

**REMOVAL OF HAZARDOUS POLLUTANTS
BY IMPREGNATION AND PARAMETRIC
OPTIMIZATION OF ADSORBENTS**

By

Amjad Farooq

A thesis submitted to the faculty of Engineering in partial
fulfillment of the requirements for the degree of

Doctor of Philosophy



Department of Chemical Engineering
Pakistan Institute of Engineering & Applied Sciences,
Nilore, Islamabad, Pakistan
January, 2012



Allah does not impose on any soul more than He
has given it. Allah will provide ease after
difficulty.

(Qur'an, 65:7)

Department of Chemical Engineering
Pakistan Institute of Engineering and Applied Sciences (PIEAS)
Nilore, Islamabad 45650, Pakistan

Declaration

I declare that all material in this thesis which is not my own work has been identified and that no material has previously been submitted and approved for the award of a degree by this or any other university.

Signature:  _____

Author's Name: **Amjad Farooq**

It is certified that the work in this thesis is carried out and completed under my supervision.




Supervisor :

Dr. Naseem Irfan

DCE (DNE),

PIEAS, Islamabad

Co-Supervisor : 

Dr. Muhammad Mansha Chaudhry

DCS (DCHE),

PIEAS, Islamabad

I dedicate this thesis to

My wife

Acknowledgement

All Praise is for Allah Almighty, the Savior of the universe, the Merciful, the Beneficent, Who always led me through the thick and thin. Humblest gratitude is due to the Holy Prophet Hazrat Muhammad (Peace be upon him) and his companions.

I must first of all acknowledge the vision and the efforts of my very kind and affectionate supervisor, Prof. Dr. Naseem Irfan, who has always been much more than a mere teacher or supervisor. He has led me through the seemingly unbelievable paths to where I am today. He is fond of using all his mental and physical resources to the height of perfection and this is what inspires me about his personality. He never hesitates to help someone even if he has to sacrifice his own interest, time and money. I pray for his health and his family's well-being. May his dreams come true. Prof. Dr. Muhammad Mansha Chaudhry, my co-supervisor, also deserves heartiest thanks and appreciation. He taught me difficult concepts of chemistry and I always found him ready to help and guide me on issues other than research also. Both of my respectable supervisors deserve a special word of thanks for the efforts they made to review this thesis. Dr. Syed Moosa Hassany (late) guided me very well in early days of my research work. May his soul rest in peace.

During the course of this research, I have been to a few foreign laboratories, where I did some experiments as a part of my PhD research. The first place is the Respirator Carbon Group at the Department of Physics and Atmospheric Science, Dalhousie University, Halifax, Nova Scotia, Canada. I thank Prof. Dr. Jeff R. Dahn, who welcomed me in his lab when I was in a desperate situation. Dr. Philippe Westreich, my immediate supervisor, Hubert Fortier and Jock W. H. Smith are also thanked and acknowledged. The second place is Laboratoire de Chimie Moléculaire et Environnement (LCME), Université de Savoie, Polytech'Savoie, Bourget du Lac, France. A heartiest thanks to Prof. Dr. Laurent Duclaux, who offered me to come in his lab and later even visited Pakistan on my invitation and conducted a short course at my institute on activated carbons. He even arranged my visit to Spain and the testing of my samples. Another thanks for that too. Dr. Laurence Reinert, Dr. Jean-Marc Leveque and Dr. Nicolas Papaiconomou also deserve special gratitude. The third place is Carbon Materials and Environment Research Group (MCMA) of the

Universidad de Alicante, Alicante, Spain. I specially thank Prof. Dr. Diego Cazorla-Amoros, head of the research group for giving me a warm welcome in his lab and Dr. Juan Pablo Marco-Lozar, a sincere friend for even completing my left-over experiments in my absence.

