## IONIC LIQUIDS FURTHER UNCOILED

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## **Critical Expert Overviews**

Edited by

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# **COIL CONFERENCES**

COIL-1	Salzburg	Austria	2005
COIL-2	Yokohama	Japan	2007
COIL-3	Cairns	Australia	2009
COIL-4	Washington	USA	2011
COIL-5	Algarve	Portugal	2013
COIL-6	Jeju Island	Korea	2015
COIL-7	Ottawa*	Canada	2017
COIL-8	Belfast*	UK	2019

<sup>\*</sup> Precise location still to be confirmed.

### **PREFACE**

This is the second of three volumes of critical overviews of the key areas of ionic liquid chemistry. The first volume is entitled *Ionic Liquids UnCOILed* (Wiley 2013), the current volume is *Ionic Liquids Further UnCOILed*, and the final volume, called *Ionic Liquids Completely UnCOILed*, will be published later this year. The history and rationale behind this trilogy was explained in the preface to the first volume, and so will not be repeated here.

Instead, we will use this space to expand on the subtitle, constant for all three volumes: Critical Expert Overviews.

### critical, adjective

- 1. Involving or exercising careful judgement or judicious evaluation
- 2. Of decisive importance in relation to an issue; decisive, crucial

*Critical* has two, rather different, meanings—both are implied in the subtitle of this book. These reviews are both decisively important *and* written by top world experts (hence the second adjective), exercising the judicious evaluation that they are uniquely qualified to do.

### overview, noun

- A general survey; a comprehensive review of facts or ideas; a concise statement or outline of a subject. Also: a broad or overall view of a subject.
- 2. A view from above.

This book includes eleven critical expert overviews of differing aspects of ionic liquids. We look forward to the response of our readers (we can be contacted at quill@qub.ac.uk). It is our view that, in the second decade of the 21st century, reviews that merely regurgitate a list of all papers on a topic, giving a few lines or a paragraph (often the abstract!) to each one, have had their day—five minutes with an online search engine will provide that information. Such reviews belong with the slide rule, the fax machine, and the printed journal—valuable in their day, but of little value now. The value of a review lies in the expertise and insight of the reviewer—and their willingness to share it with the reader. It takes moral courage to say "the work of [. . .] is irreproducible,

#### x PREFACE

or of poor quality, or that the conclusions are not valid," but in a field expanding at the prestigious rate of ionic liquids, it is essential to have this honest feedback. Otherwise, errors are propagated. Papers still appear using hexafluorophosphate or tetrafluoroborate ionic liquids for synthetic or catalytic chemistry, and calculations on "ion pairs" are still being used to rationalise liquid state properties! We trust this volume, containing eleven excellently perceptive reviews, will help guide and secure the future of ionic liquids.

NATALIA V. PLECHKOVA KENNETH R. SEDDON

### **ACKNOWLEDGEMENTS**

This volume is a collaborative effort. We, the editors, have our names emblazoned on the cover, but the book would not exist in its present form without support from many people. Firstly, we thank our authors for producing such splendid, critical chapters, and for their open responses to the reviewers' comments and to editorial suggestions. We are also indebted to our team of expert reviewers, whose comments on the individual chapters were challenging and thought provoking, and to Ian Gibson for producing the central image on the front cover. The backing from the team at Wiley, led by Dr. Arza Seidel, has been fully appreciated—it is always a joy to work with such a professional group of people. Finally, this book would never have been published without the unfailing, enthusiastic support from Deborah Poland and Sinead McCullough, whose patience and endurance never cease to amaze us.

