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2 (WEEE) using unconventional solvents based on ionic liquids

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Recovery of metals from waste electrical and electronic equipment

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10 Abstract

Waste electrical and electronic equipment (WEEE) is the fastest growing type of waste globally and is 11 12 an important challenge due to its heterogeneity, intrinsic toxicity and potential environmental impact. 13 However, WEEE also represents an important secondary source of critical and valuable elements and 14 must be viewed as a resource within the context of a circular economy. The use of ionic liquids (ILs) to 15 recover metals from WEEE is reviewed with a special emphasis on the use of ILs in leaching and solvent 16 extraction processes. The advantages of ILs as a reaction medium compared to existing conventional 17 solvents lies in their tuneability and potential environmental benefits. Many studies have used ILs for 18 the selective extraction and purification of metals from WEEE, particularly from waste fluorescent lamp 19 phosphor and batteries containing rare-earth elements with promising results. However, significant 20 barriers remain to the commercial use of ILs for treating WEEE and key barriers to progress are 21 identified. In particular more focus is required to improve upstream collection and separation of WEEE, 22 as this would greatly increase the potential to use selective, more efficient, ILs for metal recovery. 23

24 25

26 **1.** Introduction

As organisations are looking for ways to mitigate the effects of human activity on the environment, recent global trends point towards a shift from the historically prevailing linear economic model, whereby goods are manufactured, consumed and disposed, to a circular economic model promoting reuse and recycling. The primary tenet of the circular economy is the concept of value recovery: all waste possesses an inherent value, which if properly managed can be extracted. In this new context of global environmental remediation, sustainable, effective and affordable waste management is an issue of critical importance.

Keywords: Critical materials; Ionic liquids; WEEE; Solvent extraction; Leaching; Hydrometallurgy

Waste electrical and electronic equipment (WEEE) is the fastest growing waste stream globally and is an important challenge due to inherent value, heterogeneity and potential environmental impact (Ye et al., 2009; Ogondo et al., 2011). Essential constituents of many EEE products, in particular

37 information technology (IT) and communication equipment, are base metals (Fe, Cu, Al, etc.) as well 38 as important concentrations of precious metals (Au, Ag, Pd) and specialty metals (In, Sb, Te, Ta, rare 39 earth elements (REE), etc.) all highly interlinked in a complex composition (Hagulken, 2006). Many 40 WEEE streams or components of WEEE contain higher concentrations of valuable metals than their 41 primary ore and as such represent an opportunity as secondary 'ores'. This is especially true as metals 42 are finite commodities with a limited availability for human use. Mining rates of most metals are at a 43 historical maximum whilst the average ore grade has decreased over the last hundred years for both 44 base and precious metals (Mudd, 2010). The reduced availability of minable deposits and the 45 concentration of these deposits in a limited number of countries results in an imbalance between supply 46 and demand, making the current practice unsustainable in the long term. In addition, there is a growing 47 legislative pressure on the proper handling and recycling of WEEE, with the collection and recycling 48 targets recently increased as part of the EU WEEE Directive (2012/19/EU).

49 Current processes to recover value from WEEE can be complex and involve high water and energy 50 consumption and/or the production of large quantities of hazardous effluents (Cui & Zhang, 2008, 51 Tuncuk et al., 2012). WEEE recycling occurs in three major stages: (1) disassembly, (2) upgrading 52 using mechanical and/or metallurgical processing to increase the desirable materials content of the 53 waste, and (3) refining by purification of the upgraded material by metallurgical processing (Cui & 54 Forssberg, 2003). A schematic flow-diagram of the major stages in WEEE processing is given in Figure 55 1.

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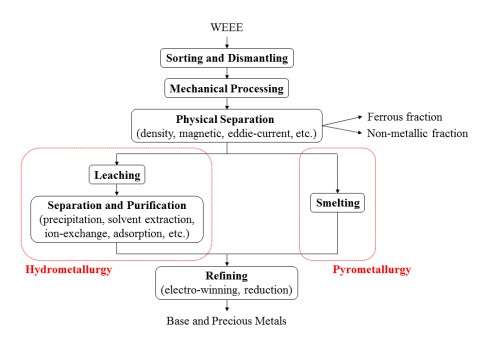




Figure 1. Simplified schematic of the major stages of WEEE recycling.

60 Although both hydro- and pyro- metallurgy have advantages, hydrometallurgy is often favoured 61 over traditional pyrometallurgy by industry due to an ability to recover metals from a variety of 62 challenging sources, including low grade ores, mixed metal ores and secondary sources such as WEEE. 63 Furthermore, hydrometallurgy allows the use of low process temperatures and recycling of reagents 64 (Wilson et al., 2014). Conventional industrial processing of WEEE requires extraction and separation 65 techniques that use aggressive acid or alkali digestion, can involve multi-stage separation procedures, 66 produce large aqueous waste streams which require further treatment, and in the solvent extraction 67 process, make use of solvents which are potentially flammable, volatile and toxic.

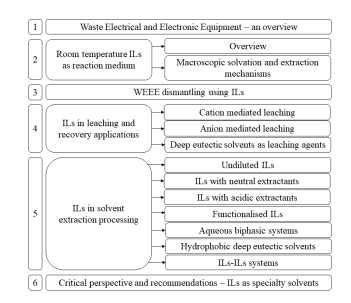
68 Ionic liquids (ILs) are solvents composed of ions with properties intermediate between molten 69 salts and organic solvents and are attracting significant interest due to potential environmental benefits 70 and tailorability compared to conventional solvents (Earle & Seddon, 2000; Rogers & Seddon, 2003). 71 They have been applied in a number of industrial processes and products (Plechkova & Seddon, 2008) 72 and if properly selected, ILs could offer an environmentally friendly alternative to current 73 hydrometallurgical WEEE recycling processes by minimizing waste generation and acid consumption 74 whilst maintaining selective leaching. Since the proposal of ILs as environmentally friendly media 75 compared to volatile organic solvents for liquid-liquid extraction (Huddleston et al., 1998), the use of 76 ILs for solvent extraction remains the most widely researched field in IL mediated metal processing. 77 Solvent extraction (or liquid-liquid extraction) is the favored option for metal separation and involves 78 the preferential two-phase distribution of a solute between mutually immiscible liquid phases usually 79 consisting of an aqueous acidic solution and an aliphatic organic solvent.

80 Due to their versatility and growing popularity, the use of ILs for metal separation has been 81 reviewed by previous authors (Dietz et al., 2006; Lee, 2006; Han & Armstrong, 2007; Abbott et al., 82 2011; Tian et al., 2010; Hiramaya, 2011; Stojanovic & Keppler, 2012; Sun et al., 2012; Kolaric, 2013) 83 whilst the use of the immobilized IL in metal extraction was reviewed elsewhere (Stojanovic & Keppler, 84 2012; Vidal et al., 2012). The use of ILs for the electro-deposition of metals was previously reviewed 85 by the research groups of Abbott et al. (2006; 2011; 2013), Smith et al. (2014) and Endres et al. (2002; 86 2008). Furthermore, a detailed description of the fundamental governing principles dictating metal 87 extraction to IL (Watanabe et al., 2014; Janssen et al., 2015; Katsuta et al., 2016; Papaiconomou et al., 88 2016) as well as specific applications for certain classes of ILs such as acidic ILs (Amarasekara, 2016) 89 or temperature responsive ILs (Qiao et al., 2017) are also available. Nevertheless, there is currently no 90 review focusing on the use of ILs for the recovery of value from WEEE despite the increasing number 91 of publications on this field in the last five years.

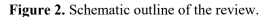
This review is divided into six sections, with the schematic review outline presented in Figure After an introduction of the current WEEE recycling and ILs, the discussion follows the typical sequence in WEEE processing, starting with dismantling, followed by leaching and finishing with solvent extraction. The first provides a brief overview of metal recycling from WEEE highlighting the metals of critical interest for recovery and the challenges to date. The second addresses the general

97 properties of ILs as reaction medium with an emphasis on metal-ILs extraction mechanisms. The third 98 and fourth sections present the application of IL-derived solvents to the solvato-metallurgical 99 processing of electronic waste, namely their use for the dismantling and leaching of WEEE. In the case 100 where the reviewed literature presents a recovery methodology, this will be presented in conjunction with its leaching to promote the application of a one solution for the elective leaching and recovery of 101 102 metals from WEEE. The fifth section reviews the use of unconventional solvents based on ILs in solvent 103 extraction processes relevant to WEEE recycling. This section represents the largest body of reviewed 104 works and covers both the use of traditional IL solvent extraction processes as well as non-conventional systems such as hydrophobic deep eutectic solvents (DES) and aqueous biphasic systems (ABS). 105 Finally, a critical assessment on the use of ILs for metal extraction from WEEE is conducted with 106 107 guidelines provided based on the reviewed literature to maximise its beneficial application. The focus 108 is on recovery of metals from WEEE rather than plastic or glass components, with emphasis on ILs used in leaching and solvent extraction. In this review, the ILs are discussed according to their 109 application. As such, one IL used both in leaching and in solvent extraction applications will be 110 111 presented in two different section although care is taken to avoid overlap. Although not within the scope 112 of this review, some relevant examples on the application of ILs for the recovery of metals from primary 113 ores are presented to enrich the discussion. Current hydrometallurgical treatment of WEEE borrows 114 heavily from principles developed by the mining industry; innovative techniques developed for the 115 latter can be transferred to the extraction of value from WEEE.