Colleagues and co-workers at PIEAS deserve sincere thanks. Islam Khattak, Abdul Hafeez and Rashid Mahmood from lab staff, Muhammad Jibran, Faran Nabeel, Muhammad Irfan, Ikram-ul-Hassan, Safia Hassan, Sabiha Javeid, Azka Mehwish, Mehwish Mumtaz, Tahira Fatima, Zain-ul-Abideen, Sheeraz Mahboob, Abdul-Mateen were co-workers who have somehow helped me in the completion of this work. My PhD colleagues and all time friends, Abdul Basit Jilani, Muhammad Arshad, Abdul Waheed, Rashid Rizwan, Azhar-ul-Haq, Rana Iqtidar Shakoor, Muhammad Zahid Rana and Muhammad Shozab Mehdi were always available to share, help and to guide. Dr. Hassan bin Awais has been very kind in allowing me to use his BET surface area machine freely. Appreciations are due to Hafiz Rub Nawaz Shahid for assisting me in BET tests. Thanks to Dr. Tariq Yasin for useful discussions on TGA results.

A special acknowledgement for all my family members, who always accommodated when I was away and not able to join them. My parents, wife and children suffered a lot because of my long absence many times.

I offer my thanks to the Higher Education Commission of Pakistan (HEC) for providing financial support through their Indigenous PhD Fellowship Program, International Research Support Initiative Program and the travel grants program. Prof. Dr. Ata-ur-Rahman, ex-chairman HEC deserves special appreciation for his vision. Thanks to the French Embassy in Islamabad for providing financial support through the sandwich fellowship program.

Publications in Peer Reviewed Journals

- Farooq, A., Reinert, L., Leveque, J.-M., Papaiconomou, N., Irfan, N., Duclaux, L. (2012) "Adsorption of Ionic Liquids onto Activated Carbons: Effect of pH and Temperature." Microporous and Mesoporous Materials 158: 55-63.
- Farooq, A., Westreich, P., Irfan, N. and Dahn, J. (2009). "Adsorption of Copper Acetate onto Pretreated Activated Carbons over a Wide Concentration Range." Industrial & Engineering Chemistry Research 48(22): 9804-9808.
- Farooq, A., Hasany, S. M., Chaudhary, M. M. and Irfan, N. (2007). "Removal of Cd(II) from Aqueous Solution Using Blue Pine Sawdust: Equilibrium, Kinetics and Thermodynamic Studies." Main Group Metal Chemistry 30(6): 345-362.
- Farooq, A., Mehboob, S., Chaudhary, M. M. and Irfan, N. (2009). "High Concentration Preferential Adsorption of Zinc Acetate onto Acid Treated Activated Carbon for Impregnation Purposes." The Nucleus 46(3): 237-240.

