mg/L. Similar UV-vis spectroscopic parameter results were detected from initial and t=0 conditions revealing insignificant surface interactions. After irradiation period of 60 min, SUVA₂₅₄ was calculated as 3.62 L/m mg representing that the remaining organic matter could be considered having hydrophilic and aliphatic character, indicating the occurrence of humic content degradation (Edzwald et al., 1985).

FI_{sync470} displayed a decrease at the end of t_{irr}=60 min due to degradation of organic matter content. In accordance with the increased NPOC, SFI_{sync470} displayed a decrease at t_{irr}=60 min, as well. Moreover, FI_{sync280} measurements of initial, t=0 and t_{irr}=60 min samples were detected as 0.00, 11.8, and 1.02, respectively. Similar to data of FI_{sync470}, FI_{sync280} demonstrated a decreasing trend between t=0 and t_{irr}=60 min conditions owing to degradation of released protein content. SFI_{sync280} of initial, t=0, and t_{irr}=60 min conditions were calculated as \leq 0.001, 1.06, and 0.0684, respectively. Moreover, FI was calculated as 1.08 at 60 min of irradiation period, representing the presence of fluorophores originating from used HA solution as an aqueous medium in the solar photocatalytic treatment, as expected. In a similar manner to the UV-vis absorbance and synchronous scan fluorescence analyses, irradiation time dependent EEM fluorescence contour plots of organic matter emerged and removed during *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment were also followed and displayed in Figure 4.160.

Figure 4.160. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in the presence of HA solution displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities slightly decreased throughout the 60 min of treatment period. It could be concluded that 0.50% Fe-SynTiO₂ has a selectivity towards bacteria in contradiction to TiO₂ having lower bacteria LRV (LRV as 1.08 for TiO₂; LRV as 3.71 for 0.50% Fe-SynTiO₂) (Tables 4.11 and 4.56) and less intense EEM contour plots (Figure 4.40).

<u>4.2.6.7. Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in HA solution. As bacteria inactivation products of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr} =60 min solutions, were presented in Table 4.62.</u>

0.50% Fe-SynTiO ₂ solar	E. coli and HA				
photocatalytic treatment	Initial	t=0	60 min		
Total K, mg/L	0.51	0.55	0.61		
Protein, mg/L	10.8	9.67	16.2		
Carbohydrate, mg/L	3.49	3.16	4.14		

Table 4.62. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in HA solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content showed an increasing trend during 60 min of solar light exposure which was also detected from SynTiO₂ solar photocatalytic treatment of *E. coli* in HA solution (Table 4.26). Release of K^+ ions from damaged bacteria cells caused increased total K concentration at the end of 60 min irradiation period.

E. coli in HA solution upon solar photocatalytic treatment displayed an increase in both protein and carbohydrate content at $t_{irr}=60$ min. Protein concentration (10.8-16.2 mg/L) showed a higher increase than carbohydrate content (3.49-4.14 mg/L). Similar pattern was also detected from solar photocatalytic treatment with 0.25% Fe-TiO₂ in the presence of HA solution (Table 4.35). Increase in protein concentrations displayed a strong correlation with NPOC data as R²=0.77 however, increased carbohydrate and NPOC contents showed a moderate correlation with each other as R²=0.61. On the other hand, no correlation was detected between protein concentrations and FI_{sync280}.

4.2.6.8. Organic matter formation and removal upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution. UV-vis absorption and synchronous scan fluorescence spectra of organic matter in solution were presented in Figures 4.161 and 4.162, respectively. Irradiation time dependent absorbance changes were different from each other in UV wavelength range of λ =200-350 nm, except HA_i solution, due to the presence of aromatic domains and color forming moieties (Color₄₃₆, UV₃₆₅, and UV₂₈₀). The reason of different absorbance characteristics of HA_i solution at λ < 240 nm could be attributed to the absence of NO₃⁻ containing WM solution in the aqueous medium.

Contrary to previous *E. coli* inactivation sets of solar photolytic (Figure 4.17) and photocatalytic treatments in the presence of "HA and WM" solution (Figures 4.41, 4.65, 4.89, 4.113, and 4.137), with the use of 0.50% Fe-SynTiO₂ a sequential irradiation time dependent increasing trend was observed in UV-vis absorption spectra. The reason could be attributed to continuing cytoplasmic

organic content release from damaged *E. coli* cells, also in accordance with low bacteria LRV as 2.23 (Table 4.26).

Figure 4.161. UV-vis absorption spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Figure 4.162. Synchronous scan fluorescence spectra of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Synchronous scan fluorescence spectra of emerged organic matter displayed intense peaks at λ_{emis} =390 nm and 470 nm (Figure 4.162) in the presence of humic content, as excepted. A drastic decrease at λ_{emis} =390 nm was detected between t=0 and t_{irr}=20 min conditions representing that the degradation of humic content in the solution. At t_{irr}=20 min and 40 min conditions, minor peaks at
λ_{emis} =280 nm were detected expressing release of protein content from bacteria. Moreover, similar to solar photolytic (Figure 4.18) and SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution (Figure 4.66), both in "HA and WM" solution and after addition of bacteria into the reaction solution (initial and t=0 conditions), increased fluorescence intensities were observed relative to sole HA_i condition at λ_{emis} < 410 nm. From a general perspective, an irradiation time dependent decreasing order in FI_{sync} data was observed.

On the other hand, $FI_{sync470}$ and $FI_{sync280}$ displayed changes in the range of 31.9-81.9 and 1.60-10.4, respectively (Figure 4.163). Due to the use of HA solution as an aqueous matrix component $FI_{sync470}$ displayed dominant presence and accepted as the key fluorophoric group. $FI_{sync280}$ of HA_i could not be detected however $FI_{sync280}$ of "HA and WM" solution was detected as 4.76.

From a general perspective, NPOC contents expressed variations in the range of 5.76-13.4 mg/L (Figure 4.163). NPOC content increased between HA_i solution (5.76 mg/L) and t=0 condition (7.10 mg/L) and decreased up to t_{irr} =20 min (6.33 mg/L) followed by a sharp increase until t_{irr} =40 min (13.4 mg/L) due to the release of organics from damaged and destructed *E. coli* cells. Subsequently a decreasing trend (11.5 mg/L) was observed during solar photocatalytic treatment.



Figure 4.163. Specified UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

In the presence of HA solution as a reaction matrix component, all specified UV-vis parameters and $FI_{sync470}$ displayed significant presence. As the most evident parameter UV₂₅₄ expressed variations in between 0.4690-0.6023, being the lowest for $t_{irr}=20$ min and the highest for $t_{irr}=60$ min conditions with a remarkable increase at $t_{irr}=40$ min (0.5855). Moreover, detected increase in all specified UV-vis parameters could be roughly correlated with decreased bacteria counts between irradiation period of 20 min and 40 min (8.17E+04 CFU/mL for $t_{irr}=20$ min; 1.01E+04 CFU/mL for $t_{irr}=40$ min) (Figure 4.145). It could be indicated that with the destruction of bacteria, release of cytoplasmic organic content could be enhanced and caused an increase in specified UV-vis parameters and also FI_{sync280}. Moreover, according to inverse relation between FI_{sync280} and FI_{sync470} at $t_{irr}=40$ min, it could be indicated that the released organic content from bacteria could probably had proteinaceous characteristics.

Based on NPOC contents, all specific UV-vis and fluorescence parameters as, SCoA₄₃₆, SUVA₃₆₅, SUVA₂₈₀, SUVA₂₅₄, SFI_{sync470}, and SFI_{sync280} parameters were calculated and presented in Figure 4.164. As the most significant parameter, SUVA₂₅₄ displayed variations in between 4.38-8.87 L/m mg during 0.50% Fe-SynTiO₂ solar photocatalytic treatment expressing differences in both UV₂₅₄ and NPOC removals.