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### **ABBREVIATIONS**

### **IONIC LIQUIDS**

GNCS guanidinium thiocyanate

GRTIL gemini room temperature ionic liquid [HI-AA] hydrophobic derivatised amino acid

IL ionic liquid

poly(GRTIL) polymerised gemini room temperature ionic liquid

poly(RTIL) polymerised room temperature ionic liquid [PSpy]<sub>3</sub>[PW] [1-(3-sulfonic acid)propylpyridinium]<sub>3</sub>[PW<sub>12</sub>O<sub>40</sub>]·

 $2H_2O$ 

RTIL room temperature ionic liquid

### CATIONS

[(allyl)mim]+ 1-allyl-3-methylimidazolium 1,3-dialkylimidazolium  $[1-C_m-3-C_n im]^+$  $[C_2 im]^+$ 1-ethylimidazolium  $[C_1 mim]^+$ 1,3-dimethylimidazolium 1-ethyl-3-methylimidazolium  $[C_2 mim]^+$ 1-propyl-3-methylimidazolium  $[C_3mim]^+$  $[^{i}C_{3}mim]^{+}$ 1-isopropyl-3-methylimidazolium 1-butyl-3-methylimidazolium  $[C_4mim]^+$  $[i-C_4mim]^+$ 1-isobutyl-3-methylimidazolium 1-secbutyl-3-methylimidazolium  $[s-C_4mim]^+$  $[^{t}C_{4}mim]^{+}$ 1-tertbutyl-3-methylimidazolium  $[C_5 mim]^+$ 1-pentyl-3-methylimidazolium  $[C_6 mim]^+$ 1-hexyl-3-methylimidazolium 1-heptyl-3-methylimidazolium  $[C_7 mim]^+$ 1-octyl-3-methylimidazolium  $[C_8 mim]^+$  $[C_9mim]^+$ 1-nonyl-3-methylimidazolium 1-decyl-3-methylimidazolium  $[C_{10}mim]^+$  $[C_{11}mim]^+$ 1-undecyl-3-methylimidazolium  $[C_{12}mim]^+$ 1-dodecyl-3-methylimidazolium 1-tridecyl-3-methylimidazolium  $[C_{13}mim]^+$  $[C_{14}mim]^+$ 1-tetradecyl-3-methylimidazolium

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$[C_{15}mim]^+$	1-pentadecyl-3-methylimidazolium
$[C_{16}mim]^+$	1-hexadecyl-3-methylimidazolium
$[C_{17}mim]^+$	1-heptadecyl-3-methylimidazolium
$[C_{18}mim]^+$	1-octadecyl-3-methylimidazolium
$[C_n \min]^+$	1-alkyl-3-methylimidazolium
$[C_1C_1mim]^+$	1,2,3-trimethylimidazolium
$[C_2C_1mim]^+$	1-ethyl-2,3-dimethylimidazolium
$[C_3C_1mim]^+$	1-propyl-2,3-dimethylimidazolium
$[C_8C_3im]^+$	1-octyl-3-propylimidazolium
$[C_{12}C_{12}im]^+$	1,3-bis(dodecyl)imidazolium

 $[C_1OC_2mim]^+$  1-(2-methoxyethyl)-3-methyl-3H-imidazolium

 $\begin{array}{ll} [C_4 dmim]^+ & \text{1-butyl-2,3-dimethylimidazolium} \\ [C_4 C_1 mim]^+ & \text{1-butyl-2,3-dimethylimidazolium} \end{array}$ 

 $[C_6C_{701}im]^+$  1-hexyl-3-(heptyloxymethyl)imidazolium  $[C_2F_3mim]^+$  1-trifluoroethyl-3-methylimidazolium

 $[C_4 \text{vim}]^+$  3-butyl-1-vinylimidazolium

 $[D_{mvim}]^+$  1,2-dimethyl-3-(4-vinylbenzyl)imidazolium

 $[C_2 mmor]^+$  1-ethyl-1-methylmorpholinium

 $[C_4py]^+$  1-butylpyridinium

 $\begin{bmatrix} C_4 m_\beta py \end{bmatrix}^+ & 1\text{-butyl-3-methylpyridinium} \\ [C_4 m_\gamma py]^+ & 1\text{-butyl-4-methylpyridinium} \\ [C_4 mpyr]^+ & 1\text{-butyl-1-methylpyrrolidinium} \\ [C_6 (dma)_\gamma py]^+ & 1\text{-hexyl-4-dimethylaminopyridinium} \\ [C_1 C_3 pip]^+ & 1\text{-methyl-1-propylpiperidinium} \\ [C_2 C_6 pip]^+ & 1\text{-ethyl-1-hexylpiperidinium}$ 