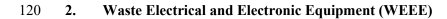
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121 Global WEEE generation is estimated at 20-50 million tonnes per year and is increasing at a 3-122 5 % rate (Ogondo et al., 2011). In 2005, WEEE already represented approximately 8 % of municipal 123 solid waste (Widmer et al., 2005). Annual WEEE production in the European Union (EU) increased 124 from 6.5 Mt in 2004 to 12 Mt in 2015 (Goosey, 2004, Baldé et al., 2015). An important distinction must 125 be made between WEEE and e-waste. Although many authors use the terms interchangeably, Robinson (2009) describes WEEE as "discarded appliances that use electricity" including traditionally non-126 127 electronic goods such as refrigerators and ovens. WEEE designates any wastes that fall under the jurisdiction of the European WEEE Directive (2002/96/EC) as set out in Table 1. E-waste is a sub-128 129 fraction of WEEE that encompasses "waste electronic goods, such as computers, televisions and cell 130 phones". These goods primarily fall under category 3 and 4 of the WEEE Directive.

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Table 1. Classification of WEEE into ten categories according to the European WEEE Directive
 (2002/96/EC). The highlighted rows correspond to the e-waste subsection of WEEE.

No.	Category	Example
1	Large household appliances	Refrigerator, oven, washing machine
2	Small household appliance	Toaster, microwave, vacuum cleaner
3	IT and telecommunications equipment	Mobile phone, computer, computer screen
4	Consumer equipment	Television, radio, speaker
5	Lighting Equipment	LED, gas discharge lamps
6	Electrical and electronic tools	Power drill, electric screw driver
7	Toys, leisure and sports equipment	Treadmill, game console
8	Medical devices	MRI, dialysis
9	Monitoring and control instruments	Thermocouple, smoke detector
10	Automatic dispenser	Vending machine, cash distributor

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Global WEEE is exponentially increasing due to (1) population growth, (2) economic growth and the resulting improvement in living standards, (3) decrease in product lifespan and (4) the increase in everyday commodities containing microprocessors. One consequence of increasing EEE on the market is the blurring of the traditional borders between classical electrical equipment and electronic equipment. Over 98% of all programmable microprocessors are now embedded in commodities that are not associated with computers (Hilty, 2005).

WEEE is a complex, heterogeneous waste stream containing over 1000 different substances (Ye et al., 2009) and is primarily composed of plastics (34.6 wt.%), electronic components (23.5 wt.%) and ferrous metals (15.6 wt.%) (Dimitrakakis et al., 2009). Essential constituents of many EEE products, in particular IT and communication equipment, are base metals (Fe, Cu, Al, etc.), precious metals (Au, Ag, Pd) and speciality metals (In, Sb, Te, Ta, rare earth elements (REEs), etc.) (Hageluken, 2006). Due

146 to their significant metal concentration, EEE production has an important impact on global demand for 147 metals and is an important driver of metal scarcity. Essential constituent of many EEE are metal with 148 reserves often concentrated in a handful of countries creating a potential supply risk. This is exemplified 149 by the substantial increase of most of rare earth oxides (REOs) prices in 2011 due to the implementation 150 during the previous year in China of a REOs export quota policy (Tan et al., 2015). Imbalances between metal supply and demand in conjunction with the importance of a metal for continued economic growth 151 152 has inspired the concept of metal criticality (Graedel et al., 2015). Multiple methodologies have attempted to assess the criticality of materials with widely varying results (Erdmann & Graedel, 2011, 153 154 Bujis et al., 2012).

155 A selection of previous studies assessing raw material criticality is presented in Table 2. These 156 studies primarily focus on the criticality of materials from a European or U.S. perspective. It is important 157 to remark that criticality varies geographically, and certain materials labelled as critical in EU are likely 158 to change if other countries such as China are considered. The most comprehensive studies are those originating from the European Commission (2014) and from Graedel et al. (2012, 2015). These reports 159 include a temporal dimension (assessment of current and future criticality) and estimate material 160 161 criticality based on multiple factors including (but not limited to) economic importance, supply risk, vulnerability to supply restriction, substitutability and environmental impact. Although material 162 criticality is a fluid concept with temporal and geographical variations, certain metal groups appear in 163 most of the presented studies, namely indium, antimony, REE, platinum group metals (PGM), 164 165 magnesium, manganese and niobium. Recovery of these elements is of major importance to minimise 166 their environmental impact and for the continued growth of the EEE industry and green industry.

168	Table 2. Selection of	previous a	studies	assessing raw	material criticality.
		1		0	5

Study	Critical Material
NRC (2008)	In, Mn, Nb, PGM, REE
Morley & Eatherley (2008)	Au, Rh, Pt, Sr, Ag, Sb, Sn
Oko-Institut (2009)	Te, In, Ga, REE, PGM, Ta, Li, Ge, Co
U.S. DoE (2010)	REE (Dy, Nd, Tb, Eu, Y), In
Erdmann & Graedel (2011)	REE, PGM, In, W
Graedel et al. (2012)	Results grouped by mineral family. Eg. for 'copper family': $As = Gd > Ag > Se \approx Cu > Te$
European Commission (2014)	Sb, Be, Co, fluorspar, Ga, Ge, graphite, In, Mg, Nb, PGM, REE, Ta, W
Graedel et al. (2015)	In, As, Tl, Sb, Ag, Se, Au, PGM, Mg, Cr, Mn, Rh, REE

170 In 2007, demand for indium, antimony, tin, copper, silver, cobalt and selenium for use in EEE 171 accounted for an estimated 79.2, 50.0, 32.7, 30.0, 30.0, 19.0 and 17.1% of total production of these 172 metals (Hagelüken & Buchert, 2008). Many WEEE streams and components of WEEE contain greater 173 concentrations of valuable metals than the primary ore. It is estimated that the annual replacement value of the materials found in WEEE disposed of in the UK alone is ~£1.3bn, with £400m either hoarded or 174 landfilled (Haig et al., 2012). The quantity of precious metal in WEEE generated solely in the UK is 175 estimated at about 50 tonnes per year. For example, phosphor powders used in fluorescent lamps 176 contains elevated concentration of REE including Eu. Binnemans et al. (2013) estimate that waste 177 178 fluorescent lamps will contain up to 25,000 tons of REEs by 2020.

179 The primary factors influencing the recyclability of a metal from WEEE are the volume of the waste, 180 the price of the metal and the purity of the element and ease of recovery. For many electrical products, 181 innovation is associated with an increase in the number of elements used. In the 1980's, 11 elements were contained in Intel® printed circuit boards (PCBs), compared to ~60 in the 2000s (Johnson et al., 182 2007). Additional elements are often present at low concentrations but have important specific 183 184 functions. New high-strength permanent magnets or new alloys with enhanced physical and chemical properties are obtained using lanthanide elements (Graedel, 2011; Binnemans et al., 2013). Recovery 185 186 of these metals from WEEE is seldom considered due to the technological and economic challenges 187 involved (Reck & Graedel, 2012). A consequence of the rising number of elements in WEEE is that 188 individual metal recycling rates can be divided into three categories. Recent global estimates of end-of-189 life recycling rates for 60 metals and metalloid (UNEP, 2011) are presented in Figure 3.

1 H Li 11 Na	4 Be 12 Mg			> 2 > 1	0% 5-50° 0-25° 10% %			No No	errou on-Fo oecia recio	errou lty	18	5 B 13 Al	6 C 14 Si	7 N 15 P	8 0 16 S	9 F 17 Cl	2 He 10 Ne 18 Ar
19 K	20 Ca	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr
37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe
55 Cs	56 Ba	*	72 Hf	73 Ta	74 W	75 Re	76 0s	77 Ir	78 Pt	79 Au	80 Hg	81 Tl	82 Pb	83 Bi	84 Po	85 At	86 Rn
87 Fr	88 Ra	**	104 Rf	105 Db	106 Sg	107 Sg	108 Hs		110 Ds	111 Rg	112 Uub	113 Uut	114 Uug	115 Uup	116 Uuh	117 Uus	118 Uuo
* Lan	thanid	es	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
** Ac	tinides	;	89 Ac	90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr

192 Figure 3. Global estimates of end-of-life recycling rates for 60 metals and metalloid in 2008 (adapted 193 from UNEP, 2011).