Furthermore, despite dominant SFI_{sync470} displaying changes in between 2.71-14.2, quite low SFI_{sync280} expressed variations in the range of 0.225-1.31. Along with the use of HA solution as an aqueous medium component, SFI_{sync470} was accepted as the main fluorophoric group. According to undetected FI_{sync280} of HA_i solution, its SFI_{sync280} could not be calculated either.



Figure 4.164. Specific UV-vis and fluroscence parameters of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

For comparative evaluation purposes, characteristic properties of organic matter under initial condition, upon introduction of 0.50% Fe-SynTiO₂ (t=0), and at the end of 0.50% Fe-SynTiO₂ solar photocatalytic inactivation of *E. coli* in "HA and WM" solution for 60 min of solar light exposure were presented in Table 4.63.

Table 4.63. Characteristic properties of organic matter present in 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution.

Irradiation	UV-vis Spectroscopic Parameters, cm ⁻¹				NPOC,	SUVA ₂₅₄ ,	EI	SEI	FI
time, min	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	L/m mg	ST 1 _{sync470}	1.1
Initial	0.0881	0.1772	0.4551	0.5451	6.15	8.87	66.1	10.7	1.15
t=0	0.0835	0.1682	0.4280	0.5119	7.10	7.21	63.6	8.95	1.16
60 min	0.1108	0.2030	0.4973	0.6023	11.5	5.25	31.9	2.78	1.25

Upon t_{irr}=60 min of solar photocatalytic treatment of *E coli* in "HA and WM" solution, at which 2.23 log *E. coli* reduction was achieved (Table 4.56), released organic matter expressed the following UV-vis parameters (cm⁻¹); Color₄₃₆: 0.1108, UV₃₆₅: 0.2030, UV₂₈₀: 0.4973, UV₂₅₄: 0.6023, and NPOC as 11.5 mg/L. With respect to increased UV-vis parameters and NPOC content at t_{irr} =60 min, it could be indicated that cytoplasmic organic content release generated throughout the irradiation period and its complete removal could not be achieved in determined irradiation period.

SUVA₂₅₄ of initial, t=0, and t_{irr}=60 min conditions were calculated as 8.87, 7.21, and 5.25 L/m mg, respectively. Calculated SUVA₂₅₄ values were > 4, therefore; it could be indicated that besides a decrease of SUVA₂₅₄ at t_{irr}=60 min, which revealing the degradation of aromatic organic matter in solution, all samples expressed hydrophobic and aromatic characteristics (Edzwald et al., 1985).

 $FI_{sync470}$ decreased at t_{irr} =60 min due to degradation of organic content originating from damaged *E. coli* cells and HA containing organic matrix. At the end of t_{irr} =60 min, calculated SFI_{sync470} showed a decrease in accordance with increased NPOC. On the other hand, FI_{sync280} of initial, t=0 and t_{irr} =60 min samples were detected as 2.79, 1.60, and 4.20, respectively. Different from FI_{sync470}, FI_{sync280} demonstrated an increase at the end of 60 min irradiation period due to released protein content from bacteria. Furthermore, SFI_{sync280} values of initial, t=0, and t_{irr} =60 min conditions were calculated as 0.454, 0.225, and 0.366, respectively. FI at 60 min irradiation period was calculated as 1.25 indicating the presence of fluorophores coming from present humic matter in solution, as expected.

In accordance with the UV-vis absorbance and synchronous scan fluorescence spectroscopic analyses, EEM fluorescence contour plots of organic matter during irradiation time dependent *E. coli*

inactivation in "HA and WM" solution under 0.50% Fe-SynTiO₂ solar photocatalytic treatment conditions were also followed and illustrated in Figure 4.165. It should be noted that both emergence and removal mechanisms were prevailing under the specified conditions.



Figure 4.165. EEM fluorescence contour plots of organic matter: Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

EEM fluorescence contour plots of *E. coli* in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment displayed the presence of Regions III, and V indicating that the fluorophores were mainly originated from fulvic-like and humic-like substances. Both humic-like and fulvic-like fluorophore intensities displayed a slight decrease throughout the treatment period of 60 min however detection of relatively high intensities at irradiation period of 60 min could be indicated that, due to the presence of WM solution as a reaction matrix constituent, a retardation effect could be observed on both degradation of organic matter content and inactivation rate of bacteria.

4.2.6.9. Mechanistic evaluation of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in "HA and WM" solution. As bacteria inactivation products of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in "HA and WM" solution, total K, protein, and carbohydrate contents present in initial, t=0, and t_{irr} =60 min samples, were presented in Table 4.64.

0.50% Fe-SynTiO ₂ solar	E. coli in "HA and WM"					
photocatalytic treatment	Initial	t=0	60 min			
Total K, mg/L	4.56	4.43	4.36			
Protein, mg/L	7.45	16.3	22.8			
Carbohydrate, mg/L	3.01	6.41	84.2			

Table 4.64. Total K, protein, and carbohydrate contents of time dependent *E. coli* inactivation in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment.

Total K content displayed an immediate adsorptive removal at t=0 condition, as 2.85 %. A slight total K content decrease was detected throughout t_{irr} =60 min. Similar pattern was also detected from previous Fe-doped TiO₂ sets carried out in "HA and WM" solution (Tables 4.37, 4.46, and 4.55).

E. coli in "HA and WM" solution upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment displayed increase in both protein (7.45-22.8 mg/L) and carbohydrate (3.01-84.2 mg/L) content at t_{irr} =60 min. No correlation detected between protein and FI_{sync280} data. Carbohydrate concentration at t_{irr} =60 min displayed the highest concentration detected (84.2 mg/L), which could be due to some experimental errors and/or interferences. Moreover, released Fe³⁺ concentration at the end of 60 min irradiation period was detected as 0.009 mg/L. On the other hand, there could be some interfering factors resulting in the detection of high carbohydrate concentration such as; *i*) the use of H₂SO₄ as a strong acid in phenol sulfuric acid method could cause some degradation products which could absorb light in the same band with carbohydrates, *ii*) H₂SO₄ treated humic matter could give extra absorbance in the same spectral band used in the method, *iii*) NO₃⁻ coming from WM solution, *iv*) the use of Fe³⁺ doped photocatalyst, and *v*) glycoprotein content founding in bacteria (Dawson and Liebezeit, 1981; Josefsson et al. 1972; Martens and Frankenberger, 1990; Meadows and Campbell, 1978; Rao and Pattabiraman, 1989; Sieburth and Jensen, 1969). Despite a number of possible interferences, NPOC content displayed strong correlations with both protein and carbohydrate concentrations as R²=0.82 and R²=0.98, respectively.

4.2.6.10. Comparative evaluation of different experimental matrixes on 0.50% Fe-SynTiO₂ solar photocatalytic inactivation mechanism of *E. coli*. Total K, protein, and carbohydrate concentrations and bacteria enumeration results of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes in comparison with initial, t=0, and t_{irr}=60 min samples were given in Figure 4.166. Total K contents varied in between 0.51-4.56 mg/L being the lowest for initial sample of *E. coli* in HA solution and the highest for initial solution of *E. coli* in "HA and WM" solution. WM solution containing conditions showed higher total K contents as expected. Total K and *E. coli* counts

of initial and t=0 conditions displayed a strong correlation as $R^2 > 0.91$ except the presence of IsoT solution. Moreover, a strong correlation was detected between t_{irr}=60 min conditions as R^2 =1.00, in the absence of sole WM solution.



Figure 4.166. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents and bacteria enumeration results of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in various aqueous matrixes.

Covering all conditions protein contents varied in between 0.68-22.8 mg/L being the lowest for initial sample of *E. coli* in IsoT solution and the highest for t_{irr} =60 min sample of *E. coli* in "HA and WM" solution. Similarly, carbohydrate contents varied in between 1.80-84.2 mg/L being the lowest for initial sample of *E. coli* in WM solution and the highest for t_{irr} =60 min condition of *E. coli* in "HA and WM" solution. Protein concentration of all treatment sets displayed an increasing order except sole HA solution which displayed a slight decrease between initial (3.49 mg/L) and t=0 (3.16 mg/L) conditions. Except t_{irr} =60 min condition of *E. coli* in sole HA solution, all carbohydrate concentrations expressed a significant increase at the end of 60 min irradiation period being the highest for "HA and WM" solution (84.2 mg/L). The possible experimental errors could be the reason of detected extremely high carbohydrate content were given in Section 4.2.6.9.

