

Original Research Article

Synthesis and characterization of ammonium ionic liquids and their antimicrobial and computational overview

Abstract

The ammonium acetate and formate ionic liquids (ILs) were synthesized in neutralization reaction with different cations of alkanol amine and alkyl amine with formic and acetic acid as anion. The reaction was monitored by thin layer Chromatography and the synthesized six ILs was characterization by FT-IR spectroscopy and ^1H , ^{13}C Nuclear Magnetic Resonance (NMR) spectroscopy. The spectroscopic analysis of the synthesized ILs gives the good agreement for the confirmation of their synthesis and purity. These compounds were tested in computing program of computational chemistry through two basis set, Mm+ and PM3 for the calculation using molecular mechanics by Hyperchem^R Release 8.0 (Hypercube, Inc.). Single point calculations were carried out to evaluate the data profile on the physical properties, such as, total energy, binding energy, hydration energy, dipole moment, heat of formation and energies for HOMO and LUMO. Furthermore the synthesized ILs was screening antimicrobial activity against both of gram positive and gram negative bacteria such as *Bacillus cereus*, *Staphylococcus aureus*, *Escherichia coli*, *Salmonella typhi*, *Pseudomonas aeruginosa* and *Shigella dysenteriae* and two phytoathogenic fungi such as *Aspergillus niger* and *Rhizopus azzahra* to establish as green and environment benign solvent.

Keywords: Ethanolamine, FT-IR, NMR, HyperchemR, Antimicrobial activity.

1. INTRODUCTION

Ionic Liquids (ILs) was introduced almost a century ago by the observation of Paul Walden in a neutralization reaction of ethanolanmonium nitrate [1]. They belong to a novel class of low temperature (typically $<100^\circ\text{C}$) molten salts, consisted of discrete anions and cations [2]. They exhibit, in most cases, relatively low viscosities, thermally stabilities, high thermal conductivities, a large electrochemical window, highly polar and non volatility [3]. Millions of ILs can be synthesized and characterized with variety of cations and anions in theoretical study which are also known as the designer solvent due to their tunable physical and chemical properties [4]. Due to their unique properties [5], ILs have become one of the most important research targets in chemistry, physics, engineering and material science and technology [6].

There is a strong interest in ILs as alternatives for volatile organic solvents in the organic synthesis [7-8]. They can act as solvents for chemical reactions, including catalytic reactions [9-13]. ILs are found as the use in electrochemical applications [7-8, 14-15], e.g. electrolytes in batteries [16], in photovoltaic devices [17], and a medium for electrode-position of metals. ILs can also find applications in separation sciences as solvents for extraction processes [18], as a stationary phase for gas chromatography [19], as well as in mass spectrometry [20]. In other hands the fancacing physio-chemical properties of alkanolanmonium based ILs including ion conducting, viscous, non-volatile and non-flammable, they are now used in various fields both of academia and industries as solvent, catalyst, reaction media, pharmaceutical ingredient and solvent, polymerization reaction [21]. In addition, the biodegradability of ammonium cations is considerably enhanced when hydroxyl groups are incorporated. To make in utilizing as the green and promising eco-friendly solvents in the separation process, ILs can serve as nonvolatile entertainers to break zoetrope and enable a more energy efficient. [22]. Ammonium based ILs especially ethanolanmonium ILs are easy to prepare and the raw materials are mostly available with cheapness. To investigate