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195 The first category of metal corresponds to 'base metals' (Fe, Cu, Sn, Ni, Zn, Al and Pb), with end-196 of-life recycling rates (EOL-RRs) at >50 wt.%. These are the main metals in WEEE, are often found in 197 relatively pure elemental form and benefit from mature recycling infrastructure (Khaliq et al., 2014). 198 The second category corresponds to 'precious metals' with EOL-RRs of ~50-60 wt.%. This includes 199 some platinum group metals, Au and Ag, which despite low concentrations have high economic value 200 (UNEP, 2011; LME, 2016). The recycling rates are positively skewed due to the high recycling rates 201 of industrial catalysts and jewellery. This significantly masks lower EOL-RR from WEEE in the EU 202 (Chancerel et al., 2009; Hagelüken, 2012). Platinum group metal recycling rates from WEEE, such as 203 Pd(II) from PCBs were only 5-10 wt.% in 2011. EOL-RRs for other elements from WEEE are restricted 204 by low collection rates due to export of WEEE to countries with primitive treatment technologies and 205 disposal of WEEE in municipal solid waste (Hagelüken, 2012). The final category consists of 'specialty 206 metals' that are characterised by low recycling rate, typically below < 1 wt.% (UNEP, 2011). Many 207 metals in this category are labelled as critical (see Table 2). These are used in low concentrations and 208 extensively mixed making separation often economically unviable (Du & Graedel, 2011; Reck & 209 Graedel, 2012). Recovery of these metals represents a major challenge to achieve a circular economy 210 due to increasing use in EEE and the current linear management system.

211 Frustratingly, a significant portion of most strategic metals in e-waste is lost during the collection, 212 sorting and mechanical separation stages. In turn, this decreases the financial viability of any recovery 213 process through the dilution of precious metal content by commingling with other types of lower-value 214 e-waste. Small high-value e-waste such as mobile phones suffer from low end-of-life recycling rate as 215 they are often either improperly disposed in municipal waste bins or stored forgotten in a drawer. Mechanical treatment options suffer from the 'concentration dilemma', i.e. the recovery yield for a 216 217 specific metal decreases with the rising purity of that metal in the output stream (Hagulken, 2006). The 218 unintended co-separation of minor metals during mechanical pre-treatment unavoidably can lead to 219 substantial losses before it ever reaches any hydrometallurgical processes. A substance flow analysis of 220 a full-scale pre-processing facility shows that 60 wt.% of the Cu reach output fractions from which it 221 may potentially be recovered. This decreases to 25.6 wt.% for Au and Pt and only 11.5 wt.% for Ag 222 (Chancerel et al., 2009).

223 The ILs covered in this review are for the most part limited to the recovery of value from PCBs, 224 waste fluorescent phosphor, nickel metal hydride batteries and permanent magnets. Research has 225 focused on these waste types due to (i) their high inherent metallic value and (ii) their ease of 226 procurement as an 'uncontaminated' waste. These e-wastes have well established collection and sorting processes, with some even separated at source due to legislative pressure. Fluorescent tubes, which contain rare-earth rich phosphor powder, are classified as a hazardous waste in the European Union due to their mercury content. As such, fluorescent lamps benefit from a dedicated collection and separation chain, ensuring a high content of REE, phosphorus and antimony. In contrast, indium and tantalum can be found in high concentration in indium-tin oxide film in flat screen and in capacitors respectively. However as these are small components of a larger waste, their current recycling barriers make the recovery of these metals from large scale WEEE treatment facilities difficult.

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235 **3. Room temperature ILs as reaction medium**

3.1. Overview

Room temperature ILs are a class of solvents consisting of organic cations and organic or inorganic anions with melting points below 100°C (Earle & Seddon, 2000). Due to their unique physical and chemical properties, ILs have generated considerable scientific interest over the last 20 years. In this section, solvation interactions within ILs are discussed prior to a more detailed assessment of ILs for the recovery of metals from WEEE with special emphasis on hydrometallurgical processes. The discussion is limited to second and third generation ILs and will not cover ILs based on chloroaluminate anions.

244 The majority of chemical reactions occur in solution, with the solvent properties influencing its 245 suitability as a reaction medium. ILs are solvents composed exclusively of ions and as such their solvation behaviour differs from that of neutral molecular solvents (hexane, acetone, DMSO, etc) and 246 247 aqueous, or partially ionized, solvents. ILs (generally) consist of large asymmetrical cations and simple 248 coordinating anions. The low degree of packing efficiency, induced by the size and lack of symmetry 249 of the cation, in conjunction with a lower cationic (and occasionally anionic) charge density, caused by 250 delocalization of the partial charge over the whole ion, drastically reduces the lattice energy of the salt resulting in lower melting points (Seddon, 1997). This low melting point distinguishes ILs from 251 252 conventional molten salts like molten sodium chloride ($T_m \ge 801$ °C), which are characterised by a high charge density and packing efficiency. Commonly used cations and anions for ILs and their 253 254 abbreviations are provided in Figure 4. For the sake of clarity and to enable meaningful comparisons 255 between different systems, ILs will be refered according to their structure and not their commercial 256 name. ILs comprised of the other anions listed in Figure 4 display varying degrees of hydrophobicity 257 depending on the cation's alkyl chain length. The hydrophobicity of an IL is of particular relevance for 258 its application as an extraction and separation media. It is important to state that all ILs are hygroscopic,

regardless of their hydrophobicity, and water may make up a significant molar percentage of an IL.

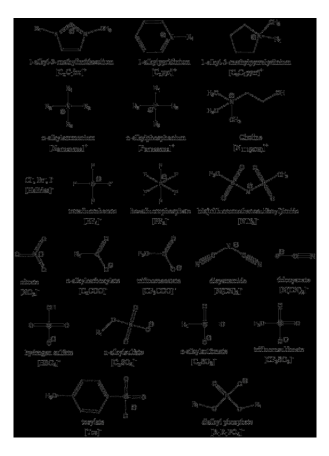


Figure 4. Schematic of selected cations and anions from common ILs and DES and their abbreviations.

Many solvent properties, including hydrophobicity, polarity and viscosity to name a few, are important to an IL's performance and applicability as a reaction medium. In ILs, these 'bulk properties' are altered by either varying the anion or the cation and this has led ILs to be labelled as 'designer solvents' (Freemantle, 1998; Huddleston et al., 2001). Many of these properties have been extensively reviewed elsewhere (Ohno, 2005; Koel et al., 2007; Endres et al., 2008, Freemantle, 2009; Hallet & Welton, 2011) and will not be discussed here.

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271 **3.2. Macroscopic solvation and extraction mechanisms**

In this section, a brief synopsis of the fundamental governing principles dictating metal-IL 272 273 interaction and extraction is given, particularly as no new extraction mechanisms were reported 274 compared to older existing work. For an in-depth analysis, the reader is referred to the review of Janssen 275 et al. (2015). The overall solvation property of a system results from the sum of the possible solute-276 solvent interactions and depends on the cation and anion of the IL, as well as the solute. Charged complexes can be highly soluble in ILs, allowing for a wider range of metal speciation compared to 277 molecular solvent in which generally only neutral complexes are soluble. The ionic nature of ILs results 278 279 in metal extraction mechanisms from an aqueous phase that are not possible in conventional organic solvents such as ion-exchange mechanisms involving ions of the IL, which can lead to the degradation

of the IL. Both charged and neutral metal complexes can be highly soluble in ILs (Jensen et al., 2002).

The most conclusive method for the determination of metallic species in IL is single-crystal XRD analysis (Nockemann et al., 2010). However, due to the low vapour pressure and melting point of ILs, acquisition of suitable crystal structure is often impossible and in-situ techniques are required. These

- are reviewed by Abbott et al. (2008) and will only be listed here. The most widely used technique to
- 286 probe metallic complexes in ILs is extended X-ray absorption fine structure (EXAFS), with certain
- 287 studies also using Raman, UV-vis and FAB-MS spectroscopic techniques.
- 288 When discussing metal-IL interaction, a distinction must be made between positively and negatively 289 charge metal complexes. For positively charged metal complexes, typically the IL anion dictates the 290 chemistry of the system because it interacts with the generally more Lewis acidic metal solute. Anion 291 selection determines the Lewis acidity and coordination ability of an IL, with metal salts being poorly 292 soluble in ILs with strongly Lewis acidic/practically non-coordinating anions. Chloride, nitrate and 293 sulphate, are strongly Lewis basic/strongly coordinating anions whilst the fluorinated anions $[BF_4]^-$, 294 $[PF_6]^-$ and $[NTf_2]^-$ are acidic and only weakly coordinating (Wasserscheid & Welton, 2008). ILs 295 incorporating strongly coordinating anions exhibit significantly greater metal dissolution potential than 296 those with acidic anions (Abbott et al., 2005; Pereiro et al., 2012). Complexation of the metal with the 297 IL anion controls the metal speciation and therefore its reactivity, solubility, redox potential and colour 298 (Abbott et al., 2011). The metal counter-anion also plays an important role in determining solubility in 299 a particular IL. The Hofmeister series (or the lyotropic series) provides a general indication of the 300 hydrophobicity of an anion. In ascending order, it reads: $[SO_4]^{2-} \le Cl^- \le Br^- \le [NO_3]^- \le I^- \le [ClO_4]^- \le I^- \le [NO_3]^- \le I^- \le [ClO_4]^- \le I^- \le I^-$ $[SCN]^{-} \leq [NTf_2]^{-}$ and reflects the charge density and hydration numbers of the different anions (Dupont 301 302 et al., 2015). Metal salts containing hydrophilic anions of high charge density are unlikely to dissolve 303 in poorly coordinating hydrophobic ILs such as $[C_4C_1im][NTf_2]$. The solubility of inorganic salts in ILs 304 with a common anion is greatly improved, both for hydrophilic and hydrophobic anions (Pereiro et al., 305 2012).