Released protein and carbohydrate contents and bacteria LRV of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in various solution matrixes were given in Figure 4.167. Released protein and carbohydrate contents were calculated by subtracting initial concentrations from t_{irr} =60 min ones. The highest released protein (15.4 mg/L) and carbohydrate contents (81.2 mg/L), and the lowest bacteria LRV (2.23) were obtained in the presence of "HA and WM" solution. While both

WM and HA solutions had similar *E. coli* LRV (3.65 for *E. coli* in WM solution; 3.71 for *E. coli* in HA solution), released carbohydrate content of *E. coli* in HA solution (0.653 mg/L) was detected as significantly lower than WM solution (45.2 mg/L); therefore, it could be mentioned that degradation of released carbohydrate content occurred in the presence of HA solution. Beside high *E. coli* LRV detected from HA solution as 3.71, degradation of released carbohydrate content could be indicated that 0.50% Fe-SynTiO₂ displayed both high bacteria inactivation efficiency and organic matter degradation at the same time. *E. coli* LRV were plotted against released protein content and a strong correlation as R^2 =0.89 was obtained, however released carbohydrate contents and *E. coli* LRV showed a moderate correlation as R^2 =0.70.



Figure 4.167. Released protein and carbohydrate contents, and bacteria LRV of *E. coli* inactivation upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment in various solution matrixes.

Upon 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in different solution matrixes, the attained protein and carbohydrate contents, FI_{sync280}, UV₂₈₀ and NPOC data under initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.168. In the presence of HA solution, high UV₂₈₀ values were obtained similar to previous solar photolytic (Figure 4.24) and photocatalytic treatments (Figures 4.48, 4.72, 4.96, 4.120, and 4.143). Moreover, the presence of HA solution caused higher protein contents for all samples. FI_{sync280} data showed a significant increase in both IsoT and WM solution for initial, t=0, and t_{irr}=60 conditions. On the other hand, initial FI_{sync280} data of HA solution was detected as ≤ 0.001 . A strong correlation as R² > 0.93 was detected between protein concentrations and FI_{sync280} of initial and t=0 samples, whereas t_{irr}=60 min showed no correlation (9.15 mg/L for initial; 11.1 mg/L for t=0; 14.9 mg/L for t_{irr}=60 min). Treatments conducted in IsoT and WM solutions displayed similar patterns for carbohydrate and NPOC contents, however higher concentration were detected for WM solution.



Figure 4.168. Initial, t=0, and t_{irr} =60 min protein and carbohydrate contents, FI_{sync280}, UV₂₈₀, NPOC, and bacteria enumeration results of 0.50% Fe-SynTiO₂ solar photocatalytic treatment of *E. coli* in various aqueous matrixes.

Except significantly different carbohydrate content of WM (47.0 mg/L) and sole HA (4.14 mg/L) solutions at irradiation period of 60 min condition, they displayed resembling NPOC contents (13.9 mg/L for *E. coli* in WM solution; 14.9 mg/L for *E. coli* in HA solution) and bacteria counts (1.33E+02 CFU/mL for *E. coli* in WM solution; 1.27E+02 CFU/mL for *E. coli* in HA solution). It could be concluded that in WM solution, only released organic content from bacteria was the source of NPOC, whereas both released organic material from bacteria and present HA content contributed to measured NPOC for sole HA solution condition. Initial carbohydrate and NPOC contents showed a rough correlation for all reaction matrixes as R^2 =0.66; however, t=0 conditions showed a strong correlations, t_{irr}=60 min conditions displayed a strong correlation as R^2 =0.93, excluding the presence of IsoT solution.

4.3.1. Comparative Efficiency Evaluation of Different Photocatalyst Specimens on *E. coli* Inactivation in IsoT Solution

In the presence of sole IsoT solution, irradiation time dependent solar photocatalytic *E. coli* inactivation profiles and first order kinetic parameters upon use of undoped and Fe-doped photocatalyst specimens were compiled from respective figures and tables and presented in Figure 4.169 and Table 4.65 in a comparative manner.



Figure 4.169. Irradiation time dependent *E. coli* inactivation in IsoT solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens.

Table 4.65. Solar photocatalytic treatment of *E. coli* in IsoT solution performed in the presence of undoped and Fe-doped TiO₂ specimens: Inactivation parameters.

<i>E. coli</i> in IsoT solution solar photocatalytic treatment	First order kinetic model				Deduction 0/
	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min	LKV	Reduction, 70
TiO ₂	0.248	2.79	1.97E+05	5.90	99.999
SynTiO ₂	0.254	2.73	1.11E+05	5.64	99.999
0.25% Fe-TiO ₂	0.174	3.98	2.05E+05	4.28	99.99
0.50% Fe-TiO ₂	0.0839	8.26	5.19E+04	2.44	99
0.25% Fe-SynTiO ₂	0.116	5.97	4.67E+04	2.86	99.7
0.50% Fe-SynTiO ₂	0.158	4.39	8.56E+04	4.15	99.99

Irradiation time dependent continuous reduction of *E. coli* in IsoT solution was achieved under all solar photocatalytic treatment conditions using all TiO₂ specimens. The effect of each photocatalyst could be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in an increasing order as; 0.50% Fe-TiO₂ < 0.25% Fe-SynTiO₂ < 0.50% Fe-SynTiO₂ < 0.25% Fe-TiO₂ < TiO₂ < SynTiO₂. In general, bacteria reduction percentages obtained from some photocatalysts showed the same efficiencies in a decreasing order as; TiO₂ = SynTiO₂ > 0.25% Fe-TiO₂ = 0.50% Fe-SynTiO₂ > 0.25% Fe-SynTiO₂ > 0.50% Fe-TiO₂. Besides the similar bacteria reductions of some photocatalysts, LRV of *E. coli* displayed differences and could be expressed in an increasing order as; 0.50% Fe-TiO₂ < 0.25% Fe-SynTiO₂ < 0.50% Fe-SynTiO₂ < 0.25% Fe-TiO₂ < SynTiO₂ < TiO₂. Except the change observed for bare TiO₂ and SynTiO₂ specimens, Fe-doped photocatalyst specimens displayed the same order for both bacteria LRV and inactivation reaction rate constants. The most effective photocatalyst was detected as TiO₂ while the least effective one 0.50% Fe-TiO₂. The highest bacteria LRV of bare TiO₂ could be due to its high BET surface area as 57.6 m²/g (Table C.2) (Birben et al., 2015).

It could be indicated that Fe-doping in differing concentrations showed no significant difference on the efficiency of bacteria inactivation, whereas the most effective Fe-doped TiO₂ specimen was 0.25% Fe-TiO₂ in IsoT solution. Yalcin and colleagues reported that 0.25% Fe-TiO₂ displayed the highest degradation rate among the other dopant concentrations upon use of 4-nitrophenol as a model organic compound (Yalcin et al., 2010). The reason for higher inactivation rate of 0.25% Fe-TiO₂ could be related to the increased amount of HO[•] formation due to generated vacancies on TiO₂ surface by Fe³⁺ doping process. Formed oxygen vacancies on the photocatalyst surface could adsorb more water molecules and during irradiation, high amount of HO[•] radicals as the main ROS could from via radical reactions as explained by basic photocatalysis mechanism in Section 2.3.1 (Equations 2.1-2.8).