51 The study of alkanolammonium based ILs, three cations such as different types of cation such as
 52 ethanolamine, diethylamine and triethanolamine were used to form ILs with anion as formate and acetate.
 53 To ensure their use as solvent, the physio-chemical properties are the greatest goal. Estimation the
 54 physio-chemical properties are a process of the time consuming and waste of chemical materials. To
 55 save the time and chemicals, the computer programming e.g. Hyperchem[®] Release 8.0 (Hypercube, Inc.)
 56 was used to calculated the total energy, binding energy, Heat of formation , HOMO, LUMO,
 57 LUMO~HOMO and different vibrational stretching of bonds [22-24]. Different cations and anions were
 58 used to make a comparison effects in physio-chemicals properties through computer programming and
 59 antimicrobial toxicity. Most of these are highly polar and well soluble in water. So there is a large scope to
 60 pollute water body and environment [25].The antimicrobial toxicological profile of these ILs was not well
 61 documented and the information are not enough for further applications. Due to the short life span,
 62 bacteria and fungi are the best starting point to investigate and estimate the toxic bio-data of synthesized
 63 formate and acetate ammonium ILs. The bacterial toxicity level in the form of antimicrobial activity against
 64 gram positive and gram negative bacteria was screened and evaluated for confirming their use in specific
 65 fields [26-27].

66 **2. EXPERIMENT**

67 **2.1 Materials and reagents**

68 All the chemicals were of research grade and used without further purification unless otherwise stated. All
 69 the solvent were obtained upon distillation before use. Formic acid (Merck KGaA), Acetic acid (Merck
 70 KGaA), 2-aminoethanol (Marck KGaA), Ethylamine (Merck KGaA), Triethanolamine (Merck KGaA) was
 71 analytical grade commercial products. Thin Layer chromatography powder (Merck KGaA), standard
 72 antibiotic Gentamycin were purchased for the reaction workup FT-IR spectrophotometer, SHIMADZU,
 73 Japan, range 600 -4500 cm⁻¹) was used with KBr disc technique. The synthesis, computer programming
 74 and the characterization were done at department of chemistry in University of Chittagong, and
 75 Chittagong, Bangladesh. The antimicrobial activity was at department of Microbiology in University of
 76 Chittagong, and Chittagong, Bangladesh. The ¹H, ¹³C NMR Spectroscopy was done in Iwate University,
 77 Japan.

78 **2.2 Ionic Liquids Synthesis and Purification**

79 The synthesis of alkanol or alkyl ammonium ILs
 80 consists in an acid-basic neutralization
 81 reaction[28]. The base, in this case
 82 ethanolamine, diethylamine and triethanol
 83 amine were added under stirring in a slow drop
 84 wise about 20-25 minute maintaining the
 85 temperature using ice bath from exothermic
 86 heat release on a glass flask with

87 the acid (formic and acetic acid respectively).
 88 Then the mixture was stirred for 24 hours at
 89 room temperature, to obtain a viscous clear
 90 liquid. The reaction was monitored by Thin
 91 layer chromatography (TLC).The reaction
 92 products are an ester and a salt of
 93 ethanolamine [28-30] (The ILs purification
 94 process consists in a strong agitation and slight
 95 heating, at 323.15K, for the vaporization of
 96 impurities (residual non reacted and water)
 97 under vacuum of 20 kPa. Humidity below 0.1%
 98 was obtained after this purification process[28,
 99 30] (Alvarez et al., 2010; Iglesias et al. 2010),
 100 and the liquids presented a limp and viscous
 101 appearance. The ammonium salt formation was
 102 proved in by FT-IR spectroscopy by using a Shimadzu IR.
 103 The structure of synthesized IL is shown in figure 1.
 104
 105