Presentations in International Conferences

- Farooq, A., Hassan, S., Reinert, L., Leveque, J. M., Irfan, N. and Duclaux, L. (04-07 April, 2011). Adsorption de Liquides Ioniques en Solution Aqueuse sur des Carbones Activés. GFEC 2011, Orbey, France.
- Farooq, A., Reinert, L., Leveque, J. M., Papaiconomou, N., Irfan, N. and Duclaux, L. (11-16 July, 2010). Removal of Ionic Liquids and Methylene Blue using Various Activated Carbons. CARBON 2010, Clemson, SC, USA.
- Farooq, A., Irfan, M., Mehboob, S., Mahmood, A., Chaudhary, M. M. and Irfan, N. (11-16 July, 2010). High Concentration Preferential Adsorption of Zinc Chloride onto Acid Treated Activated Carbon and Comparison to Copper Acetate and Zinc Acetate. CARBON 2010, Clemson, SC, USA.
- Farooq, A., Reinert, L., Leveque, J. M., Papaiconomou, N., Irfan, N. and Duclaux, L. (18-22 Oct, 2010). Adsorption de Liquides Ioniques en Solution Aqueuse sur des Carbones Activés. MATERIAUX 2010, Nantes, France.
- Farooq, A., Reinert, L., Leveque, J. M., Papaiconomou, N., Irfan, N. and Duclaux, L. (29 Mar-01 Apr, 2010). Adsorption de Liquides Ioniques et de Bleu de Méthylène en Solution Aqueuse sur des Carbones Activés. GFEC 2010, Gréoux-les-Bains, France.
- Farooq, A., Westreich, P., Irfan, N. and Dahn, J. (13-18 July, 2008). The Adsorption of Copper Acetate onto Pretreated Activated Carbons over a Wide Concentration Range. CARBON 2008, Nagano, Japan.
- Farooq, A., Hasany, S. M., Chaudhary, M. M. and Irfan, N. (26-30 May, 2007). Blue Pine Sawdust as a Cheap Adsorbent for the Removal of Cadmium from Aqueous Solutions. 90th Canadian Chemistry Conference and Exhibition, Winnipeg, Manitoba, Canada.
- Farooq, A., Irfan, N., Chaudhary, M. M., Duclaux, L., Reinert, L., Lozar, J. P. M., Amoros, D. C., Jibrán, M., Nabeel, F., Mumtaz, M., Irfan, M., Hassan, L., Hassan, S. (17-22 June, 2012). Impregnation of TEDA onto activated carbon at pilot scale level. CARBON 2012, Krakow, Poland.

Table of Contents

Chapter	Description	Page
1	Introduction	1
	1.1 Control of Hazardous Pollutants	2
	1.1.1 Hazardous Air Pollutants	2
	1.1.2 Hazardous Water Pollutants	3
	1.2 Environmental and Health Impacts	5
	1.3 Scope and Layout of Thesis	8
2	Literature Review	9
	2.1 Adsorbents for Environmental Remediation	10
	2.1.1 Low-Cost Adsorbents	11
	2.1.1.1 Development and Application of Low-Cost Porous Adsorbents	11
	2.1.2 Microporous Adsorbents	13
	2.1.2.1 Historical Development of Activated Carbon	14
	2.1.2.2 Applications of Activated Carbon	16
	2.2 General Mechanism, Kinetics and Energetics	17
	2.2.1 Physisorption and Chemisorption	18
	2.2.2 Kinetics of Adsorption	18
	2.2.2.1 Mass Transfer Zone	22
	2.2.2.2 Frequently used Kinetic Models	23
	2.2.3 Thermodynamics of Adsorption	24
	2.3 Specific Hazardous Pollutants and their Control using Adsorbents	29
	2.3.1 Selective Adsorption of Gaseous Pollutants	30
	2.3.1.1 Impregnation of Dopants on Activated Carbon (AC)	31
	2.3.2 Selective Adsorption of Water Pollutants	41
	2.3.2.1 Adsorption of Ionic Liquids (ILs) on Activated Carbon	41
	2.3.2.2 Removal of Heavy Metals from Industrial Waste	44
	2.4 Summary	46
3	Experimental Setups and Procedures	49
	3.1 Pilot Scale Impregnation Setup	49
	3.1.1 Concept and Requirement of Experimental Setup Design	50
	3.1.2 Design of Pilot Scale Setup	51
	3.1.2.1 Design of Fluidized Bed Adsorbing Tower (FBT)	51
	3.1.2.2 TEDA Vessel Design	54
	3.1.2.3 Blower Design	55
	3.1.2.4 Design of Cross Flow Heat Exchanger	56
	3.1.3 Sequence and Procedures for an Experimental Run on Setup	60
	3.1.4 Experimental Parameters Studied for Optimization	61
	3.1.4.1 Process Time Optimization	61
	3.1.4.2 Process Temperature Optimization	61
	3.1.4.3 Percentage TEDA Input optimization	61
	3.1.5 Characterization of TEDA impregnated AC samples	63
	3.1.5.1 Analysis of TEDA Impregnated AC	63
	3.1.5.2 Nitrogen Adsorption Isotherms, TGA, FTIR and SEM Analysis	63
	3.1.5.3 Performance Testing for Adsorption of SO ₂ and CH ₃ I	63