 $[C_8quin]^+$  1-octylquinolinium

[DMPhim]<sup>+</sup> 1,3-dimethyl-2-phenylimidazolium

 $\begin{array}{ll} [EtNH_3]^+ & ethylammonium \\ [Hmim]^+ & 1\text{-methylimidazolium} \\ [H_2NC_2H_4py]^+ & 1\text{-}(1\text{-aminoethyl})\text{-pyridinium} \end{array}$ 

 $[H_2NC_3H_6mim]^+$  1-(3-aminopropyl)-3-methylimidazolium

 $[N_{1 \ 1 \ 1 \ 2OH}]^+$  cholinium

 $[N_{1122OH}]^+$  ethyl(2-hydroxyethyl)dimethylammonium

 $\begin{array}{ll} \left[N_{1\,1\,1\,4}\right]^{+} & trimethylbutylammonium \\ \left[N_{1\,4\,4\,4}\right]^{+} & methyltributylammonium \\ \left[N_{1\,8\,8\,8}\right]^{+} & methyltrioctylammonium \\ \left[N_{4\,4\,4\,4}\right]^{+} & tetrabutylammonium \end{array}$ 

 $[N_{6\,6\,6\,14}]^+ \hspace{1cm} trihexyl(tetradecyl)ammonium$ 

[NR<sub>3</sub>H]<sup>+</sup> trialkylammonium

 $\begin{array}{ll} \left[P_{2\,2\,2(101)}\right]^{+} & triethyl(methoxymethyl)phosphonium \\ \left[P_{4\,4\,4\,3a}\right]^{+} & (3\text{-aminopropyl})tributylphosphonium \\ \left[P_{6\,6\,6\,14}\right]^{+} & trihexyl(tetradecyl)phosphonium \\ \left[P_{8\,8\,8\,14}\right]^{+} & tetradecyl(trioctyl)phosphonium \end{array}$ 

 $[P_n mim]^+$  polymerisable 1-methylimidazolium

[PhCH<sub>2</sub>eim]<sup>+</sup> 1-benzyl-2-ethylimidazolium

 $\begin{array}{ll} [pyH]^+ & pyridinium \\ [S_{2\,2\,2}]^+ & triethylsulfonium \end{array}$ 

### ANIONS

[Ala]<sup>-</sup> alaninate  $[\beta Ala]^ \beta$ -alaninate

[Al(hfip)<sub>4</sub>]<sup>-</sup> tetra(hexafluoroisopropoxy)aluminate(III)

[Arg] arginate
[Asn] asparaginate
[Asp] asparatinate

[BBB] bis[1,2-benzenediolato(2-)-*O*,*O*']borate

 $\begin{array}{lll} [C_1CO_2]^- & \text{ethanoate} \\ [C_1SO_4]^-, [O_3SOC_1]^- & \text{methyl sulfate} \\ [C_8SO_4]^-, [O_3SOC_8]^- & \text{octyl sulfate} \\ [C_nSO_4]^- & \text{alkyl sulfate} \end{array}$ 

 $[(C_n)(C_m)SO_4]^-$  asymmetrical dialkyl sulfate  $[(C_n)_2SO_4]^-$  symmetrical dialkyl sulfate

[CTf<sub>3</sub>]<sup>-</sup> tris{(trifluoromethyl)sulfonyl}methanide

[Cys]<sup>-</sup> cysteinate

[FAP] tris(perfluoroalkyl)trifluorophosphate

glutaminate [Gln] [Glu]glutamate glycinate anion [Gly] [His] histidinate [Ile] isoleucinate [lac] lactate [Leu] leucinate [Lys] lysinate [Met] methionate [Nle] norleucinate