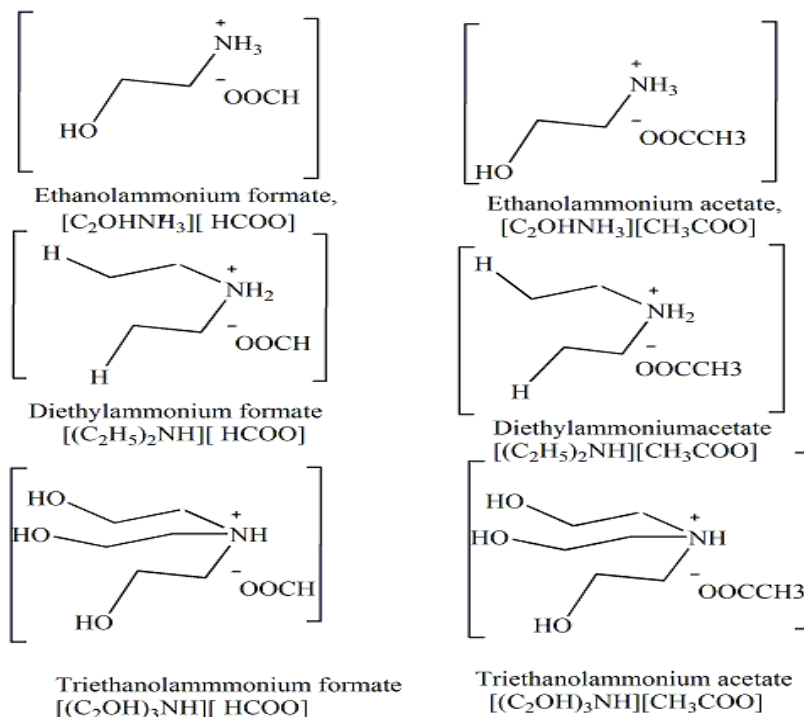


Figure 1: synthesized ammonium based IL

106 3. RESULT AND DISCUSSION

107 Ethanolammoniumformate (IL01), $[\text{C}_2\text{OHNH}_3][\text{HCOO}]$, Yield: 92 %, MW 107.11gm;
 108 Found: %C = (33.62-34.02), %H= 8.47- 8.23; %N= 13.08-12.96, Calculated: ^1H NMR: 2.0 (s, 1H, OH),
 109 3.52(t, 2H, NCH_2), 4.27 (t, 2H, OCH_2), 7.0 (bs, 3H, NH_3), 9.6 (s, 1H, CH).
 110 ^{13}C NMR: 63.3 ($-\text{CH}_2$), 42.9 ($-\text{CH}_2$), 47.0 ($-\text{CH}_3$).

111 FTIR: 3572 ($-\text{OH}$), 3147 (N-H) asymmetry, 3012 (C=C) in benzene ring, 2966 (N-H) symmetry, 2931 (C-
 112 H) asymmetry, 2897 (C-H) symmetry, 2360, 1589 (C-O) asymmetry, 1485 ($-\text{CO}$) symmetry cm^{-1} .

113 Ethanolammoniumacetate (IL02), $[\text{C}_2\text{OHNH}_3][\text{CH}_3\text{COO}]$, Yield: 83.0 %, MW121.09 gm.

114 Found: %C = 39.64-39.43, %H= 9.16- 9.51, % N= 11.56- 11.27. Calculated: ^1H NMR: 2.0 (s, 1H, OH), 2.20 (s,
 115 3H, CH_3), 3.52 (t, 2H, NCH_2), 4.21 (t, 3H, OCH_2), 7.0 (bs, 3H, NH_3).

116 ^{13}C NMR: 63.3 ($-\text{CH}_2$), 42.9 ($-\text{CH}_2$), 56.0 (CH_2), 17.0 ($-\text{CH}_3$).

117 FTIR: 3437 ($-\text{OH}$), 3414(N-H asymmetry), 3390 (N-H) asymmetry, 2939 (C-H) asymmetry, 2893 (C-H)
 118 symmetry, 1697 (C-O) symmetry, 1723 (C-O) asymmetry, cm^{-1} .

119 Diethylammoniumformate (IL03), $[(\text{C}_2\text{H}_5)_2\text{NH}][\text{HCOO}]$, Yield%: 93.0 %, MW: 119.10 gm

120 Found: %C = 50.38 – 50.22, %H= 11.00- 11.76, % N= 11.75- 12.03, Calculated:

121 ^1H NMR: 1.61 (t, 6H, $2\times\text{CH}_3$), 3.27 (q, 4H, $2\times\text{CH}_2$), 7.0 (bs, 2H, NH_2), 9.2 (s, 1H, CH)

122 ^{13}C NMR: 11.08 ($-\text{CH}_3$), 42.2 ($-\text{CH}_2$), 30.6 ($-\text{CH}_3$), 47.0 ($-\text{CH}_3$), 6.8 ($-\text{CH}_3$), 6.8 ($-\text{CH}_3$).