306 In many cases, metal ions, including the precious metals present in WEEE, form negatively charged 307 complex species once dissolved in an acidic solution. For instance, Pt(IV) forms various polychloroplatinate(IV) complexes such as $PtCl_5(H_2O)^-$ or $PtCl_6^{2-}$ in aqueous solutions of HCl. Pt(II), 308 309 Pd(II) and Rh(III) form similar polychlorometallate complexes, depending on the concentration of HCl 310 in water. In such cases, ILs are much more efficient at extracting these ions. Precious metals are 311 successfully extracted using undiluted ILs such as [CnC1im][NTf2] and their pyridinium or 312 pyrrolidinium homologues (Papaiconomou et al. 2012; 2013; 2014; 2015; Watanabe et al., 2015). 313 Functionalized ILs, also known as task specific ILs (TSILs), are ILs with an added functional group 314 covalently bonded to either the cation or anion which confer certain desired properties useful for 315 selective metal extraction (Fagnant et al, 2013). Derivatization of the IL cation to incorporate taskspecific functionalities such as metal ligating groups provide an efficient way to maximise metal solvation (Visser et al., 2001). For example, substitution of the cationic alkyl chain in $[C_nC_1im][PF_6]$ with either urea-, thiourea-, and thioether- alkyl groups can result in an increase in the extraction of Cd(II) from aqueous solution by several orders of magnitude (Visser et al., 2002).

320 Major developments in the use of ILs in solvent extraction can be divided into three main categories: 321 ILs used as the diluting hydrophobic phase in conjunction with a traditional extractant; selective 322 extraction of metals through IL anion interaction; and task specific ILs (Abbott et al., 2011). 323 Electroneutrality is the governing principle behind the multiple potential extraction mechanisms in ILs 324 as the net charge of the aqueous and IL phases must remain constant (Janssen et al., 2015). A brief 325 overview of the potential extraction mechanisms for a metal cation (M^{n+}) from an aqueous solution containing an anion (X⁻) to an IL phase (C⁺ - IL cation; A⁻ - IL anion) containing a ligand (L or HL if 326 327 protonated) is described in Table 3. A commonly used category of hydrophobic ILs in solvent extraction 328 are those incorporating the $[NTf_2]^-$ anion. These ILs often require the addition of an extractant to increase the hydrophobicity of the metal complex to facilitate the extraction of a hydrated metal cations 329 330 (Zhao et al., 2005). For example, many studies have reported enhanced extraction efficiency and selectivity for REEs on substitution of conventional organic diluents by hydrophobic ILs (Nakashima 331 et al., 2004; Shimojo et al., 2008; Shen et al., 2011). The structure and abbreviation for the main 332 333 extractants discussed in this work are presented in Figure 5.

Table 3. Multiple extraction schemes for the transfer of metal cations from an aqueous phase to an ILphase.

Extraction mechanism General extraction scheme				
	ILs + extractants			
NI	$M_{aq}^{n+} + nHL_{IL} \rightarrow [M \cdot (L)_n]_{IL} + nH_{aq}^+$	(1)		
Neutral extraction	$M_{aq}^{n+} + mL_{aq} + nX_{aq}^{-} \rightarrow [M \cdot (L)_m \cdot (X)_n]_{IL}$	(2)		
Ion exchange	$M^{n+}_{aq} + mL_{aq} + nC^+_{IL} \rightarrow [M \cdot (L)_m]^{n+}_{IL} + nC^+_{aq}$	(3)		
	$\begin{split} M^{n+}_{aq} + (n+1)HL_{aq} + [C][A]_{IL} &\to [M \cdot (L)_{n+1}]^{IL} \cdots [C]^+_{IL} + \\ (n+1)H^+_{aq} + A^{aq} \end{split}$	(4)		
	$\begin{split} M^{n+}_{aq} + mL_{aq} + (n+1)X^{-}_{aq} + [C][A]_{IL} \rightarrow \\ [M \cdot (X)_{n+1} \cdot (L)_m]^{-}_{IL} \cdots [C]^{+}_{IL} + A^{-}_{aq} \end{split}$	(5)		
	Neat ILs			
Ion pair	$M_{aq}^{n+} + mX_{aq}^{-} + (m-n)C_{aq}^{+} \to [C_{m-n}MX_m]_{IL}$	(6)		

$$M_{aq}^{n+} + mX_{aq}^{-} + (m-n)CA_{IL} \to [C_{m-n}MX_m]_{IL} +$$
(7)

$$(m-n)A_{aq}$$

$$M_{aq}^{n+} + mX_{aq}^{-} + (m-n)A_{IL} \to [MX_m]_{IL} + (m-n)A_{aq}$$
(8)



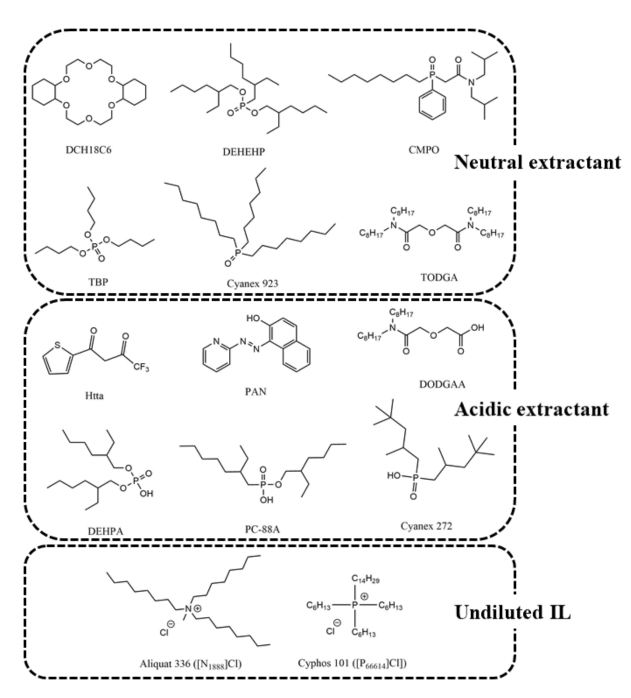


Figure 5. Schematic of selected neutral and acidic extractants and primary undiluted ILs presented inthis work and their abbreviations.

342 These equations do not represent all potential extraction mechanisms in ILs but rather provide an 343 overview of the various extraction schemes possible whilst preserving electroneutrality. The extraction 344 mechanisms in solvent extraction systems with a ligand dissolved in the IL can be broadly divided into 345 two categories: neutral extraction and ion-exchange extraction. Neutral extraction can proceed through 346 the formation of a neutral metal-ligand complex either via deprotonation of the ligand (Ext. 1), 347 formation of a charge neutral complex with aqueous solution anions and extraction of a neutral metal-348 cation/anion ion-pair (Ext. 2). For metal extraction proceeding through an ion exchange mechanism, 349 the IL cation is released to the aqueous phase to compensate for the incoming positive charge of the 350 metal complex (Ext. 3-5). In the case of TSILs, the functionilized component of the IL acts as the ligand 351 (Jensen et al., 2003). It is possible for multiple extraction mechanisms to co-exist within similar solvent 352 extraction systems. The factors determining the relative importance of an extraction scheme include hydrophobicity of the IL cation (Jensen et al., 2002; Dietz et al., 2003) and anion (Jensen et al., 2003; 353 354 Luo et al., 2006) and the hydration energy of the aqueous metal complex (Garvey et al., 2012; 355 Wankowski & Dietz, 2016). The composition of the aqueous phase can also influence the extraction 356 efficiency for a particular solvent extraction. The water miscibility of four hydrophobic ILs was found to increase with addition of the cations $NH_4^+ > K^+ > Na^+ > Li^+ > Ca^{2+} > Mg^{2+}$ and anions $[ClO_4]^- > I^- >$ 357 $[NO_3]^- > Cl^- > F^- > [SO_4]^{2-}$ as predicted by the Hofmeister series (Dupont et al., 2015). 358

The most widely used solvent extraction systems involve use of either ion-exchange extractant 359 classified into acidic (eg. carboxylates, sulfonates, phosphates) and basic extractant (primary to 360 361 quaternary amines), solvating extractant (or neutral extractant) possessing ketone, ether, ester or alcohol 362 functional groups and/or coordinating extractant (or chelating extractant) (Free, 2013). These will be 363 discussed in the following section. Whilst a single extractant can often saturate the coordination sphere 364 of a metallic cation as well as produce a charge neutral and hydrophobic complex, on occasion a 365 combination of two extractants is required. If the resulting distribution ratio is greater for the combined 366 extractant than the sum for the individual extractants employed, the system benefits from a synergistic 367 extraction mechanism (Stepinski et al., 2005). In addition to increasing the extraction efficiencies, synergistic extraction systems can also improve the separation factors between two different metals. 368 369 Such systems have been used effectively in conjunction with ILs, and the presence of two extractants 370 can result in a change of extraction mechanism compared to when the extractants are individually 371 employed (Zhu et al., 2015; Petrova, 2016).