 $[NPf_2]^-, [BETI]^- \\ bis\{(pentafluoroethyl)sulfonyl\} amide \\ bis\{(trifluoromethyl)sulfonyl\} amide \\ bis\{(trifluoromethyl)sulfonyl] amide \\ bis\{(trifluoromethyl)sulfonyl]$ 

 $\begin{array}{ll} [O_2CC_1]^- & \text{ethanoate} \\ [O_3SOC_2]^-, [O_3SOC_2]^- & \text{ethylsulfate} \end{array}$ 

[OMs]^methanesulfonate (mesylate)[ONf]^perfluorobutylsulfonate[OTf]^trifluoromethanesulfonate

[OTs]<sup>-</sup> 4-toluenesulfonate, [4-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>SO<sub>3</sub>]<sup>-</sup> (tosylate)

[Phe] phenylalaninate [Pro] prolinate [Ser] serinate

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[Suc] succinate

[tfpb] tetrakis(3,5-bis(trifluoromethyl)phenyl)borate

[Thr]<sup>-</sup> threoninate [Tos]<sup>-</sup> tosylate

[Trp] -tryphtophanate[Tyr] -tyrosinate[Val] -valinate

### **TECHNIQUES**

AES Auger electron spectroscopy
AFM atomic force microscopy

AMBER assisted model building with energy refinement

ANN associative neural network

ARXPS angle resolved X-ray photoelectron spectroscopy

ASM Associated-Solution Model

ATR-IR attenuated total reflectance infrared spectroscopy

BPNN back-propagation neural network CADM computer-aided design modelling

CC Cole–Cole model

CCC counter-current chromatography

CD Cole–Davidson model CE capillary electrophoresis

CEC capillary electrochromatography

CHARMM Chemistry at HARvard Molecular Mechanics
COSMO-RS **CO**nductor-like**S**creening**MO**del for Real Solvents

COSY COrrelation SpectroscopY

CPCM conductor-like polarisable continuum model

CPMD Car–Parrinello molecular dynamics

DFT density functional theory

DMH dimethylhexene

DRS dielectric relaxation spectroscopy
DSC differential scanning calorimetry

ECSEM electrochemical scanning electron microscopy
EC-XPS electrochemical X-ray photoelectron spectroscopy

EFM effective fragment potential method

EI electron ionisation

EMD equilibrium molecular dynamics

EOF electro-osmotic flow

EPSR empirical potential structure refinement

ES electrospray mass spectrometry

ESI–MS electrospray ionisation mass spectrometry EXAFS extended X-ray absorption fine structure

FAB fast atom bombardment FIR far-infrared spectroscopy FMO fragment molecular orbital method FTIR Fourier transform infrared spectroscopy

GAMESS general atomic and molecular electronic structure

system

GC gas chromatography

GGA generalized gradient approximations

GLC gas-liquid chromatography GSC gas-solid chromatography

HM heuristic method

HPLC high-performance liquid chromatography

HREELS high-resolution electron energy loss spectroscopy

IGC inverse gas chromatography

IR infrared spectroscopy

IRAS infrared reflection absorption spectroscopy IR-VIS SFG infrared visible sum frequency generation

ISS ion scattering spectroscopy

L-SIMS liquid secondary ion mass spectrometry
MAES metastable atom electron spectroscopy
MALDI matrix-assisted laser desorption
MBSS molecular beam surface scattering

MC Monte Carlo

MD molecular dynamics

MIES metastable impact electron spectroscopy

MLR multi-linear regression
MM molecular mechanics
MS mass spectrometry

NEMD non-equilibrium molecular dynamics

NMR nuclear magnetic resonance

NR neutron reflectivity
NRTL non-random two liquid

OPLS optimized potentials for liquid simulations

PCM polarisable continuum model
PDA photodiode array detection
PES photoelectron spectroscopy
PGSE-NMR pulsed-gradient spin-echo
PPR projection pursuit regression