123 FTIR: 3433(N-H) asymmetry, 3059(N-H) symmetry, 2993 (C-H) asymmetry, 2951 (C-H) symmetry, 2885
 124 and 2796 alkyl C-H stretch asymmetry and symmetry of ethylamine, 1650 (C-O) asymmetry , 1589 (C-
 125 O) symmetry, cm^{-1} .

126 Diethylammoniumacetate (IL04), $[(\text{C}_2\text{H}_5)_2\text{NH}][\text{CH}_3\text{COO}]$, Yield%: 96.0 %, MW: 133.12 gm, Found: %C =
 127 54.09- 53.69, %H= 11.36- 10.96, % N= 10.52. 10.28.

128 ^1H NMR: 1.61 (t, 6H, $2\times\text{CH}_3$), 2.20 (s, 3H, CH_3), 3, 37 (q, 4H, $2\times\text{CH}_2$), 7.20 (bs, 2H, NH_2)

129 ^{13}C NMR: 11.08 ($-\text{CH}_3$), 42.2 ($-\text{CH}_2$), 30.6 ($-\text{CH}_3$), 56.0 ($-\text{CH}_3$), 17.0 ($-\text{CH}_3$), 6.8 ($-\text{CH}_3$), 6.8 ($-\text{CH}_3$).

130 FTIR: FTIR: 3437(N-H) asymmetry, 3414 (N-H) symmetry, 3035 (C-H) asymmetry, 2893 (C-H) symmetry,
 131 2885 and 2796 alkyl C-H stretch asymmetry and symmetry of ethylamine, 1535 (C-O) asymmetry , 1500
 132 (C-O) symmetry, cm^{-1} .

133
 134 Triethanolammoniumformate, Code no: (IL05), $[(\text{C}_2\text{OH})_3\text{NH}][\text{HCOO}]$, Yield%: 81 %, MW: 195.52

135 Found: %C = (45.90- 45.70), %H= (9.44- 9.66), % N= (7.65-7.40). Calculated: ^1H NMR: 2.0 (s, 3H, $3\times\text{OH}$),
 136 3.43 (t, 6H, $3\times\text{NCH}_2$), 3.77 (t, 6H, $3\times\text{OCH}_2$), 7.08 (s, 1H, NH), 9.62 (s, 1H, CH).

137 ^{13}C NMR: 58.6 ($-\text{CH}_2$), 59.5 ($-\text{CH}_2$), 38.1 ($-\text{CH}_3$), 59.5 ($-\text{CH}_2$), 58.6 ($-\text{CH}_2$), 47 ($-\text{CH}_3$), 16.9 ($-\text{CH}_3$), 55.8(
 138 CH_2).

139 FTIR: 3417 for N-H bond of ammonium ion, 2947 (C-H) asymmetry, 2893 (C-H) asymmetry, 1666 (C-O)
 140 symmetry, 1593(C-O) symmetry.

141 Triethanolammoniumformate(IL06), $[(\text{C}_2\text{OH})_3\text{NH}][\text{CH}_3\text{COO}]$, Yield%: 84.0 %, MW: 210.52 Found: %C=
 142 (45.60- 45.28), H%= (9.74- 9.89), % N= (6.65- 6.52).

143 Calculated: ^1H NMR: 2.05 (s, 3H, $3\times\text{OH}$), 2.26 (s, 3H, CH_3), 3.21 (t, 6H, $3\times\text{NCH}_2$), 3.87 (t, 6H, $3\times\text{OCH}_2$),
 144 7.0 (s, 1H, NH).

145 ^{13}C NMR: 58.6 ($-\text{CH}_2$), 59.5 ($-\text{CH}_2$), 38.1 ($-\text{CH}_3$), 59.5 ($-\text{CH}_2$), 58.6 ($-\text{CH}_2$), 56.0 ($-\text{CH}_2$), 17.0 ($-\text{CH}_3$), 55.8
 146 ($-\text{CH}_2$).