3.2 Pretreatment and Copper Acetate Impregnation on AC for Adsorption of Chemical Gases and Toxicants	65
3.2.1 Activated Carbons	65
3.2.2 Heat Treatment of Commercial Activated Carbons	66
3.2.3 Acid Treatment of Commercial Activated Carbons	66
3.2.4 Boehm titrations, Surface Area, XRD, TGA and pH measurement	66
3.2.5 Copper Acetate Adsorption on Activated Carbons	67
3.2.5.1 Isotherms Fitting to Adsorption Data	67
3.3 Testing of Cyclohexane Breakthrough Times	68
3.3.1 Cyclohexane Testing Setup	68
3.4 Setup for contaminant removal studies from aqueous streams	70
3.4.1 Removal of Ionic liquids using AC	70
3.4.1.1 Activated carbons used for this study	71
3.4.1.2 Morphological structure and EDS characterization	71
3.4.1.3 N ₂ adsorption-desorption at 77 K	71
3.4.1.4 p <i>H</i> _{PZC} measurement	72
3.4.1.5 Boehm titrations	72
3.4.1.6 Ionic liquids used in this study	73
3.4.2 Cadmium removal using low cost adsorbent	74
3.4.2.1 Reagents/Buffers and Materials	74
3.4.2.2 Equipment	75
3.4.2.3 Sawdust	75
3.4.2.4 Procedure	76
3.5 Summary	77
4 Results & Discussion	78
(Modification of AC Surfaces and Removal of Selected Hazardous Air Pollutants)	
4.1 TEDA impregnated activated carbon	78
4.1.1 Results of Impregnation of TEDA on AC at pilot scale setup	78
4.1.1.1 Process Time Optimization	79
4.1.1.2 Process Temperature Optimization	80
4.1.1.3 Optimization of TEDA Input Weight	81
4.1.1.4 Summary of optimized parameters for TEDA impregnation	82
4.1.2 Characterization of the selected TEDA impregnated AC samples	84
4.1.2.1 Estimation of doped TEDA % by UV-visible spectroscopy	84
4.1.2.2 Nitrogen adsorption isotherms and porosimetry tests	85
4.1.2.3 Elemental analysis	86
4.1.2.4 Thermogravimetric Analysis (TGA)	87
4.1.2.5 FTIR analysis	87
4.1.2.6 Scanning electron microscopy	89
4.1.3 Performance Testing of the Impregnated Activated Carbon	90
4.1.3.1 Sulfur Dioxide (SO ₂) Adsorption Capacity	90
4.1.3.2 Methyl Iodide Removal Efficiency	92
4.2 Pretreatment and copper acetate impregnation for adsorption of chemical gases and toxicants	93
4.2.1 Characterization of raw and modified activated carbons	94
4.2.1.1 Boehm titrations	94
4.2.1.2 Porosimetry and surface area analysis	95
4.2.1.3 X-Ray diffraction analysis	98
4.2.1.4 Breakthrough time testing for cyclohexane adsorption	99
4.2.2 Copper acetate adsorption on activated carbons	101
4.2.2.1 Double Langmuir isotherm fitting	101
4.2.2.2 Combined isotherm fitting	102
4.2.2.3 Effect of pH on adsorption	106
4.3 Summary	107