On the other hand, 0.25% Fe-TiO₂ exhibited the lowest crystallite particle size among other Fedoped photocatalysts as 16.4 nm which could generate various agglomerate sizes resulted in higher photocatalytic activity (Table C.1.) (Yalcin et al., 2010). Moreover, 0.25% Fe-doped TiO₂ specimen displayed heterogenous aggregate formations, smaller crystallite particles containing straight edges and sharp corners, which indicates the presence of large variety of shape and distribution causing high grade of porosity, therefore; it could be indicated that previously reported crystallite particle size properties of 0.25% Fe-doped TiO₂ could cause mechanic inactivation effect on *E. coli* (Yalcin et al., 2010). The second effective Fe-doped photocatalyst was detected as 0.50% Fe-SynTiO₂, exhibiting comparatively less agglomeration behavior (Turkten and Cinar, 2019). Agglomeration could lead to altered surface attractions prevailing in between *E. coli* and photocatalyst specimen resulting in attachment of particles onto comparatively higher cell surface area (6.0 μ m²) to be exposed to irradiation. Under these conditions, light harvesting capacity of the 0.50% Fe-SynTiO₂ could be affected leading to lower efficiency on the ROS formation.

4.3.1.1. Comparative mechanistic evaluation of *E. coli* inactivation in IsoT solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens. As inactivation products of bare and Fe-doped TiO₂ solar photocatalytic treatments of *E. coli* in the presence of IsoT solution, total K, protein, and carbohydrate concentrations of initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.170.



Figure 4.170. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents of *E. coli* inactivation in IsoT solution upon solar photocatalytic treatments performed in the presence of bare and Fe-doped TiO₂ specimens.

From a general perspective, all total K contents displayed variations in the range of 0.79-1.94 mg/L. Instantaneous adsorptive removal of total K at t=0 condition expressed changes in between 7.4-56 %. Maximum adsorptive removal of total K was detected upon use of 0.25% Fe-TiO₂ however, 0.50% Fe-TiO₂ displayed the lowest initial adsorptive removal at t=0 condition. Moreover, varying trend of total K contents could be related most probably to the passive and active transport mechanisms of *E. coli* as well as by attractive surface interactions of the charged sites.

Protein contents displayed variations in the range of 0.683-6.44 mg/L. Under all experimental conditions, protein concentrations increased at the end of t_{irr} =60 min irradiation, however maximum protein release was detected upon use of 0.25% Fe-TiO₂ as 6.44 mg/L. The presence of proteinaceous organic compounds was very clearly presented by respective UV-vis absorbance (Figures 4.74 and 4.76) synchronous scan fluorescence spectral features (Figure 4.75) as well as EEM fluorescence contour plots (Figures 4.78).

Under all conditions, carbohydrate contents displayed variations in the range of 0.891-78.0 mg/L. All concentrations showed an increase in carbohydrate contents at irradiation period of 60 min however maximum carbohydrate release was detected as 78.0 mg/L in case that 0.25% Fe-SynTiO₂ was utilized as the photocatalyst specimen.

As presented, the most efficient photocatalyst specimen in IsoT solution could be visualized as undoped TiO₂ specimen expressing the effective degradation mechanism upon solar photocatalysis as was also represented by NPOC contents (Figure 4.28, Table 4.12). On the other hand, it should be noted that, related to measured high carbohydrate contents at t_{irr} =60 with the use of SynTiO₂ (45.0 mg/L) and 0.25% Fe-SynTiO₂ (78.0 mg/L), and possible experimental errors due to the present interfering factors which widely explained in Section 3.2.3.3, comparative evaluation of photocatalyst specimens according to carbohydrate contents could cause erroneous conclusions.

4.3.2. Comparative Efficiency Evaluation of Different Photocatalyst Specimens on *E. coli* Inactivation in WM Solution

In the presence of sole WM solution, irradiation time dependent solar photocatalytic *E. coli* inactivation profiles and first order kinetic parameters upon use of undoped and Fe-doped photocatalyst specimens were compiled from respective figures and tables and presented in Figure 4.171 and Table 4.66 in a comparative manner. Solar irradiation time dependent continuous reduction of *E. coli* in WM solution was displayed upon solar photocatalysis using both undoped and Fe-doped

TiO₂ specimens. As could be visualized from logarithmic decay profiles, *E. coli* cells were effectively inactivated upon solar photocatalytic treatment processes using the specified specimens either as undoped or Fe doped forms in WM solution. Although variations were considerably evident in between the efficiencies of the employed photocatalyst specimens, upon irradiation period of 60 min, 2.95E+03 CFU/mL *E. coli* cells were still present upon use of 0.25% Fe-TiO₂ whereas only 2.85E+01 CFU/mL *E. coli* cells were present upon use of SynTiO₂. Therefore, the effect of reaction medium could be successfully expressed in terms of inactivation reaction rate constant (k, min⁻¹) in an increasing order as; 0.25% Fe-TiO₂ < TiO₂ < 0.50% Fe-TiO₂ < 0.25% Fe-SynTiO₂ ≤ 0.50% Fe-SynTiO₂ < SynTiO₂ (Table 4.66).



Figure 4.171. Irradiation time dependent *E. coli* inactivation in WM solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens.

Table 4.66. Solar photocatalytic treatment of *E. coli* in WM solution performed in the presence of undoped and Fe-doped TiO₂ specimens: Inactivation parameters.

<i>E. coli</i> in WM solution solar photocatalytic treatment	First order kinetic model				Deduction 0/
	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min	LKV	Reduction, 70
TiO ₂	0.132	5.25	9.44E+04	3.73	99.9
SynTiO ₂	0.180	3.85	1.14E+05	4.35	99.99
0.25% Fe-TiO ₂	0.0891	7.78	8.17E+04	2.49	99
0.50% Fe-TiO ₂	0.135	5.13	9.06E+04	3.49	99.9
0.25% Fe-SynTiO ₂	0.146	4.75	3.26E+04	3.75	99.9
0.50% Fe-SynTiO ₂	0.146	4.75	8.69E+04	3.65	99.9

The effect of photocatalyst specimens on *E. coli* inactivation in WM solution could also be displayed in terms of reduction percentage of *E. coli* in an increasing order as; 0.25% Fe-TiO₂ < 0.50% Fe-TiO₂ = 0.50% Fe-SynTiO₂ = TiO₂ = 0.25% Fe-SynTiO₂ < SynTiO₂. However, although *E coli* reduction percentages (e.g., 99.9 %) were the same, respective bacteria LRV displayed differences in the range of 3.49-3.75. LRV of *E. coli* inactivation followed an increasing order as; 0.25% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-SynTiO₂ < TiO₂ < 0.25% Fe-SynTiO₂ < SynTiO₂. It could be indicated that, except 0.25% Fe-TiO₂ specimen, other Fe-doped photocatalysts showed no significant efficiency difference. The most effective photocatalyst upon *E. coli* inactivation in WM solution was observed as SynTiO₂ while the least effective one 0.25% Fe-TiO₂ specimen.

<u>4.3.2.1.</u> Comparative mechanistic evaluation of *E. coli* inactivation in WM upon solar photocatalytic treatment with undoped and Fe-doped TiO₂ specimens. As solar photocatalytic inactivation products of bare and Fe-doped TiO₂ solar photocatalytic treatments of *E. coli* in the presence of WM components, total K, protein, and carbohydrate contents under initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.172.

Total K contents displayed variations in the range of 3.86-4.38 mg/L. Except 0.25% Fe-TiO2, all photocatalyst specimens displayed an immediate adsorptive removal of total K content at t=0 condition in the range of 1.7-9.5%. Maximum adsorptive removal % of total K content was attained in the case of 0.50% Fe-SynTiO₂. Moreover, all experiment conditions showed decreased total K concentrations at the end of 60 min irradiation period. Depending on the resultant cell counts depicted in respective Figure 4.171, the role of K⁺ ion transportation mechanisms in and out of the cell wall via passive and active transport should also be considered under all conditions. It should be noted that excessive amount of K ions present in WM solution could be used by bacteria to compensate released K⁺ ion content and protect bacteria from destruction by retarding of cell wall damage.