In the case where no extractant is used and the metal ion forms negatively charged complex ions with anions present in the aqueous phase, either originating from the acid used, or added on purpose in the solution, the extraction mechanisms can be discussed in a similar way as that detailed above. Because electroneutrality must be respected, two main possible extraction mechanisms, hydrophobic ion pair formation or anion exchange, occur (Ext. schemes 6 to 8). In the case of the formation of hydrophobic ion pairs, two versions exist in the literature, one using cations from the IL that are in the 378 IL phase, and another one using cations that are dissolved in the aqueous phase (Ext. 6 and 7). 379 Strikingly, even though anion exchange and ion pair formation were discussed as two distinct 380 mechanisms, most recent studies show that these are actually linked together because a hydrophobic 381 IL, as any salt, exhibits a limited, though non negligible, solubility in water. The typical solubility of $[C_8C_1im][NTf_2]$ is around 10⁻³ mol.L⁻¹. In other terms, the solubility product of hydrophobic ILs is not 382 negligible (Chapeaux et al., 2007; Mazan et al., 2014). In an anion exchange mechanism, an anion of 383 384 the IL is transferred to the aqueous phase. Because the solubility of the IL is then exceeded, this 385 necessarily yields the transfer of an IL ion pair initially dissolved in the aqueous phase towards the IL 386 phase. This results as the co-extraction of a metal complex with a cation of the IL, that is, the formation 387 of a hydrophobic ion pair (Ext. 8).

388 In the following sections, the use of ILs at different stages of WEEE recycling is reviewed, with 389 particular emphasis on the use of ILs as solvents for leaching and liquid-liquid extraction processes. 390 The use of IL for the electro-deposition of metals has previously been reviewed by Abbott et al. (2006; 391 2011; 2013; Smith et al., 2014) and Endres et al. (2002; 2008) and will only be briefly addressed due to 392 the potential for IL-based solvents as a medium for the 'one-pot' leaching and deposition of metals. It 393 is important to note that not all reported studies use actual waste but often focus on simulated solutions 394 or mixtures containing a few elements. The use of ILs covered in this review is for the most part focused 395 on the recovery of value from PCBs, waste fluorescent phosphor, nickel metal hydride batteries and 396 permanent magnets. From the literature reviewed, optimised process flowsheets are proposed for these 397 wastes and are presented in Section 7.

398

399 4. WEEE dismantling using ILs

400 Current research on the use of ILs as WEEE dismantling agent is limited to the disassembly of 401 electrical components and epoxy resin from waste PCBs. PCBs are complex and challenging materials 402 due to their heterogeneous nature. PCBs are comprised of either: (1) an epoxy resin-based plastic 403 reinforced with glass fibres coated with a copper layer (FR-4 type PCB) or (2) a single layer of fibreglass 404 or cellulose paper and phenolic resins coated with a copper layer (FR-2 type PCB). The resin board is 405 overlaid with microelectronic components such as semiconductor chips and capacitors plus solder, 406 which is a Pb-Sn alloy (Zeng et al., 2013; Zhu et al., 2012c).

407 Previous research focused on the separation of the electrical components from the waste PCBs and/or 408 dissolution of the epoxy resin using the water soluble ILs $[C_2C_1im][BF_4]$ and $[C_4C_1im][BF_4]$. Separation 409 of the electrical components from waste PCBs relies on the high thermal stability of the IL used. The 410 welding point temperature for Pb and Pb-free solders is 215 °C (Zeng et al., 2013). By heating PCB 411 chips in the IL at 240 to 250 °C for 10 to 30 minutes, at 45 to 150 rpm, the solder was melted and 412 recovered from the board along with the previously attached electrical components. Under these

- 413 conditions, over 90 wt.% of the electronic components were separated from the PCBs and XRD analysis
- of the recovered solder indicates no structural changes (Zhu et al., 2012b; Zhu et al., 2012c; Zeng et al.,
- 415 2013). The IL can be successfully recycled, and cost analysis of the process indicates that the process
- 416 becomes financially viable for waste PCB amounts over 3 kt (Zeng et al., 2013). Heating of the bare
- 417 board in $[C_2C_1im][BF_4]$ for 10 minutes results in the complete dissolution of the polymeric materials
- 418 and the delamination of the glass and copper foils. H¹-NMR analysis indicates that H-bond formation
- 419 plays an important role in the dissolution of the bromine epoxy resins by $[C_2C_1im][BF_4]$. The IL was
- successfully regenerated by the addition of water, resulting in the formation of a precipitate which was
 confirmed as bromine epoxy resin by FTIR (Zhu et al., 2012a; Zhu et al., 2012c).
- 422

423 5. ILs in leaching and recovery applications

424 In this section, several methods for leaching metal ions and recovering them under a solid form 425 (either as oxide or salts) will be detailed. The methods presented here will not deal with liquid-liquid extraction. The dissolution of metals and metal salts is the first step in hydrometallurgical processes 426 427 (Figure 1). The state of elements in WEEE is diverse, with elements present as elemental metals, oxides, sulphates, phosphates, carbonates and others. The selection of an appropriate leaching agent depends 428 429 on the waste matrix and value of the final product and should effectively act as a concentrating step by only selectively solubilising elements of interest prior to further separation and recovery steps. Selection 430 431 of a suitable leaching medium can help reduce the complexity and number of downstream processing 432 steps. Research on metal dissolution using unconventional solvents based on ILs has focused on DES 433 and ILs incorporating Brønsted acidic functional groups and/or ILs based on strongly Lewis basic 434 anions. However, some metal oxides including Ag₂O, NiO, CuO and ZnO were found to dissolve in 435 imidazolium-based ILs with a hydrogen atom in the C² position of the imidazolium ring through the 436 formation of carbene complexes (Wellens et al., 2014b). Selected example of IL used in leaching and 437 recovery applications are summarised in Table 4.

- 438
- Table 4. Selected example of ILs used in leaching and recovery applications of metals from WEEE andores.

Waste matrix	Metals investigated	Ionic liquid / oxidant	Reference
Electric arc furnace dust	ZnO, PbO, Fe ₂ O ₃ , Al ₂ O ₃	[N _{111(20H)}]Cl:urea	Abbott et al. (2009)
Brass ash	ZnO, CuO, Cu ⁰	IL: [C4C1im][HSO4] Oxidant: H2O2 or KHSO5	Kilicarslan et al. (2014)

Metals and alloys	Fe ⁰ , Cu ⁰ , In ⁰ , Zn ⁰ , Ga ⁰ , Sb ⁰ , Au ⁰ , Pt ⁰ , Ge ⁰ , Ta ⁰ , Sm ⁰ , Dy ⁰ , GaAs, InAs	$[P_{44414}][Cl_3]$	Li et al. (2018)
Ore	Sulfidic ore containing Au, Ag, Cu, Zn, Fe, Pb	<u>IL</u> : [C ₄ C ₁ im][HSO ₄] w/ thiourea <u>Oxidant:</u> Fe ₂ (SO ₄) ₃	Whitehead et al (2004a)
Sulfidic ore / synthetic oxide ore	CuFeS ₂ , FeS ₂ , ZnS / SiO ₂ , CaSO ₄ , Fe ₂ O ₃ , Al ₂ O ₃ , MnO ₂ , TiO ₂ , AuO, AgO	<u>IL</u> : $[C_4C_1im][X]$ (X = Cl ⁻ , $[C_1SO_3]^-$, [SCN] ⁻ , [HSO ₄] ⁻) w./ thiourea <u>Oxidant</u> : Fe ₂ (SO ₄) ₃	Whitehead et al. (2007)
Ore	Sulfidic ore containing Au, Ag, Cu, Zn, Fe, Pb	<u>IL:</u> $[C_4C_1im][X] (X = Cl^-, [HSO_4]^-)$ w./ thiourea, chloride, bromide or iodide <u>Oxidant</u> : Fe ₂ (SO ₄) ₃ or KHSO ₅	Whitehead et al. (2009)
Ore	CuFeS ₂ , FeS ₂ , ZnFeS	$\frac{\text{IL:}}{\text{Oxidant:}} \begin{bmatrix} C_4 C_1 \text{im} \end{bmatrix} \begin{bmatrix} \text{HSO}_4 \end{bmatrix}$	Dong et al. (2009)
Fluoride ore / synthetic ore	RECO ₃ F, RE ₂ O ₃ , RE ₂ (CO3) ₃	$[N_{11(2(OH))}][NTf_2]$ in $[C_4C_1im][NTf_2]$	Freiderich et al. (2015)
Synthetic ore	Cu-Zn powder and crushed Ga- As wafer	<u>IL:</u> [N _{111(20H)}]Cl:EG (1:2) <u>Oxidant:</u> Iodine	Abbott et al. (2015)
Ore	Sulfidic ore containing Au, Ag, Te, Fe	IL: [N _{111(20H)}]Cl:EG (1:2) Oxidant: Iodine	Jenkin et al. (2016)
Ore	Chalcopyrite	$\underline{\text{IL:}} [C_nC_1\text{im}][\text{HSO}_4] \text{ (n=1,4)}$ <u>Oxidant:</u> O ₂	Carlesi et al. (2016)
Ore	Bauxite ore containing REE(III), Ti(IV), Si(IV), Al(III), Fe(III), Ca(II)	$[N_{111(2(0)OH)}][NTf_2]$	Davris et al. (2016)
Ore	Pyrite, arsenopyrite	[N _{111(20H)}]Cl:EG (1:2)	Abbott et al. (2017)
PCB	Cu ⁰	<u>IL</u> : [C ₄ C ₁ im][HSO ₄] <u>Oxidant:</u> H ₂ O ₂	Huang et al. (2014)
РСВ	Cu ⁰ , Zn ⁰ , Pb ⁰	<u>IL:</u> $[(C_4SO_3H)C_1im][X] (X = [OTf]^-, [HSO_4]^-), [C_nC_1im][HSO_4] (n = 1, 4)Oxidant: H_2O_2$	Chen et al. (2015a,b,c)
РСВ	Cu ⁰	IL: [(SO ₃ H)C ₄ pyr][HSO ₄] Oxidant: O ₃	Zhang et al. (2018)
NdFeB magnets	Nd ₂ O ₃ , Dy ₂ O ₃ , CoO, Fe ₂ O ₃	[N _{111(2(O)OH)}][NTf ₂]	Dupont & Binnemans (2015a)
NdFeB magnets	CoO, Fe ₂ O ₃ , B ₂ O ₃ , Nd ₂ O ₃ , Dy ₂ O ₃ , Pr ₂ O ₃ , Gd ₂ O ₃	[N _{111(20H)}]Cl:lactic acid (1:2)	Riaño et al. (2017)
Synthetic phosphor mix	$\begin{array}{l} Y_{2}O_{3}:Eu^{3+} (YOX),\\ (Sr,Ca)_{10}(PO_{4})_{6}(Cl,F)_{2}:Sb^{3+},Mn^{2+}\\ (HALO) \end{array}$	$[N_{111(2(O)OH)}][NTf_2]$	Dupont & Binnemans (2015b)