147 FTIR: 3363(N-H asymmetry), 3151(N-H asymmetry), 3016 (C-H) asymmetry, 2970 (C-H)
 148 symmetry, 2931, and 2897, alkyl C-H stretch asymmetry and symmetry of ethanolamine, 2839.22 C-H
 149 asymmetry and C-H symmetry, 1566 (C-O) asymmetry, 1485 (C-O) symmetry cm^{-1} .

150 4. COMPUTATIONAL OVERVIEW OF THE SYNTHESIZED ILS

151 Computational chemistry is a branch of chemistry that uses computers to assist in solving chemical
 152 problems uses the results of theoretical chemistry, incorporated into efficient computer programs, to
 153 calculate the structures and properties of molecules and solids[31]. There are two broad areas within
 154 computational chemistry devoted to the structure of molecules and their reactivity as molecular
 155 mechanics (MM) and quantum mechanics (QM) [32]. They both perform the same basic types of
 156 calculations. Molecular mechanics uses the laws of classical physics to explain and interpret the structure
 157 and properties of molecules. Molecular mechanics methods are available in many computer programs,
 158 including MM3, HyperChem, Gaussian, Quanta, Sybyl and Alchemy.

159 The tested compounds were constructed using Hyperchem^R Release 8.0 (Hypercube, Inc.) drawing
 160 platform and are geometrically optimized for further calculations. For the calculation using molecular
 161 mechanics, these compounds were tested in two basis set, Mm+ and PM3. Single point calculations were
 162 carried out to create the data profile on the physical properties, such as, total energy, binding energy,
 163 hydration energy, dipole moment, heat of formation and energies for HOMO and LUMO.
 164 In the calculation, total energy and free energy for the compounds are same due to zero value of entropy.
 165 According to Gibb's equation, $A = E - TS$ (where, A=Free energy, T=Temperature, E=Total energy,
 166 S=Entropy); If S=0, then A=E-0 and A=E.

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 168 Table01: Properties calculation using optimized Mm+ set.
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Chemicals	Calculation type	Total energy (kcal/mol)	Dipole moment (debyes)	Hydration energy (kcal/mol)
IL01	Mm+	0.50945	3.50600	3.900
IL02	Mm+	-0.69262	1.53409	-11.670
IL03	Mm+	3.74434	2.18200	-2.620
IL04	Mm+	7.14584	4.75700	-14.100
IL05	Mm+	14.66050	2.40697	-8.660
IL06	Mm+	17.62987	6.28500	-17.940

170

171 Table02: Properties calculation using optimized PM3 set.

Chemicals	IL01	IL02	IL03	IL04	IL05	IL06
Calculation type	PM3	PM3	PM3	PM3	PM3	PM3
Total energy (kcal/mol)	-34720.769	-38168.335	-48382.8	-51832.872	-61706.87	-65498.4
Binding energy (kcal/mol)	-1262.9246	-1542.4632	-1916.36	-2198.4278	-2533.345	-2855.42
Heat of formation (kcal/mol)	-93.863591	-98.308165	-137.554	-144.52623	-196.9071	-191.771
HOMO (eV)	-10.54155	-10.06375	-10.41844	-10.24847	-3.97568	-8.92506
LUMO (eV)	1.13607	0.90474	0.93535	0.94809	0.83742	-1.65566
LUMO~HOMO (eV)	11.677621	10.968487	11.353785	11.196557	4.813104	7.269402

172 The properties estimated by the basis set Mm+ and PM3, can be used to explain the molecular properties
 173 and variations can be attributed by the hydrogen bonding, binding energy calculations. According to the
 174 results found by the software, it is obvious that the total energy is more for higher alkyl chain ILs compare
 175 to the lower alkyl chain ILs. Standard methods were employed for all the test and experiments. Lower
 176 carboxylate such as formate and acetate ILs were tested in molecular programming software
 177 Hyperchem^R Release 8.0 and found good result for properties calculation.