Furthermore, due to the presence of common anions and cations in WM solution, various electrostatic attractive and repulsive forces between charged species should take place resulting in varying surface interactions both onto photocatalyst specimens and bacteria surface. The effect of pH_{zpc} of photocatalyst specimens (TiO₂: 6.25; SynTiO₂: 6.40; 0.25% Fe-TiO₂: 4.80) played the dominant role specifically on t=0 conditions and during solar photocatalytic treatment under irradiation conditions covering the formation of charged cell fractions (Birben et al., 2015, 2017b).

From a general perspective, protein contents displayed variations in the range of 0.710-14.8 mg/L. All protein concentrations showed an increase following $t_{irr}=60$ min however the highest

protein content was detected as 14.8 mg/L upon use of SynTiO₂ specimen. Under these conditions, both UV-vis absorbance (Figures 4.55 and 4.57) and synchronous scan fluorescence spectral features (Figure 4.56) as well as EEM fluorescence contour plots (Figure 4.59) proved the existence of protein like components in reaction medium.



Figure 4.172. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents of *E. coli* inactivation in WM solution upon solar photocatalytic treatments performed in the presence of bare and Fe-doped TiO₂ specimens.

From a general perspective, carbohydrate contents displayed variations in the range of 0.618-74.7 mg/L. However, upon use of undoped and Fe-doped TiO₂ specimens almost all carbohydrate contents were quite low as < 4 mg/L for initial and t=0 conditions, expressing insignificant role of Fe doping on TiO₂. Since carbohydrate contents could not be followed by spectroscopic techniques, it could only be related to the selectivity of photocatalysis towards the organic fractions exhibiting carbohydrate structures mainly composed of hydrophilic groups. Deprotonation of these groups could

be neutralized by counter ions e.g., K⁺ ions formed through cell lysis or by cations present as WM constituent (Bythell et al., 2018). Although an analogous approach could also be related to the behavior of SynTiO₂ (< 2 mg/L at t_{irr}=60 min), Fe-doping significantly affected the reaction mechanism excluding the degradation of carbohydrate components. Increasing Fe dopant concentration of SynTiO₂ from 0.25% to 0.50% did not reflect as a dopant concentration role, therefore lower carbohydrate content was found as 47 mg/L. The reason could be attributed to the surface morphological properties of Fe-doped Syn TiO₂ specimens rather than the selectivity or efficiency of photocatalysis. Similar trend was also observed upon use undoped and 0.25% Fe-doped TiO₂ specimens during solar photocatalytic inactivation of E. coli in IsoT solution revealing carbohydrate contents as $\sim < 5$ mg/L (Figure 4.170). Further increase in Fe doping of TiO₂ resulted in substantial increase in carbohydrate content as 25 mg/L. The reason could only be related to the sole presence of Cl⁻ anion in solution exerting considerably less ion effect. Contrary to these findings, SynTiO₂ either as undoped or doped forms expressed varying carbohydrate contents under all conditions proceeding through different reaction mechanism. Interestingly, upon use of 0.25% Fe-SynTiO₂ in the presence of either IsoT solution (78.0 mg/L) or WM solution (74.7 mg/L) almost similar carbohydrate contents were still present in reaction medium (Figures 4.170 and 4.172).

4.3.3. Comparative Efficiency Evaluation of Different Photocatalyst Specimens on *E. coli* Inactivation in HA Solution

In the presence of sole HA solution, irradiation time dependent solar photocatalytic *E. coli* inactivation profiles and first order kinetic parameters upon use of undoped and Fe-doped photocatalys specimens were compiled from respective figures and tables and presented in Figure 4.173 and Table 4.67 in a comparative manner.

Irradiation time dependent continuous reduction of *E. coli* in HA solution was achieved under all solar photocatalytic treatment conditions using undoped and Fe-doped TiO₂ specimens. Although differences were considerably evident in between the efficiencies of the employed photocatalyst specimens, upon $t_{irr}=60 \text{ min}$, 3.85E+04 CFU/mL E. coli cells were still present upon use of bare TiO₂ whereas only 3.35E+01 CFU/mL E. coli cells were present upon use of undoped SynTiO₂. Therefore, the effect of photocatalyst specimens could be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in an increasing order as; TiO₂ < 0.25% Fe-TiO₂ < 0.50% Fe-TiO₂ < 0.50% Fe-SynTiO₂ < SynTiO₂ < 0.25% SynTiO₂. The effect of different photocatalysts on *E. coli* inactivation in the presence of sole HA solution could also be displayed in terms of *E. coli* reduction percentage in a decreasing order as; SynTiO₂ = 0.50% Fe-TiO₂ = 0.25% Fe-SynTiO₂ = 0.50% Fe-SynTiO₂ > 0.25% Fe-TiO₂ > TiO₂. Besides the same bacteria reduction percentage of some photocatalysts (e.g., 99.9 %), LRV of *E. coli* displayed changes in between 3.36-3.82. *E. coli* LRV of all conditions could be displayed in an increasing order as; TiO₂ < 0.25\% Fe-TiO₂ < 0.50\% Fe-TiO₂ < SynTiO₂ < 0.50% Fe-SynTiO₂ < 0.25\% Fe-SynTiO₂.



Figure 4.173. Irradiation time dependent *E. coli* inactivation in HA solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens.

Table 4.67. Solar photocatalytic treatment of *E. coli* in HA solution performed in the presence of undoped and Fe-doped TiO₂ specimens: Inactivation parameters.

<i>E. coli</i> in HA solution solar photocatalytic treatment	First order kinetic model				De la stiene 0/
	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min	LKV	Keduction, 70
TiO ₂	0.0672	10.3	7.26E+04	1.45	90
SynTiO ₂	0.150	4.62	2.54E+04	3.70	99.9
0.25% Fe-TiO ₂	0.0774	8.95	4.74E+04	2.09	99
0.50% Fe-TiO ₂	0.125	5.54	7.81E+04	3.36	99.9
0.25% Fe-SynTiO ₂	0.152	4.56	3.98E+04	3.82	99.9
0.50% Fe-SynTiO ₂	0.148	4.68	9.53E+04	3.71	99.9

The most effective photocatalyst in the presence of HA solution was detected as 0.25% Fe-SynTiO₂ and the least effective one as bare TiO₂ specimen. Related to detected inactivation rate constants and LRV upon use of various photocatalyst specimens, it could be concluded that SynTiO₂ and its Fe-doped species displayed selectivity towards *E. coli* however TiO₂ and its Fe-doped species expressed selectivity against humic matter in solution. The reason could be attributed that the different crystallite particle sizes of photocatalyst specimens (Table C.1). Besides lower crystallite particle sizes of bare TiO₂ (22.3 nm) and Fe-doped TiO₂ specimens (16.4 nm for 0.25% Fe-TiO₂; 19.9 nm for 0.50% Fe-TiO₂), bare SynTiO₂ (33.1 nm) and Fe-doped SynTiO₂ specimens (20.7 nm for 0.25% Fe-SynTiO₂; 23.7 nm for 0.50% Fe-SynTiO₂) displayed higher crystallite particle sizes (Turkten and Cinar, 2019; Turkten, 2016; Turkten et al., 2019; Yalcin et al., 2010). It could be indicated that photocatalyst specimens having lower crystallite particle sizes, namely TiO₂ species, display selectivity toward the removal of organic matter however higher crystallite particle sized SynTiO₂ species showed selectivity against bacteria inactivation.

Furthermore, findings of Birben and colleagues indicating that 0.25% Fe-TiO₂ showed a retardation effect on HA removal rates compared to undoped TiO₂, supporting the order of 0.25% Fe-TiO₂ and bare TiO₂ in the aspect of inactivation rate constants and bacteria LRV. It could be indicated that 0.25% Fe-TiO₂ has slightly more selectivity towards bacteria than TiO₂. Researchers reported that the surface interactions between HA and surface of Fe-doped photocatalysts could be the cause of slower degradation rates of HA upon solar photocatalytic treatment (Birben et al., 2017b).