Synthetic phosphor mix & oxides	CuO, NiO, ZnO, MnO, MnO ₂ , Co ₃ O ₄ , Fe ₂ O ₃ , La ₂ O ₃ , Nd ₂ O ₃ , Dy ₂ O ₃ LaPO ₄ :Ce ₃ ⁺ ,Tb ₃ ⁺ (LAP) BaMgAl ₁₀ O ₁₇ (BAM)	$[N_{R1R2(H-SO3H)}][NTf_2]$	Dupont et al. (2016b)
Waste phosphor	Y ₂ O ₃ :Eu ³⁺ (YOX), (Sr,Ca) ₁₀ (PO ₄) ₆ (Cl,F) ₂ :Sb ³⁺ ,Mn ²⁺ (HALO)	[C ₆ C ₁ im][HSO ₄]	Schaeffer et al. (2017)
Crude In(OH) ₃	In(III), As(V), Ca(II), Cd(II), Fe(III), Pb(II), Zn(II), trace elements	$[N_{111(2(0)OH)}][NTf_2]$	Deferm et al. (2018)

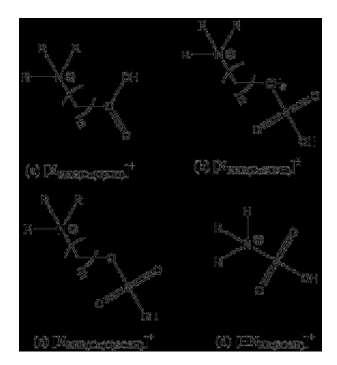
441 RE – rare earth, EG – ethylene glycol,

442

443 **5.1. Cation mediated leaching**

444 Many metal salts are insoluble in common hydrophobic ILs due to the hydrophilic nature of the 445 metallic cation and the weakly coordinating nature of the anion; metals and metal oxides are only 446 generally soluble in strong acidic or alkali aqueous solutions. First synthesised by Nockemann et al. 447 (2006), [N_{R11(2(O)OH})][NTf₂] (R - imidazolium, pyridinium, pyrrolidinium, piperidinium, morpholinium, 448 and quaternary ammonium) are a group of betaine-based TSIL bearing a carboxyl group capable of 449 selectively solubilising REO, UO₂, ZnO, CdO, HgO, NiO, CuO, PdO, PbO, MnO, Ag₂O and some 450 metal hydroxides. Insoluble or very poorly soluble oxides are Fe₂O₃, CoO, Al₂O₃ and SiO₂ (Nockemann 451 et al., 2008a). The solubility of metal oxides in [N_{R11(2(O)OH)}][NTf₂] is high due to the formation IL-RE stoichiometric compounds as indicated by single crystal XRD studies (Nockemann et al., 2008b; 452 Nockemann et al., 2009). Another interesting property of [N_{111(2(O)OH)}][NTf₂] is the presence of an upper 453 454 critical solution temperature (UCST) for the binary mixture of [N_{111(2(O)OH)}][NTf₂]/H₂O (Nockemann et 455 al., 2006). ILs with an UCST are said to exhibit thermomorphic behaviour, with the two immiscible 456 phases forming one homogeneous phases above the UCST. A two-phase system is re-established by 457 lowering the solution temperature to below the UCST. ¹H-NMR investigation attributes the presence of a UCST to the temperature-dependent hydration and hydrogen-bond formation between the carboxylic 458 group of the $[N_{111(2(O)OH)}]^+$ cation and the nitrogen atom of the $[NTf_2]^-$ anion (Nockemann et al., 2008a). 459 Conditions above the UCST reduce the viscosity of the solution and accelerate mass transfer, resulting 460 461 in higher extraction efficiencies. Extracted metals were stripped from $[N_{111(2(0)OH)}][NTf_2]$ by contacting 462 with an acidic aqueous solution, allowing the IL to be recycled for reuse.

The thermomorphic properties of $[N_{111(2(O)OH)}][NTf_2]$ were used for the recovery of Nd₂O₃, Dy₂O₃ and CoO from waste NdFeB magnets in a combined leaching-extraction step (Dupont & Binnemans, 2015a). After roasting in a microwave, waste NdFeB magnets were leached at 80°C in a $[N_{111(2(O)OH)}][NTf_2]-H_2O$ solution. The leach solution was cooled and returned to a biphasic system whereby the REE and Co were distributed in the aqueous phase whilst Fe remained in the IL phase. 468 Addition of oxalic acid to the solution precipitated the REE and Co while transferring Fe to the aqueous 469 phase as a soluble oxalate complex. The REEs and Co were separated by selectively leaching Co from 470 the oxalate precipitate using aqueous ammonia before calcining the RE oxalate to yield a 99.9 % pure Nd₂O₃ and Dy₂O₃ mix. Dupont and Binnemans (2015b) also used [N_{111(2(O)OH)}][NTf₂] for the recovery 471 472 of yttrium and europium oxide from simulated waste phosphor powder yielding a 99.9% pure final 473 Y_2O_3 : Eu³⁺ product after oxalate precipitation and calcination. The water content of the TSIL played a 474 crucial role in the effectiveness and selectivity of the leaching process. Under water-poor conditions 475 (<5 wt.% H₂O in the IL) YOX phosphor dissolves much more than HALO phosphor, but YOX leaching efficiency also decreases with decreasing IL water content. The leaching selectivity of water-poor 476 [N_{111(2(0)OH)}][NTf₂] was attributed to the inability of the IL to solvate anions due to the lack of suitable 477 coordination mechanisms. As the dissolution of oxides generates no anions, YOX phosphor easily 478 479 dissolves in [N_{111(2(O)OH)}][NTf₂] whilst the phosphate and chloride metal salt of HALO phosphor do not. The selectivity of [N_{111(2(0)OH)}][NTf₂] for REEs was also exploited for their recovery from bauxite 480 residue containing the impurities Ti(IV), Si(IV), Al(III), Fe(III) and Ca(II) (Davris et al., 2016). In 481 482 addition, [N_{111(2(O)OH)}][NTf₂] was shown to be an effective leaching agent for the purification of the critical element In(III) (Deferm et al., 2018). At the optimal conditions of 3 hrs leaching time at 80°C 483 in a 1:1 $[N_{111(2(O)OH)}][NTf_2]:H_2O$ mixture, 99 wt.% of $In(OH)_3$ was leached and separated (D > 100) 484 485 from Al(III), Ca(II), Cd(II), Ni(II) and Zn(II). Additional pre-hydrolysis and hydrolysis steps including 486 the reduction of Fe(III) to Fe(III) were required to separate In(III) from the co-extracted Fe(III), As(V) 487 and Pb(II). MgCl₂ was added as a salting-out agent to minimise the loss of the hydrophilic cation to 488 aqueous phase during extraction but its addition was found to negatively impact the separation of In(III) 489 from the other elements suggesting a compromise is required.



491

492 Figure 6. Structures of Brønsted acidic TSIL-derived from (a) betaine, (b) sulfonic acid, (c) sulfuric
493 acid and (d) sulfamic acid.