178 5. EXPERIMENT FOR ANTIMICROBIAL ACTIVITY

179 5.1 Preparation of IL solutions in different concentrations

180 The required amount of the sample was measured in Digital balance with highly carefully so that no
 181 impurities were obtained. Then the required 1.5 mL distilled water was added and well shake for well
 182 soluble.

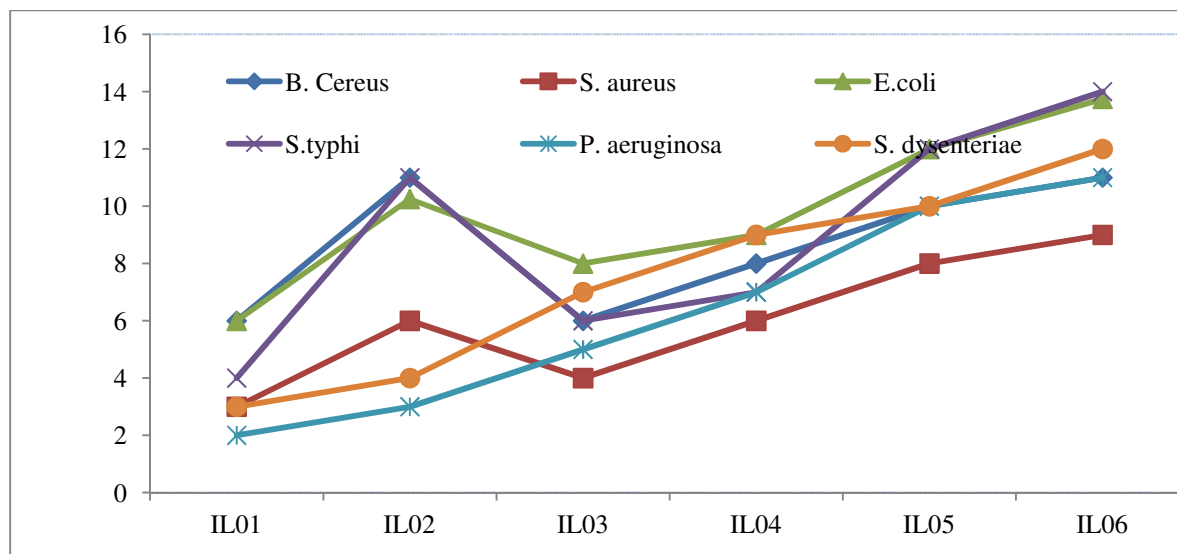
183 **5.2 Antimicrobial assay**

184 A preliminary investigation on the antibacterial activities of pure ILs was performed through
 185 measurements of primary screening both the gram-positive and gram- negative bacteria. Antibacterial
 186 screening of the test ILs were carried out with six bacterial pathogens, such as, *Bacillus cereus*,
 187 *Staphylococcus aureus*, *Escherichia coli*, *Salmonella typhi*, *Pseudomonas aeruginosa* and *Shigella*
 188 *dysenteriae*. This method was carried on via well diffusion method[33-34]. The bacterial inhibition zone
 189 (subtracting the well diameter 5.0 mm) was measured in mm scale with consideration ± 1.0 with all taking
 190 value. All the measurements were done in triplicate and the averages were listed in table 03. The initial
 191 concentration was maintained for all ILs in 1000 mM/L, 500 mM/L and 100 mM/L in distilled water or
 192 methanol. A control plate is always observed for the ILs if there is any significant inhibition occurred for
 193 the solvent. The results showed that all compounds had antimicrobial activity against bacterial pathogens
 194 used in this study.