4.3.3.1. Comparative mechanistic evaluation of *E. coli* inactivation in HA solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO_2 specimens. As inactivation products of bare and Fe-doped TiO_2 solar photocatalytic treatments of *E. coli* in the presence of sole HA solution, total K, protein, and carbohydrate contents of initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.170.

From a general view, total K contents expressed changes in the range of 0.44-0.73 mg/L. In the presence of HA solution, total K contents displayed lower presence from IsoT solution conditions (Figure 4.170). Different from IsoT and WM solution conditions, in the presence of HA solution, immediate adsorptive removal of total K content at t=0 condition was only calculated upon use of bare TiO₂ and 0.25% Fe-TiO₂ specimens as 7.5 and 25 %, respectively. Moreover, except 0.25% Fe-TiO₂, all other conditions, showed increased total K concentrations at the end of 60 min irradiation period, which was a pattern that did not detected in the presence of IsoT and WM solutions (Figures 4.170 and 4.172).

Under all conditions, protein contents expressed changes in the range of 3.75-16.2 mg/L. The use of bare TiO₂ as a photocatalyst resulted in a degradation in protein content at t_{irr}=60 min (13.4-3.75 mg/L). It could be indicated that, expressed protein degradation could be associated with

obtained lower bacteria inactivation rate of TiO_2 and could also support the suggestion about selectivity of TiO_2 against organic content rather than bacteria. On the other hand, solar photocatalytic treatments conducted in the presence of bare SynTiO₂ and all Fe-doped photocatalyst specimens displayed increased protein contents at the end of 60 min irradiation period, being the highest for 0.50% Fe-SynTiO₂ condition as 16.2 mg/L. Under these conditions, despite the masking effect of humic-like matter on protein-like ones, both UV-vis absorbance (Figures 4.156 and 4.158) and synchronous scan fluorescence spectral features (Figure 4.157) as well as EEM fluorescence contour plots (Figure 4.160) proved the existence of protein like components in reaction medium.



Figure 4.174. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents of *E. coli* inactivation in HA solution upon solar photocatalytic treatments performed in the presence of bare and Fe-doped TiO₂ specimens.

Carbohydrate contents displayed variations in the range of 2.24-77.9 mg/L. However, upon use of undoped and Fe-doped TiO₂ specimens almost all carbohydrate contents were quite low as < 5 mg/L for initial and t=0 conditions, expressing insignificant role of Fe doping on TiO₂. Similar trend

was also observed for initial and t=0 conditions upon use undoped and Fe-doped TiO₂ photocatalyst specimens during solar photocatalytic inactivation of *E. coli* in IsoT and WM solutions revealing carbohydrate contents as < 4 mg/L (Figure 4.170).

On the other hand, similar to degradation of protein content in the presence of undoped TiO_2 after irradiation period of 60 min, degraded carbohydrate content (2.77-2.24 mg/L) was detected upon use of TiO_2 . All other conditions showed an increase in carbohydrate contents at the end of 60 min solar light exposure period however maximum carbohydrate content was detected as 77.9 mg/L in case that 0.50% Fe-TiO₂ was used as the photocatalyst specimen.

4.3.4. Comparative Efficiency Evaluation of Different Photocatalyst Specimens on *E. coli* Inactivation in "HA and WM" Solution

In the presence of "HA and WM" solution, irradiation time dependent solar photocatalytic *E. coli* inactivation profiles and first order kinetic parameters upon use of undoped and Fe-doped photocatalyst specimens were compiled from respective figures and tables and presented in Figure 4.175 and Table 4.68 in a comparative manner.

Irradiation time dependent continuous reduction of *E. coli* in "HA and WM" solution was achieved under all solar photocatalytic treatment conditions using undoped and Fe-doped TiO₂ specimens. However, after irradiation time of 40 min, TiO₂, SynTiO₂, and 0.25% Fe-SynTiO₂ displayed a retardation in *E. coli* reduction (Figure 4.175). Although variations were considerably evident in between the efficiencies of the employed photocatalyst specimens, upon irradiation period of 60 min, 5.50E+04 CFU/mL *E. coli* cells were still present upon use of bare TiO₂ whereas only 4.67E+02 CFU/mL *E. coli* cells were present upon use of SynTiO₂. Based on these results, the effect of photocatalysts could be expressed in terms of inactivation reaction rate constant (k, min⁻¹) in an increasing order as; TiO₂ < 0.50% Fe-TiO₂ < 0.25% Fe-TiO₂ < 0.50% Fe-SynTiO₂ < 0.25% SynTiO₂

The effects of photocatalyst specimens on bacteria inactivation in "HA and WM" solution could also be expressed in terms of reduction percentage of *E. coli* in a decreasing order as; SynTiO₂ > 0.25% Fe-SynTiO₂ > 0.50% Fe-SynTiO₂ = 0.25% Fe-TiO₂ > 0.50% Fe-TiO₂ > TiO₂. However, although bacteria reduction percentages (e.g., 99 %) were the same, respective bacteria LRV displayed variations in between 2.14-2.23. LRV of *E. coli* inactivation in "HA and WM" solution followed an increasing order as; TiO₂ < 0.50% Fe-TiO₂ < 0.25% Fe-TiO₂ < 0.50% Fe-SynTiO₂ < 0.25% Fe-SynTiO₂ < SynTiO₂. It could be indicated that, 0.25% Fe-SynTiO₂ expressed higher bacteria inactivation efficiency among other Fe-doped photocatalyst specimens. On the other hand, the most effective photocatalyst upon *E. coli* inactivation in "HA and WM" solution was observed as SynTiO₂ while the least effective one bare TiO₂ specimen.



Figure 4.175. Irradiation time dependent *E. coli* inactivation in "HA and WM" solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens.

Table 4.68. Solar photocatalytic treatment of *E. coli* in "HA and WM" solution performed in the presence of undoped and Fe-doped TiO₂ specimens: Inactivation parameters.

<i>E. coli</i> in "HA and	First	LDV			
photocatalytic treatment	k, min ⁻¹	t _{1/2} , min	R, CFU/mL min	LRV	Reduction, %
TiO ₂	0.0488	14.2	3.22E+04	1.08	90
SynTiO ₂	0.117	5.92	6.21E+04	3.06	99.9
0.25% Fe-TiO ₂	0.0834	8.31	7.72E+04	2.14	99
0.50% Fe-TiO ₂	0.0734	9.44	3.92E+04	1.84	96.8
0.25% Fe-SynTiO ₂	0.0943	7.35	7.38E+04	2.50	99.7
0.50% Fe-SynTiO ₂	0.0850	8.15	4.88E+04	2.23	99

From a general perspective, related to inactivation rate constants and LRV of *E. coli* upon use of different photocatalyst specimens, it could be concluded that $SynTiO_2$ and Fe-doped $SynTiO_2$ specimens, having higher crystallite particle sizes, displayed selectivity towards *E. coli* however TiO_2 and Fe-doped TiO_2 specimens, having lower crystallite particle sizes, expressed selectivity against

humic matter present in reaction solution (Table C.1.). Similar trend was also observed from sole HA solution conditions.

Moreover, Birben and colleagues reported that 0.25% Fe-TiO₂ showed a retardation effect on HA removal rates compared to bare TiO₂, which complies with higher bacteria inactivation rate constant and LRV obtained in case that 0.25% Fe-TiO₂ was used as the photocatalyst specimen (Birben et al., 2017b). Therefore, it could be indicated that 0.25% Fe-TiO₂ has selectivity towards bacteria rather than HA content on contrary to bare TiO₂ specimen.