5 Results & Discussion	109
(Removal of Selected Hazardous Water Pollutants)	
5.1 Adsorption of ionic liquids on various activated carbons	109
5.1.1 Characterization of activated carbons	110
5.1.1.1 Surface chemistry	110
5.1.1.2 Porosity characterization	111
5.1.2 Kinetics of adsorption	114
5.1.3 Factors affecting adsorption	115
5.1.3.1 Evolution of adsorption isotherms with pH	115
5.1.3.2 Evolution of adsorption isotherms with activated carbon type	117
5.1.3.3 Evolution of adsorption isotherms with IL type	119
5.1.4 Effect of temp on ILs adsorption (thermodynamic parameters)	120
5.2 Cadmium removal using sawdust as a low cost adsorbent	123
5.2.1 Optimization of Cd(II) ions adsorption	123
5.2.2 Adsorption isotherms	126
5.2.3 Thermodynamics of adsorption	128
5.2.4 Influence of common ions on the adsorption	129
5.2.5 Kinetics	130
5.2.6 Sorption mechanism	131
5.3 Summary	132
6 Conclusion and Recommendations	134
References	138
Appendix-I	162
Appendix-II	163
Appendix-III	166
Appendix-IV	168
Appendix-V	183
Appendix-VI	186
Appendix-VII	189
Appendix-VIII	190

List of Figures

Figure 2.1:	(a) Relative concentration of toxicant in gas phase within carbon bed plotted against position along the carbon bed, (b) Shape of mass transfer zone is mirrored in the detected concentration at the end of bed, plotted against time [Fortier, 2007]	22
Figure 2.2:	Types of adsorption isotherms according to IUPAC [Sing, 1985]	25
Figure 3.1:	Structure of triethylenediamine (TEDA)	50
Figure 3.2:	Schematic flow sketch of required TEDA pilot scale impregnation Setup	51
Figure 3.3:	Drawings and as-built details of adsorbing tower (FBT)	53
Figure 3.4:	Drawings and as-built details of TEDA vessel	55
Figure 3.5:	Assembly drawings and as built details of Blower	56
Figure 3.6:	Drawings and as-built details of heat exchanger	58
Figure 3.7:	Pilot Scale TEDA Impregnation Setup for Nuclear Grade Activated Carbon Production	59
Figure 3.8:	Flow diagram of cyclohexane breakthrough test setup	69
Figure 3.9:	Infrared based cyclohexane adsorption test setup installed in a fume hood	69
Figure 4.1:	Time Optimization for (a) BET Surface Area and (b) Percent TEDA Impregnated	79
Figure 4.2:	Temperature Optimization for (a) BET Surface Area and (b) % TEDA Impregnated	80
Figure 4.3:	TEDA Input versus (a) BET Surface Area and (b) Percent TEDA Impregnated	81
Figure 4.4:	N ₂ adsorption isotherms for FNS00, FNS09, FNS12, FNS27, FNS28 and FNS29	86
Figure 4.5:	Pore size distribution curves for FNS00, FNS09, FNS12, FNS27, FNS28 and FNS29	86
Figure 4.6:	TGA curves of FNS00, FNS09, FNS12, FNS27, FNS28 and FNS29	87
Figure 4.7:	FTIR spectra of FNS00, FNS09, FNS12, FNS27, FNS28 and FNS29	88
Figure 4.8:	Expanded FTIR spectra of FNS00, FNS09, FNS12, FNS27, FNS28 and FNS29	88
Figure 4.9:	SEM micrographs of (a) FNS00, (b) FNS09, (c) FNS12, (d) FNS27, (e) FNS28 and (f) FNS29 at x 5000 magnification	89
Figure 4.10:	SO ₂ adsorption capacity of raw and TEDA doped AC samples	91