495 Three groups of zwitterionic, strongly Brønsted acidic TSILs with functionalized cations incorporating sulfonic acid groups (pKa \sim -2.0), alkylsulfuric acid groups (pKa \sim -3.5) and sulfamic 496 497 acid groups (pKa ~ 2.0) can solubilise large stoichiometric amounts for a wide range of metal oxides 498 (Dupont et al., 2015; Dupont et al., 2016a, Dupont et al., 2016b). The general structure for these cations 499 is presented in Figure 6. This includes the inert oxides CoO, Co₃O₄, Fe₂O₃, TiO₂, Al₂O₃, Cr₂O₃ and WO3, which were not previously soluble in [N111(2(O)OH)][NTf2]. A 40 wt.% solution of the sulfamic IL 500 501 [HN_{22(SO3H)}][NTf₂] diluted in [C₂C₁im]Cl was able to leach over 90 wt.% of the notoriously inert 502 fluorescent lamp phosphors LaPO₄:Ce³⁺,Tb³⁺ (LAP) and BaMgAl₁₀O₁₇ (BAM) (Dupont et al., 2016b). 503 The natural hydrophilicity of ILs incorporating these polar and very acidic functional groups was 504 overcome by using long alkyl chains in combination with the $[NTf_2]$ anion. Alkylsulfuric and sulfamic ILs with cationic alkyl chains shorter than n=3 were found to be fully miscible in water. Alkyl chains 505 506 containing 6 or more carbons were required to compensate for the greater hydrophilicity of the sulfonic 507 group. Interestingly, sulfonic and alkylsulfuric IL with short alkyl chains were shown to form aqueous 508 biphasic system in the presence of sulfuric acid and NaCl respectively. Similarly to [N_{111(2(0)OH)}][NTf₂], 509 metal extraction in these super-acidic TSIL with long alkyl chains is controlled by a proton exchange 510 mechanism and is therefore pH sensitive. As the acidic proton is exchanged for the metal ion, the 511 electronic balance is maintained, and extraction occurs generally without transfer of IL cations or anions 512 to the water phase. Comparison of hydrophobic sulfonic and alkylsulfuric IL metal extraction shows 513 that the extraction efficiency of the latter is intermediate between carboxyl-functionalized ILs such as

514 $[N_{111(2(O)OH)}]$ [NTf₂] (poor extraction for most metal ions), and sulfonic acid ILs which fully extract most 515 studied metal ions from chloride solutions (>80%) (Dupont et al., 2016a). This trend follows the 516 functional group charge density $R-COO^{-} < R-OSO_{3}^{-} < R-SO_{3}^{-}$. Interestingly in these systems, RE ions 517 are extracted in the reverse order (La(III) > Nd(III) > Dy(III) > Lu(III)) than the one expected based on 518 their charge density. This is attributed to the sterical hindrance from the functional groups and long alkyl side chains which causes REEs with smaller radius to be less efficiently coordinated (Dupont et 519 520 al., 2015). The doubly acidic task-specific pyridinium-based IL incorporating a sulfonic acid moiety 521 [(SO₃H)C₄pyr][HSO₄] was used to replace H₂SO₄ in a slurry electrolytic system for the one step 522 leaching and recovery of Cu(II) from waste PCBs (Zhang et al., 2018). A 20% increase in Cu(II) recovery to 90.9 wt.% was obtained when 10% H₂SO₄ was replaced by [(SO₃H)C₄pyr][HSO₄]. 523 524 However, a further increase in the IL concentration to 40 wt.% resulted in the decrease in the Cu(II) 525 recovery rate and current efficiency due to the increase in solution viscosity. In addition, lower quality 526 deposits were obtained at these high concentrations due to the strong competitive absorption of the IL 527 cation at the cathode inhibiting deposition. The deposit was identified as a mixture of Cu⁰, CuO and 528 Cu₂O for a 10 wt.% concentration of [(SO₃H)C₄pyr][HSO₄] with H₂SO₄ and O₃ as oxidant.

529 Although the hydrophobic IL trihexyl(tetradecyl)phosphonium chloride ([P₆₆₆₁₄]Cl) is insoluble in 530 water, [P₆₆₆₁₄]Cl can adsorb up to 13.5 wt.% of water at room temperature. [P₆₆₆₁₄]Cl saturated with 531 concentrated HCl was used to dissolve CaO, NiO, MnO, CoO, CuO, ZnO and Fe₂O₃. Addition of HCl 532 resulted in a large decrease in IL viscosity and allowed the solubilisation of metal oxides. UV-Vis 533 analysis indicated the presence of Ni(II), Cu(II), Fe(III), and Co(II) in [P₆₆₆₁₄]Cl as anionic 534 tetrachlorometallate complexes, whilst Ca(II) is unable to form anionic complexes and remains as a 535 hydrated calcium ion. Separation of the solubilised elements using a stripping HCl solution was based 536 on the tendency of Ni(II) and Ca(II) to form aqua complexes regardless of HCl concentration and the 537 partial stripping of Co(II) an Mn(II) at low HCl concentrations. Ni(II) and Ca(II) were first stripped 538 using 6 M HCl before subsequent removal of Co and Mn from Fe, Zn and Cu by contacting the saturated 539 IL with a water and ammonia solution (Wellens et al., 2014a).

540

541 **5.2.** Anion mediated leaching

542 $[HSO_4]^-$ -based IL including $[C_4C_1im][HSO_4]$ were successfully used to recover Cu from waste 543 PCBs (Huang et al., 2014) and chalcopyrite ore (Dong et al., 2009; Chen et al., 2015a), Cu and Zn from 544 brass ash (Kilicarslan et al., 2014), leaching of Au, Ag, Cu and base metals from ores (Whitehead et al., 545 2004a; Whitehead et al., 2007; Whitehead et al., 2009) and YOX phosphor from waste fluorescent tubes 546 (Schaeffer et al., 2017).

547 The recovery of yttrium europium oxide from waste fluorescent tube phosphor by a simple efficient 548 low-cost IL process was developed using $[C_4C_1im][HSO_4]$ (Schaeffer et al., 2017). The waste phosphor

- 549 containing Y(III), Eu(III), Tb(III), La(III) and Gd(III) was first pre-treated to concentrate its rare earth 550 content. Larger glass particles were removed by sieving through a 25 µm sieve whilst soluble Ca(II), 551 Ba(II), Sr(II) were partially removed by contacting the phosphor with a dilute HCl solution for 1hr. 552 Leaching of REEs was optimised by systematically varying the leaching temperature, the 553 $[C_4C_1]$ [HSO₄]:H₂O ratio, leaching time and solid to liquid ratio. Under the optimised conditions of 554 1:1 wt. [C₄C₁im][HSO₄]:H₂O solution, a solid:liquid ratio of 5%, a mixing speed of 300 rpm, a leaching 555 temperature of 80°C and leaching time of 4 h, the percentage of Y, Eu and Ca leached is 91.6wt.%, 97.7wt.% and 24.9wt.%, respectively. The low leaching efficiency of Tb(III), La(III) and Gd(III) 556 557 compared to Y(III) and Eu(III) is due to the former being as highly stable phosphate salts whilst the 558 latter are found as oxides. The REEs in the leachate were precipitated by the addition of oxalic acid at 559 a molar ratio (OA:REE) of 1.5 and a solution pH of 0.9. Luminescence analysis of the recovered oxide 560 product after calcination indicates that the recovered yttrium europium oxide has the potential to be 561 directly reused as YOX phosphor. The stability and recyclability of the IL was demonstrated over four 562 leaching/precipitation cycles.
- 563 A recent comparison of six ILs incorporating $[HSO_4]^-$ and $[CF_3SO_3]^-$ on the leaching efficiency of Cu, Zn and Pb respectively from waste PCBs found that greater leaching efficiency could be achieved 564 565 using [HSO₄]⁻ and that inclusion of an acidic group on the alkyl chain of the cation increased Cu and 566 Zn leaching (Chen et al., 2015a; Chen et al., 2015b). Leaching results indicate that all six IL acids could 567 successfully recover Cu and Zn under the test conditions with almost 100 wt.% recovery except for 568 $[C_1C_1im]$ [HSO₄]. Kinetic analysis of the dissolution process using the shrinking core set of equations 569 indicates that in the case of Cu and Zn dissolution by $[HSO_4]^-$ and $[CF_3SO_3]^-$ ILs, the reaction rate is 570 controlled by diffusion processes and not by the surface chemical reaction. However, the study of Pb is 571 of minimal interest because of the aqueous insolubility of PbSO₄, with little to no recovery recorded for 572 the $[HSO_4]^-$ and $[CF_3SO_3]^-$ ILs tested (Chen et al., 2015c). Quantitative Cu extraction from waste PCBs 573 using $[C_4C_1im][HSO_4]$ was also achieved under the leach conditions: 1 g PCB powder (particle size of 574 0.1–0.25 mm), 25 mL 80% (v/v) [C₄C₁im][HSO₄], 10 mL 30 wt.% H₂O₂, solid: liquid ratio of 1:25, 575 70°C and 2 hours. Kinetic analysis of the leaching process determined that this was diffusion controlled 576 with an activation energy of 25.4 kJ.mol⁻¹ (Huang et al., 2014). None of these cited studies investigated 577 the concomitant leaching of other elements or the recovery of Cu once in solution despite the interesting 578 work of Zhang and Hua, which extensively studied the deposition of Zn (Zhang & Hua, 2009; Zhang 579 et al., 2009a; Zhang & Hua, 2011; Zhang & Hua, 2012) and Cu (Zhang et al., 2009b) from aqueous 580 sulphate solutions with [C₄C₁im][HSO₄] or [N₄₄₄₄][HSO₄] (Zhang et al., 2015) as additives. Both ILs 581 were found to inhibit electro-reduction, leading to finer and more levelled deposits. In contrast, 582 dissolved ZnO in [C₁C₁im][CF₃SO₃] was recovered by electro-deposition, with high current efficiencies 583 both in neat and aqueous IL solutions. FTIR and Raman spectroscopy indicated that Zn(II) is stabilised 584 by 1-methylimidazole rather than the IL anion (Liu et al., 2015).