QM quantum mechanics

QSAR quantitative structure–activity relationship
QSPR quantitative structure–property relationship
RAIRS reflection absorption infrared spectroscopy

RI refractive index

RNEMD reverse non-equilibrium molecular dynamics

RNN recursive neural network

RP-HPLC reverse phase high-performance liquid

chromatography

RST regular solution theory

### xx ABBREVIATIONS

SANS small-angle neutron scattering SEM scanning electron microscopy SFA surfaces forces apparatus

SFC supercritical fluid chromatography

SFG sum frequency generation

SFM systematic fragmentation method
SIMS secondary ion mass spectrometry
soft-SAFT soft statistical associating fluid theory
STM scanning tunnelling microscopy

SVN support vector network

TEM tunnelling electron microscopy TGA thermogravimetric analysis

THz-TDS terahertz time-domain spectroscopy

TLC thin layer chromatography

tPC-PSAFT truncated perturbed chain polar statistical

associating fluid theory

TPD temperature programmed desorption

UHV ultra-high vacuum

UNIFAC UNIversal Functional Activity Coefficient

UNIQUAC UNIversal QUAsiChemical

UPLC ultra-pressure liquid chromatography
UPS ultraviolet photoelectron spectroscopy

UV ultraviolet

UV-Vis ultraviolet-visible

XPS X-ray photoelectron spectroscopy

XRD X-ray powder diffraction

XRR X-ray reflectivity

### MISCELLANEOUS

 $m \mathring{A}$  1  $m \mathring{A}ngstrom = 10^{-10}~m$  ACS American Chemical Society

ATMS acetyltrimethylsilane ATPS aqueous two-phase system

BASF<sup>TM</sup> Badische Anilin- und Soda-Fabrik

BASIL Biphasic Acid Scavenging utilizing Ionic Liquids

BE binding energy

BILM bulk ionic liquid membrane
BNL Brookhaven National Laboratory

b.pt. boiling point

BSA bovine serum albumin
BT benzothiophene
calc. calculated

CB Cibacron Blue 3GA

CCD charge coupled device

CE crown ether

CEES 2-chloroethyl ethyl sulphide

CFC MC "continuous fractional component" Monte Carlo

CLM charge lever momentum
CMC critical micelle concentration

CMPO octyl(phenyl)-*N*,*N*-diisobutylcarbamoylmethylphosp

hine oxide

[C<sub>n</sub>MeSO<sub>4</sub>] alkyl methyl sulfate CNTs carbon nanotubes

COIL Congress on Ionic Liquids
CPU central processing unit
CWAs chemical warfare agents

d doublet (NMR)

D°<sub>298</sub> bond energy at 298 K
2D two-dimensional
3D three-dimensional
DBT dibenzothiophene
DC direct current

DC18C6 dicyclohexyl-18-crown-6 DF Debye and Falkenhagen

DH Debye–Hückel DIIPA diisopropylamine

4,6-DMDBT 4,6-dimethyldibenzothiophene

DMF dimethylmethanamide (dimethylformamide)

DNA deoxyribonucleic acid

2DOM two-dimensional ordered macroporous 3DOM three-dimensional ordered macroporous

DOS density of states
DPC diphenylcarbonate
DRA drag-reducing agent
DSSC dye-sensitised solar cell

E enrichment

EDC extractive distillation column EE expanded ensemble approach

EOR enhanced oil recovery EoS equation of state

EPA Environmental Protection Agency
EPSR empirical potential structure refinement

eq. equivalent

FCC fluid catalytic cracking
FFT fast Fourier transform
FIB focussed ion beam
FSE full-scale error

ft foot

#### xxii ABBREVIATIONS

GDDI generalised distributed data interface

GEMC Gibbs ensemble Monte Carlo

HDS hydrodesulfurisation

HEMA 2-(hydroxyethyl) methacrylate HOMO highest occupied molecular orbital HOPG highly oriented pyrolytic graphite