Figure 4.11:	Nitrogen adsorption isotherms of GC, HTGC, H2TGC, GG, HTGG, H2TGG, GAC, HTGAC and H2TGAC	96
Figure 4.12:	BET and Langmuir surface areas (m^2/g) of GC, HTGC, H2TGC, GG, HTGG, H2TGG, GAC, HTGAC and H2TGAC	96
Figure 4.13:	Pore size distribution (PSD) of GC, HTGC, H2TGC and NITGC and NITGC	97
Figure 4.14:	XRD results of HTGC, HTGG, HTGAC, H2TGC, H2TGG and a TGA sample (heated and cooled thrice in argon, hydrogen, argon)	98
Figure 4.15:	Typical breakthrough curves of cyclohexane adsorption onto GC in dry/dry & in wet/wet conditions	99
Figure 4.16:	Bar graph showing breakthrough times (minutes) of raw and modified AC samples for adsorption of cyclohexane in dry/dry & in wet/wet conditions	100
Figure 4.17:	Adsorption Isotherm for copper acetate on H2TGC, fitted to a sum of two Langmuir Isotherms	101
Figure 4.18:	Adsorption Isotherm for copper acetate on NITGC, fitted to a combination of Langmuir and Freundlich Isotherms (AAS data only)	102
Figure 4.19:	Adsorption isotherm for copper acetate on GC and H2TGC fitted to a sum of two Langmuir isotherms, and on NITGC fitted to a combination of Langmuir and Freundlich isotherms	103
Figure 5.1:	SEM images of (a) Chinese AC, (b) Fabric AC, (c) Artichokes AC and particle size distribution of (d) Chinese & Artichokes ACs	111
Figure 5.2:	N_2 adsorption (full symbol)/desorption (empty symbol) isotherms at 77 K on activated carbons: Chinese (■), Zorflex Fabric (◆), and Artichokes (▲)	111
Figure 5.3:	Pore Size Distribution obtained by bidimensional Non-Local Density Functional Theory (2D-NLDFT) method based on carbon finite slit pore (with an asymmetric ratio equal to 6) of (a) Artichoke AC, (b) Zorflex fabric AC, and (c) Chinese AC	113
Figure 5.4:	Kinetics of adsorption of Octyl Pyridinium cation (OPy^+) on activated carbons: Chinese (◆), Zorflex (■), and Artichokes (▲)	114
Figure 5.5:	Adsorption of $BMIm^+$ (▲), $OMIm^+$ (◇), and OPy^+ (■) onto Chinese (dotted line), Fabric (solid line) and artichokes (dashed line) ACs at pH = 2	116
Figure 5.6:	Adsorption of $BMIm^+$ (▲), $OMIm^+$ (◇), and OPy^+ (■) onto Chinese (dotted line), Fabric (solid line) and artichokes (dashed line) ACs at pH = 9	116
Figure 5.7:	Adsorption of $BMIm^+$ (▲), $OMIm^+$ (◇) and OPy^+ (■) onto Artichoke AC at pH = 2 (dotted line), 7 (dashed line) and 9 (solid line)	116

Figure 5.8:	Difference between maximum adsorption uptakes (Q_m) for BMIm ⁺ (◆), OMIIm ⁺ (■) and OPy ⁺ (▲) measured at pH = 9 and 2 as a function of the oxygen content of activated carbons (Chinese, Fabric and Artichokes)	118
Figure 5.9:	Experimental and simulated adsorption isotherms of OPyBr (a) at pH=7 for T=25°C (◇), 40°C (■), and 55°C (▲); and (b) at pH=9 for T=20°C (◇), 30°C (■), and 50°C (▲). The adsorption isotherms were simulated using the Langmuir-Freundlich models (solid line). The parameters of the fits are reported in Table 5.5	120
Figure 5.10:	Adsorption uptake dependence (Q_0 in mmol/g), at pH = 7 (dotted line) and pH = 9 (solid line), of the isosteric Gibbs free energy for BMIm ⁺ (▲), OMIIm ⁺ (■), and OPy ⁺ (◆), and of entropy of adsorption of the ILs for BMIm ⁺ (▽), OMIIm ⁺ (□), and OPy ⁺ (◇)	122
Figure 5.11:	The distribution ratio of Cd(II) ions (18μM) onto sawdust (0.2 g/10 mL) as a function of shaking time	124
Figure 5.12:	The distribution ratio of Cd(II) ions (18μM) as a function of dosage of sawdust after 30 min shaking	125
Figure 5.13:	The effect of initial concentration of Cd(II) ions on its adsorption onto sawdust (0.2g/10mL)	125
Figure 5.14:	Langmuir adsorption isotherm of Cd(II) ions onto sawdust	126
Figure 5.15:	Freundlich isotherm of Cd(II) ions adsorption onto sawdust	127
Figure 5.16:	D-R adsorption isotherm of Cd(II) ions onto sawdust	127
Figure 5.17:	The variation of equilibration constant for the adsorption of Cd(II) ions (18 μM) onto sawdust (0.2 g/10 mL) as a function of temperature	128
Figure 5.18:	Lagergren plot for the adsorption of Cd(II) ions onto sawdust	131
Figure 5.19:	Morris-Weber profile of Cd(II) ions (18 μM) adsorption onto sawdust (0.2 g/10 mL)	132