Based on inactivation rate constants and LRV of *E. coli* inactivation in "HA and WM" solution, undoped and Fe-doped SynTiO₂ specimens displayed quite close results with each other however bare SynTiO₂ expressed higher *E. coli* LRV and bacteria inactivation rate constant than 0.50% Fe-SynTiO₂ specimen. However, Turkten and Cinar reported that 0.50% Fe-SynTiO₂ displayed a slight agglomeration and higher degradation percentage of 4-nitrophenol as a model organic compound than SynTiO₂ and 0.50% Fe-SynTiO₂ (Turkten and Cinar, 2019). Therefore, it could be concluded that 0.50% Fe-SynTiO₂ most probably tend to display selectivity towards organic matter degradation rather than bacteria inactivation on contrary to bare SynTiO₂ specimen.

<u>4.3.4.1.</u> Comparative mechanistic evaluation of *E. coli* inactivation in "HA and WM" solution upon solar photocatalytic treatments performed in the presence of undoped and Fe-doped TiO₂ specimens. As solar photocatalytic inactivation products of bare and Fe-doped TiO₂ solar photocatalytic treatments of *E. coli* in "HA and WM" solution components, total K, protein, and carbohydrate contents under initial, t=0, and t_{irr}=60 min conditions were given in Figure 4.176.

Under all conditions, total K contents displayed variations in the range of 4.07-4.61 mg/L. Due to the presence of WM solution as a reaction matrix constituent, total K contents expressed higher presence than IsoT (Figure 4.170) and sole HA solution (Figure 4.174) conditions as expected. All photocatalyst specimens displayed immediate adsorptive removal of total K the range of 0.44-4.1 % at t=0 condition. Moreover, all conditions showed decreased total K concentrations at the end of 60 min irradiation period.

Protein contents expressed changes in between 4.35-22.8 mg/L under all conditions. In the presence of bare TiO₂, protein content displayed a decreasing trend at t_{irr}=60 min (3.93-1.48 mg/L) on contradiction to the increasing trend obtained upon use of other photocatalyst specimens. Degradation of proteinaceous matter could be associated with obtained lower bacteria inactivation

rates in case that TiO_2 used as bare photocatalyst and could support the suggestion about selectivity of TiO_2 against organic content rather than bacteria. Similar protein degradation and release patterns were also detected solar photocatalytic treatments conducted in sole HA solution conditions (Figure 4.174).



Figure 4.176. Initial, t=0, and t_{irr} =60 min total K, protein, and carbohydrate contents of *E. coli* inactivation in "HA and WM" solution upon solar photocatalytic treatments performed in the presence of bare and Fe-doped TiO₂ specimens.

Upon with the use of bare SynTiO₂ and all Fe-doped photocatalyst specimens, protein contents showed an increase at the end of 60 min irradiation period and maximum protein concentration was detected as 22.8 mg/L in the presence of 0.50% Fe-SynTiO₂. Under these conditions, despite the masking effect of humic-like fluorophores on protein-like ones, the presence of proteinaceous organic compounds was clearly presented by respective UV-vis absorbance (Figures 4.161 and 163), synchronous scan fluorescence spectral features (Figure 4.162) as well as EEM fluorescence contour

plots (Figures 4.165). Interestingly, solar photocatalytic treatments upon use of 0.50% Fe-SynTiO₂ in the presence of either sole HA solution (16.2 mg/L) or "HA and WM" solution (22.8 mg/L) high protein contents were still present in reaction medium (Figures 4.174 and 4.176).

From a general view, carbohydrate contents displayed variations in the range of 1.48-84.2 mg/L. Upon use of undoped and Fe-doped TiO₂ specimens all carbohydrate concentrations were quite low as < 6.5 mg/L for initial and t=0 conditions, expressing insignificant role of Fe doping on TiO₂. On the other hand, bare TiO₂ specimen displayed a degradation effect on carbohydrate content at t_{irr} =60 min (3.93-1.48 mg/L) similar to detected decrease in protein content. Similar trend was also detected upon use of undoped TiO₂ in sole HA solution condition (Figure 4.174). All other photocatalyst specimens showed increased carbohydrate concentrations at the end of 60 min solar light irradiation period and maximum carbohydrate content was detected as 84.2 mg/L in the case that 0.50% Fe-SynTiO₂.

4.4. Endotoxin Release Upon Solar Photolytic and Photocatalytic Inactivation of *E. coli* in Various Aqueous Solutions

Endotoxins are noxious substances present outside of the cell wall of Gram-negative bacteria. LPS which are a component of outer membrane of Gram-negative bacteria e.g., *E. coli*, consists of three parts as: O polysaccharide, core polysaccharide, and lipid A (Figure 4.177). The lipid A portion of LPS is the endotoxin of *E. coli*, therefore; it should be noted that endotoxins are not metabolic products, they are a natural part of the cell wall. Endotoxin release could occur when bacteria die, cell wall undergo to lysis and during the bacterial multiplication (Prescott et al., 1993; Tortora et al., 2018). Moreover, it should be indicated that endotoxins are not necessarily specific to infectious *E. coli* serotype O157:H7 due to presence of endotoxins in the outer membrane structure of all Gramnegative bacteria.

Major health related impacts could be expressed as that endotoxins have a great spectrum of biological activities on mammals and some of the main ones are pyrogenicity, leukopenia and septic shock. Basically, pyrogens are fever inducing agents of Gram-negative bacteria represented by endotoxins and pyrogenicity is probably the first recognized activity of endotoxins (Galanos, 1998).

During solar photolytic and photocatalytic inactivation processed, endotoxins could be released as inactivation by-products of bacteria. In this study, potential release of *E. coli* endotoxins was followed under different irradiation period conditions as well as control conditions as sole HA solution, IsoT solution, WM solution, and distilled-deionized water. Limiting value of LAL analysis kit was determined as 0.125 EU/mL according to pure dialysis water having pyrogen-free characteristics (Glorieux et al., 2012).



Figure 4.177. Schematic diagram of LPS structure and positioning on Gram-negative bacteria cell wall (Cardoso et al., 2006).

All selected solutions were subjected to LAL analysis and displayed endotoxin content as > 0.125 EU/mL including control conditions. Due to the detection of endotoxin content in sole HA, IsoT and WM solution conditions in the absence of E. coli, it could be considered that the probability of false positive determinations. Moreover, even distilled-deionized water expressed endotoxin content more than 0.125 EU/mL, therefore; positive endotoxin contents of sole reaction matrixes could be due to the lack of pyrogen-free water conditions rather than false positive endotoxin determinations. On the other hand, it should be noted that sterile and pyrogen-free indicates different conditions, sterile means absence of any viable microorganism while pyrogen-free means nonexistence of fever causing substances. Consequently, it could be indicated that endotoxin analysis should be cautiously interpreted in the absence of pyrogen-free water conditions. However, degradation of endotoxin content originated from E. coli destruction under UV and solar irradiation conditions upon use of various photocatalyst specimens were reported in literature (Gadgil and Kodialbail, 2021; Sunada et al., 1998; Sreeja and Shetty, 2016, 2017). It should be emphasized that by-products released upon cell lysis e.g., carbohydrates were not considered in all of the aforementioned publications contrary to the findings reported in this study. Therefore, a comprehensive approach covering all aspects of cell destruction resulting in utmost removal of any live bacteria should be followed by characterization of complex organic matrix unless otherwise complete mineralization is achieved.

5. CONCLUSION

In this study, solar photolytic and photocatalytic inactivation of *E. coli* in various reaction matrixes was successfully achieved and characterized. Photocatalysis was performed under simulated solar irradiation conditions (wavelength range of λ =290 nm to 800 nm and light intensity, I₀=250 W/m²) using different TiO₂ based photocatalyst specimens i.e., TiO₂ (P-25), SynTiO₂ (sol-gel synthesized) and Fe-doped counterparts as 0.25% and 0.50%. HA solution was used to represent NOM in aqueous medium. Saline solution (IsoT) was considered as baseline medium and aqueous solution containing common anions and cations defined water matrix (WM) was also employed.