195 Table03: Zone of inhibition (in mm) observed in three different concentrations (X=1000mM/L,
 196 Y=500mM/L, Z=100mM/L).

pathogens and ILs	<i>B. Cereus</i>			<i>S. aureus</i>			<i>E.coli</i>			<i>S.typhi</i>			<i>P. aeruginosa</i>			<i>S. dysenteriae</i>		
	X	Y	Z	X	Y	Z	X	Y	Z	X	Y	Z	X	Y	Z	X	Y	Z
IL01	6.0	0	0	3.0	0	0	6.0	0	0	4.0	0	0	2.0	0	0	3.0	0	0
IL02	11.0	0	0	6.0	0	0	10.25	6.0	0	11.0	5.2	0	3.0	0	0	4.0	0	0
IL03	6.0	0	0	4.0	0	0	8.0	0	0	6.0	5.0	0	5.0	0	0	7.0	0	0
IL04	8.0	0	0	6.0	0	0	9.0	0	0	7.0	0	0	7.0	0	0	9.0	0	0
IL05	10.0	6.0	0	8.0	6.0	0	12.0	8.0	0	12.0	0	0	10.0	0	0	10.0	14.0	0
IL06	11.0	0	0	9.0	0	0	13.75	0	0	14.0	0	0	11.0	0	0	12.0	0	0

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199 **Figure 2: A comparison study of toxicity for ammonium bases ILs.**

200 From the above graph, it was shown that the antimicrobial toxicity of primary, secondary and tertiary is not
 201 same. In all case, primary ammonium based ILs has lower toxicity than secondary ammonium based ILs.
 202 Again secondary ammonium based ILs has higher toxicity than tertiary ammonium based ILs. But in all

203 case, anion also has an activity on the toxicity. The ammonium formate has lower toxicity than
 204 ammonium acetate.

205 **5.3 Antifungal Screening Test**

206 *Aspergillus niger* and *Rhizopus azzahra* were used for evaluating the antifungal activity of all synthesized
 207 compounds. The antifungal activity was evaluated by Well diffusion method [35]. The media was altered
 208 Potato dextrose broth (abbreviated "PDB") is formulated identically to PDA, omitting the agar. Common
 209 organisms that can be cultured on PDB are molds such as *Aspergillus niger* and *Rhizopus azzahra*. All
 210 synthesized compounds were dissolved in water or methanol basis on their solubility for making the
 211 concentration 1000 mM/L. The 100 µL solution of ILs were taken in Petri-plate. The Media of PDB was
 212 dispersed and solidified. A well of 5 mm were made in the middle of Petri-plate using cork-borer and the
 213 fungal lead were place there. The plates were then kept in incubator for 96 h at 37°C. After 3 days, the
 214 fungal growth in presence of ILs, were measured and analyzed.

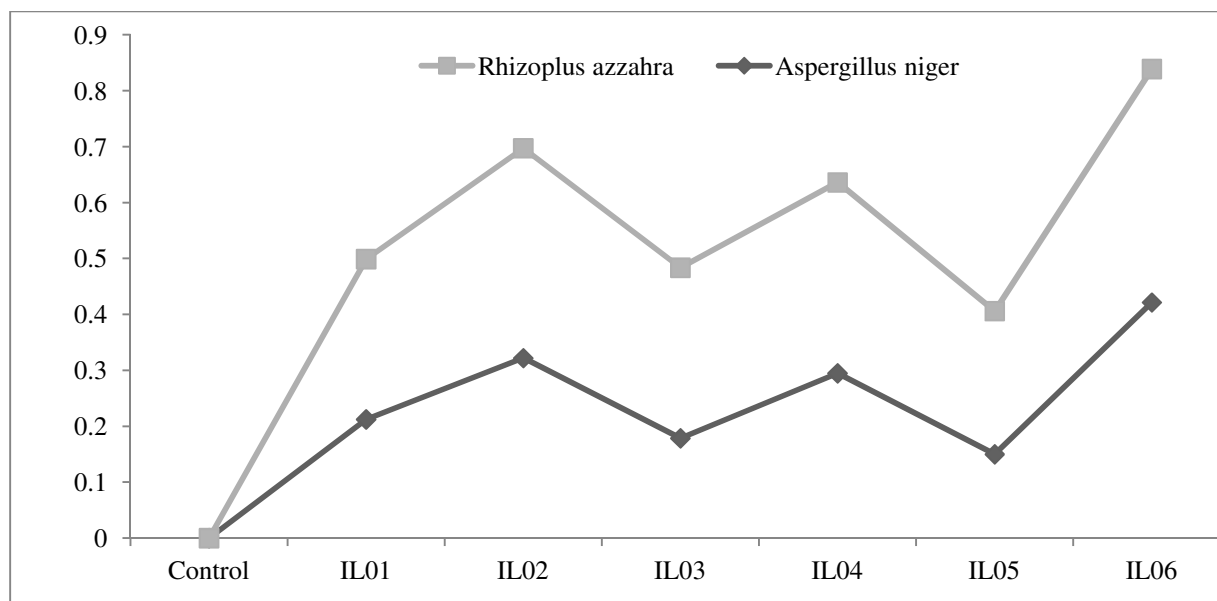
215 The antifungal test was completed and calculated the growth percentage compared with the control
 216 where the growth of control is 100% percent. The growth percentage is deduced as the following
 217 equation:

218
$$\%Growth = \frac{\text{Growth of fungus with IL solution}}{\text{Growth of fungus without IL solution as control}} \times 100$$

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Table no 04: Result and data for antifungal test.

Chemicals tested	Zone of growth (in mm)		Percentage of Growth		Percentage of Inhabitation	
	<i>Aspergillus niger</i>	<i>Rhizopus azzahra</i>	<i>Aspergillus niger</i>	<i>Rhizopus azzahra</i>	<i>Aspergillus niger</i>	<i>Rhizopus azzahra</i>
Control	28 mm	41 mm	100.0%	100.0%	--	--
IL01	20.5±1	29.2±1.0	78.57%	71.34%	21.25%	28.66%
IL02	19.0±1.0	25.2±1.0	67.80%	62.43%	32.20%	37.53%
IL03	23.0±1.0	28.5±1.0	82.14%	69.51%	17.86%	30.49%
IL04	19.7±1.0	27.0±1.0	70.53%	65.85%	29.47%	34.15%
IL05	24.0±1.0	30.5±1.0	85.00%	74.39%	15.00%	25.61%
IL06	16.2±1.0	24.0±1.0	57.85%	58.53%	42.15%	41.75%



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Figure 3: Comparative study of 1^o, 2^o and 3^o ammonium based ILs

234 From the above data, it is found that, the antifungal activity against *Aspergillus niger* and *Rhizopus*
235 *azzahra* is similar relation as like bacteria. The increasing order of antifungal activity is as of 1^o < 2^o < 3^o
236 ammonium based ILs. The anions have an effect on antimicrobial toxicity where acetate has higher
237 antifungal activity than formate. The antifungal activity of *Rhizopus azzahra* is higher than *Aspergillus*
238 *niger*.

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6. Conclusion

241 The synthesized ILs was characterized by Fourier Transform Infrared Spectroscopy (FT-IR), Nuclear
242 Magnetic Resonance (¹H-NMR and ¹³C NMR) Spectroscopy and elemental analysis. The synthesized
243 quaternary ammonium formate and acetate based ILs is the room temperature ionic liquids. Using
244 HyperchemR computer programming, the HOMO-LUMO gap of IL01 to IL06 is 11.677621, 10.968487,
245 11.353785, 11.196557, 4.813104 and 7.269402 respectively that indicates the low reactivity of ILs. The
246 low reactivity of ILs is the good agreements to use as solvents in chemical process. To safe use in
247 chemical industries, the toxicity profile is the most imperative key point of chemical substance. To
248 evaluate the antimicrobial toxicity profiles, bacteria is the first choice due to have short life. Antibacterial
249 screening was conducted using the well-diffusion technique with ILs solutions of three different
250 concentrations. The results were then used to select the initial higher concentration for dilution method, to
251 obtain the concentrations as 1000, 500 and 100 mM/L and the antifungal activity was tested against two
252 phytopathogenic fungi, such as, *Aspergillus niger* and *Rhizopus azzahra* using well diffusion
253 susceptibility test. All of synthesized ILs showed poor inhibition as potential antimicrobial agents, can
254 proceed to future study. It also shows the information about the comparative study of primary, secondary
255 and tertiary based ILs. They have very near toxicity.

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