HV high vacuum

IgG Immunoglobulin G
IPBE ion-pair binding energy

IPE Institute of Process Engineering, Chinese Academy

of Sciences, Beijing

ITO indium–tin oxide

IUPAC International Union of Pure and Applied

Chemistry

J coupling constant (NMR)
KWW Kohlrausch–Williams–Watts
LCEP lower critical end point

LCST lower critical separation temperature
LEAF Laser-Electron Accelerator Facility
LF-EoS lattice-fluid model equation of state

LLE liquid-liquid equilibria
LMOG low molecular weight gelator

LUMO lowest unoccupied molecular orbital

m multiplet (NMR)
M molar concentration
MBI 1-methylbenzimidazole
MCH methylcyclohexane

MDEA methyl diethanolamine; bis(2-hydroxyethyl)

methylamine

MEA monoethanolamine; 2-aminoethanol
MFC minimal fungicidal concentrations
MIC minimal inhibitory concentrations

MMM mixed matrix membrane

MNDO modified neglect of differential overlap

m.pt. melting point

MSD mean square displacement

3-MT 3-methylthiophene MW molecular weight

MWCNTs multi-walled carbon nanotubes

*m/z* mass-to-charge ratio NBB 1-butylbenzimidazole

NCA N-carboxyamino acid anhydride
NE equation Nernst-Einstein equation
NES New Entrepreneur Scholarship

NFM *N*-formylmorpholine

NIP neutral ion pair NIT neutral ion triplet NMP *N*-methylpyrrolidone NOE nuclear Overhauser effect NRTL non-random two liquid

NRTL-SAC non-random two liquid segmented activity

coefficients

OKE optical Kerr effect

pressure

PAO polyalphaolefin **PDMS** polydimethoxysilane

PEDOT poly(3,4-ethylenedioxythiophene)

poly(ethyleneglycol) **PEG** 

**PEM** polymer-electrolyte membrane

poly(ethylene-2,6-naphthalene decarboxylate) PEN

**PES** polyethersulfone

 $-\log_{10}([H^+])$ ; a measure of the acidity of a solution рН

PID proportional integral derivative

 $\mathfrak{p}K_{\mathtt{h}}$  $-\log_{10}(K_{\rm b})$ 

PPDD polypyridylpendant poly(amidoamine) dendritic

derivative

Peng-Robinson equation of state (PR)-EoS

PS polystyrene

**PSE** process systems engineering

1 pound per square inch = 6894.75729 Pa psi

phase transfer catalyst PTC poly(tetrafluoroethylene) PTFE

pressure-temperature composition PTx

bond length

rotating disc contactor RDC

Registration, Evaluation, Authorisation and REACH

restriction of CHemical substances

(RK) EoS Redlich-Kwong equation of state root mean square deviation **RMSD** 

RT room temperature

singlet (NMR) S

S entropy

supercritical carbon dioxide scCO<sub>2</sub> sodium dodecyl sulphate SDS

SED Stokes-Einstein-Debye equation

solvent-to-feed ratio S/F

supported ionic liquid membrane SILM SILP supported ionic liquid phase solid liquid equilibrium SLE SLM supported liquid membrane

### xxiv ABBREVIATIONS

t triplet (NMR)
TBP 4-(t-butyl)pyridine

TCEP 1,2,3-tris(2-cyanoethoxy)propane

TEA triethylamine

TEGDA tetra(ethyleneglycol) diacrylate

THF tetrahydrofuran

TIC toxic industrial chemical

TMB trimethylborate
TMP trimethylpentene
TOF time-of-flight

UCEP upper critical end point

UCST upper critical solution temperature

UHV ultra-high vacuum

VFT Vogel-Fulcher-Tammann equations

VLE vapour-liquid equilibria

VLLE vapour–liquid–liquid equilibria VOCs volatile organic compounds

v/vvolume for volumew/wweight for weightwt%weight pe reentXmolar fraction $\gamma$ surface tension

 $\delta$  chemical shift in NMR