List of Tables

Table 3.1:	Designed specifications of FBT	53
Table 3.2:	Specifications of designed TEDA Vessel	54
Table 3.3:	Specifications of Blower	56
Table 3.4:	Specifications of heat exchanger	57
Table 3.5:	Size, Shape & Structure of Ionic Liquids used	73
Table 3.6:	Proximate analysis of sawdust (<i>Pinus Wallichiana</i>)	75
Table 3.7:	Fractionation of crude fiber of sawdust (<i>Pinus Wallichiana</i>)	76
Table 4.1:	Surface Area and Micropore Volume of Raw and TEDA doped AC samples	85
Table 4.2:	Elemental analysis results of Raw and TEDA doped AC samples	87
Table 4.3:	Adsorption capacity of raw and TEDA doped AC samples for SO ₂	91
Table 4.4:	Removal Efficiency (% E) of raw and TEDA doped AC samples for radioactive methyl iodide (CH ₃ I ¹³¹)	93
Table 4.5:	Boehm titration results of raw and heat treated AC samples (mmol/g AC)	94
Table 4.6:	BET surface areas, micropore volumes and total pore volumes of GC, HTGC, H2TGC	97
Table 4.7:	Adsorption parameters from Langmuir fit for GC and H2TGC	103
Table 4.8:	Adsorption parameters from combined Langmuir and Freundlich fits for NITGC	104
Table 4.9:	pH values at different initial concentrations as measured for copper acetate solution before and after stirring with NITGC	106
Table 4.10:	pH values at different initial concentrations as measured for copper acetate solution before and after stirring with H2TGC	107
Table 4.11:	Summary of suggested interactions at low and high concentrations in the cases of GC, H2TGC and NITGC	107
Table 5.1:	Boehm Titration results and pH of point of zero charge (pH _{pzc}) of Chinese, Fabric and Artichokes ACs	110
Table 5.2:	Textural properties of Chinese, Fabric and Artichokes ACs obtained by N ₂ adsorption/desorption at 77K	112
Table 5.3:	Results of the best kinetics fitting (R-square correlation coefficient > 0.98) for the three activated carbons (Chinese, Fabric and Artichokes), among the four tested models: pseudo first order, pseudo second order, Elovich and diffusion.	115

Table 5.4:	Comparison of IL volume to the Micropore Volume of Chinese, Fabric and Artichokes ACs	119
Table 5.5:	Isosteric Gibbs free energy, enthalpy and entropy of adsorption of the ILs (at constant value of adsorption uptake) at pH = 7 and pH = 9	122
Table 5.6:	Effect of pH on the adsorption of Cd(II) ions onto sawdust (0.2 g/10 mL) using 30 min agitation time	124
Table 5.7:	The effect of anions on the retention of Cd(II) ions (18 mM) onto sawdust (0.2 g/10 mL) after 30 min equilibration time	129
Table 5.8:	The effect of cations on the retention of Cd(II) ions (18 mM) onto sawdust (0.2 g/10 mL) after 30 min equilibration time	130