Most significantly, simultaneous reduction of *E. coli* cell counts and degradation of organic matter could be attained under different experimental conditions upon solar irradiation. A retardation effect in the presence of "HA and WM" solution as mimicking natural surface water conditions was observed upon all solar photolytic and photocatalytic conditions regardless of changing photocatalyst specimens either bare or Fe-doped types. However, the significant role of "HA and WM" solution occurring via both photo initiated self-reactions and adsorptive interactions should be cautiously interpreted.

Photocatalyst specimens displayed different effectiveness upon use of IsoT, WM, sole HA, and "HA and WM" solutions. Solar photocatalytic inactivation of E. coli in IsoT solution under the presence of bare TiO₂ specimen was effectively achieved leading LRV as 5.90 excluding all possible adverse effects on inactivation process. However, in the presence of "HA and WM" solution, undoped TiO₂ was observed as the least effective photocatalyst specimen expressing *E. coli* LRV as 1.08. Attained enhanced retardation of E. coli inactivation in "HA and WM" solution should be evaluated considering present binary and ternary interactions between reaction medium components. In the presence of common anion and cation containing WM solution, solar photocatalytic E. coli inactivation was achieved successfully upon use of bare SynTiO₂ expressing LRV as 4.35. Moreover, upon use of sole HA solution as a reaction medium, the most effective E. coli reduction was attained in the presence of 0.25% Fe-SynTiO₂ specimen expressing LRV as 3.82. On the other hand, undoped SynTiO₂ was observed as the most effective photocatalyst specimen on E. coli inactivation in "HA and WM" solution revealing bacteria LRV as 3.06. As an overall evaluation expressed by first order kinetic rate constants, except upon use of 0.25% Fe-TiO₂, all solar photolytic and photocatalytic treatments displayed the least effectiveness on *E. coli* reduction in "HA and WM" solution conditions. Therefore, it should be noted that, bacteria inactivation efficiencies of different photocatalyst specimens were directly related to present aqueous medium conditions.

Organic matter released and removed upon solar photolytic and photocatalytic inactivation of *E. coli* could be effectively characterized by UV-vis and fluorescence spectroscopic techniques and quantified by NPOC contents. Based on applied EEM fluorescence contour plots relating to *E. coli* inactivation upon solar irradiation, properties of in-situ produced organic matter could be deeply investigated. Application of EEM fluorescence spectroscopy displayed detailed information about qualification of organic inactivation products by distinguishing them from humic features originating from background reaction matrixes. Moreover, emergence of microbial by-products and protein-like fluorophores were evidently identified by EEM fluorescence features.

Bacteria inactivation products as total K⁺, protein and carbohydrate contents significantly revealed the destruction of *E. coli* cell wall. However, K⁺ ion release from damaged cells should be carefully distinguished from already present K⁺ ion content originating from WM condition especially while working with low initial bacteria counts. In the presence of high NOM contents, the accuracy of inactivation product concentrations (protein and carbohydrate) could considerably be affected via cage effect and/or encapsulation of lower molecular weight fractions by higher molecular weight fractions of humic substances due to re-conformational arrangements. On the other hand, it should be kept in mind that effect of filtration applied prior to analysis could cause elimination of some inactivation products due to prevailing electrostatic interactions between matrix constituents, therefore; remaining organic and inorganic substances in solution could not be regarded as representing exact content prior to filtration. It should be emphasized that membrane filtration did not exert any effect on *E. coli* cell counts thereby inactivation efficiencies due to the applied sampling procedure prior to filtration step.

Considering diverse physicochemical features of natural water systems containing NOM along with inorganic components, due to probable retardation effects on microorganism inactivation efficiencies of various photocatalyst specimens, direct solar light initiated reactions should be taken into consideration as well. Thus, simultaneously operating reactions ensued remarkable efficiencies on both *E. coli* cell count reduction as well as degradation of complex organic matter using bare TiO_2 and VLA TiO_2 specimens.

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APPENDIX A: COMPOSITION OF WATER MATRIX SOLUTION

Common anions and cations	Concentration, mg/L
Anions [*]	
Cl-	88.3
NO ₃ ⁻	6.11
SO4 ²⁻	59.5
$H_2PO_4^-$	3.23
Cations**	
Na ⁺	12.3
K^+	4.26
Ca ²⁺	61.0
Mg^{2+}	9.05
-1-	

Table A.1. Composition of WM solution.

Reference methodology: *Ion chromatography (IC), **Flame atomic absorption spectroscopy (FAAS).

APPENDIX B: SPECTROSCOPIC CHARACTERIZATION OF 100 kDa HUMIC ACID SOLUTION



Figure B.1. UV-vis absorption spectra of 100 kDa HA solution.



Figure B.2. Synchronous scan fluorescence spectra of 100 kDa HA solution.

Table B.1. Specified and specific UV-vis and fluorescence spectroscopic properties and NPOC of100 kDa HA solution.

	UV-vis	UV-vis Spectroscopic Parameters, cm ⁻¹			NPOC,	SUVA ₂₅₄ ,	FIsync470	SFI _{sync470}	FI
	Color ₄₃₆	UV ₃₆₅	UV ₂₈₀	UV ₂₅₄	mg/L	L/m mg	591101170	Synetro	
100 kDa HA Solution	0.0960	0.2100	0.5337	0.6295	4.96	12.7	67.8	13.7	1.03



Figure B.3. EEM fluorescence contour plot of 100 kDa HA solution.

APPENDIX C: CHARACTERIZATION OF PHOTOCATALYST SPECIMENS



Figure C.1. X-Ray Diffraction (XRD) diffractograms of undoped and Fe-doped photocatalyst specimens: a. TiO₂, b. 0.25% Fe-TiO₂, c. 0.50% Fe-TiO₂ (Yalcin et al., 2010),
d. SynTiO₂, e. 0.25% Fe-SynTiO₂, f. 0.50% Fe-SynTiO₂ (Turkten and Cinar, 2019).

Table C.1. Crystallite particle sizes, absorption wavelengths and band-gap energies (E_{bg}) of undoped and Fe-doped TiO₂ specimens.

Photocatalysts	Crystallite Particle Size, nm	E _{bg} , eV	Wavelength, nm	References
TiO ₂	22.3	3.18	390	Turkten et al., 2019
SynTiO ₂	33.1	2.91	410	Turkten et al., 2019
0.25% Fe-TiO ₂	16.4	2.55	486	Yalcin et al., 2010
0.50% Fe-TiO ₂	17.9	2.48	500	Yalcin et al., 2010
0.25% Fe-SynTiO ₂	20.7	2.82	440	Turkten, 2016
0.50% Fe-SynTiO ₂	23.7	2.67	466	Turkten and Cinar, 2019

Table C.2. BET surface areas of photocatalyst specimens.

Photocatalysts	BET surface area, m ² /g	References
TiO ₂	57.6	Birben et al., 2015
SynTiO ₂	50.3	Turkten et al., 2019
0.25% Fe-TiO ₂	33.0	Birben et al., 2017
0.25% Fe-SynTiO ₂	55.4	Turkten, 2016
0.50% Fe-SynTiO ₂	34.0	Turkten and Cinar, 2019



Figure C.2. SEM micrographs and Energy-Dispersive X-Ray (EDX) spectra of undoped and Fedoped photocatalyst specimens: a. TiO₂, b. 0.25% Fe-TiO₂ (Yalcin et al., 2010);

c. SynTiO₂, d. 0.50% Fe-SynTiO₂ (Turkten and Cinar, 2019).

APPENDIX D: CALIBRATION CURVE OF BOVINE SERUM ALBUMIN STANDARD



Figure D.1. Calibration curve of bovine serum albumin (BSA) standard.

$$y = 179.5x - 1.043$$

Where,

y = Concentration of total protein in solution, mg/L

x = Measured absorbance of the solution at λ =750 nm

APPENDIX E: CALIBRATION CURVE OF GLUCOSE STANDARD



Figure E.1. Calibration curve of glucose standard.

y = 96.02x + 1.889

Where,

y = Concentration of total carbohydrate in solution, mg/L

x = Measured absorbance of the solution at λ =490 nm