585 Many of the species of interest are present in their elemental form and must be oxidised prior to 586 solubilisation. The solubility of oxygen in ILs is of crucial interest as it may determine the leaching rate 587 and efficiency. The solubility of oxygen in neat imidazolium-based IL with [HSO4]⁻ anion is similar or 588 even superior to that in water, decreases with increasing temperature, and is anion dependent. Larger 589 anion-cation pairings leave larger 'holes' appropriate for oxygen residency compared to more compact 590 ion-pairings. In addition, dipolar interactions between oxygen and the IL anion, influenced by the shape 591 and surface charge density on the anion, are also likely to result in increased oxygen solubility 592 (Whitehead et al., 2007). Common oxidants used with $[C_nC_1im][HSO_4]$ include hydrogen peroxide 593 (H_2O_2) , potassium peroxymonosulfate (KHSO₅) and Fe(III). Iron(III) is a mild oxidant and unlikely to 594 affect the stability of the IL. However, none of the reported studies using $[C_nC_1im][HSO_4]$ as lixiviant 595 have investigated the potential denaturalization of the IL after a leaching cycle in the presence of strong 596 oxidants such as H_2O_2 . Results from previous research suggest that $[C_nC_1im][HSO_4]$ is readily degraded 597 by of H_2O_2 (Domínguez et al., 2014). The length of the alkyl chain has a dramatic effect on the stability 598 of imidazolium-based ILs to degradation by stoichiometric amounts of H₂O₂, with greater resistance 599 conferred by longer alkyl chains. Furthermore, ILs bearing methanesulfonate and methylsulfate anions 600 are oxidised faster than those with chloride or acetate anions. A compromise between metal extraction 601 and IL recovery is required under such circumstances

602 An innovative class of ILs containing trichloride anions (Cl_3) with oxidative properties capable of 603 releasing chlorine was synthesised by bringing chlorine gas into contact with the corresponding chloride 604 (Cl⁻) ILs at room temperature (Li et al., 2018). These were prepared with a range of cations with no 605 degradation of the latter observed. An additional benefit from the conversion of the IL anion Cl⁻ to Cl₃⁻ 606 is the accompanying reduction in the IL viscosity due to the lower electrostatic attraction between the 607 cation and the Cl_3^- ion. Taking $[P_{44414}]Cl_3$ as a representative IL, dissolution experiments were 608 performed for 12 metals and 2 alloys in solutions at 25 °C containing molar ratio of metal:IL of 1:8. 609 Whilst the dissolution time was found to greatly vary between metals tested, 8 metals commonly found 610 in WEEE Cu, Fe, In, Zn, Ga, Au, Ge and Sb along with the alloys InAs and GaAs could be dissolved 611 in the tested IL. EXAFS analysis revealed that these metals exist as anionic chloride complexes in the 612 IL.

613

614 5.3. Deep eutectic solvents (DES) as leaching agents

DES are usually composed of two chemicals, typically one hydrogen bond donor (HBD) and one hydrogen bond acceptor (HBA), exhibiting high melting points. Once combined in specific stoichiometric ratios, the mixture melts and forms a homogeneous liquid phase. The DES under consideration in this review are those based on at least one IL homologues, namely a quaternary ammonium salt, choline chloride ($[N_{111(20H)}]Cl$) being the most common, and a HBD such as urea, 620 carboxylic acids or polyols (Smith et al., 2014). DES exhibit unusual solvent properties that are 621 dependent on the selection of the HBD and HBA and their respective molar ratio. The considerable 622 attraction of DES stems from low synthesis cost, low environmental toxicity and solubility for a wide 623 range of metal oxides (Abbott et al., 2002; Abbott et al., 2004).

624 Metal oxide solubility in DES is influenced by the extent of H-bonding interactions, with compounds 625 able to donate or accept electrons or protons to form H-bonds displaying high solubility (Abbott et al., 626 2002). The solubility of 17 metal oxides in the elemental mass series Ti through to Zn was reported, 627 with [N_{111(20H)}]Cl:malonic acid displaying the greatest solubility for the tested elements after HCl 628 (Abbott et al., 2006). Most ionic transition metal oxides are soluble in DES whilst covalent metal oxides such as SiO₂ and Al₂O₃ are insoluble in all tested DES to date. Judicious selection of an appropriate 629 630 HBD allows for the selective leaching of certain elements from a mixed oxide matrix. For example, Fe₃O₄ is soluble and CuO insoluble in [N_{111(2OH)}]Cl:oxalic acid eutectic whilst the opposite behaviour 631 is observed when the HBD is changed to phenylpropionic acid (Abbott et al., 2004). Furthermore, 632 633 compounds that are known to form DES with choline chloride were successfully used for the leaching 634 of Mn, Li, Co, and Ni using either L-tartaric acid (He et al., 2017) and lactic acid (Li et al., 2017) 635 highlighting the untapped potential of DES as leaching agents. In addition to suitability for metal oxide 636 leaching, a growing number of studies employ DES as electrolytes for metal electro-deposition, 637 allowing for the development of a 'one pot' solvatometallurgical process. A large number of metals 638 including Zn, Ni, Cu, Co, Sn, Se, In, Sm, Pd, Fe, Ag and Pb were successfully deposited from DES 639 (Abbott et al., 2013; Smith et al., 2014; Zhang et al., 2016).

640 The establishment of a comparable electrochemical series for 17 redox couples referenced to the 641 $Fe(CN)_6^{3-/4-}$ couple in the DES [N_{111(20H)}]Cl:urea and [N_{111(20H)}]Cl:EG shows that some redox potentials shift considerably in DESs compared to aqueous solution, particularly the oxophilic p-block elements 642 643 such as Ga and Sb, Cu and the chlorophilic late transition elements including Ag, Pd and Au. The high 644 chloride concentration in DES electrolytes acts as an environmentally benign stabilising ligand used to adjust metal speciation, thereby controlling solubility and deposition properties, making DES excellent 645 646 separation and recovery media (Abbott et al., 2011b). The determination of the electrochemical series 647 served as the basis for the recovery of value from synthetic wastes matrices and ores (Abbott et al., 648 2015; Jenkin et al., 2016). Iodine was selected as the electrocatalyst due to its fast electron transfer, 649 reversibility, high rates of mass transport, high solubility in DES, and environmental compatibility in 650 reduced form (Abbott et al., 2011b; Abbott et al., 2015). The redox potential of I_2/I^- in a [N_{111(20H)}]Cl:EG 651 DES was found to be more positive than that of most common metals. The developed system depicted in Figure 7 could oxidize Cu and Zn from 1:1 Cu-Zn powders, As and Ga from crushed Ga-As wafer 652 653 and Au and Ag from gold-bearing ores (Abbott et al., 2015, Jenkin et al., 2016).

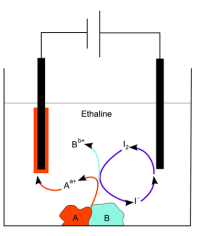


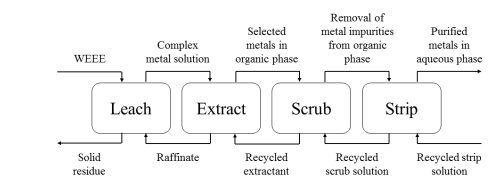
Figure 7. Schematic diagram of the separation and recovery of two elements, A and B, using iodine
 as an electrocatalytic oxidation agent in [N_{111(20H)}]Cl:EG. Reproduced from Abbott et al. (2015) with
 permission from The Royal Society of Chemistry.

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660 6. ILs in solvent extraction processing

661 Solvent extraction (or liquid-liquid extraction) is the favored option for metal separation and 662 involves the preferential two-phase distribution of a solute between mutually immiscible liquid phases, 663 usually consisting of an aqueous acidic solution and an aliphatic organic solvent. The ILs used in solvent 664 extraction must be water immiscible to promote recovery of the metal and prevent the loss of the IL to 665 the aqueous phase. Metal extraction from the aqueous phase results from the formation of hydrophobic 666 Mⁿ⁺-ligand complexes and their migration to the hydrophobic phase (Fray, 2000; Rydberg et al., 2004). A schematic of the typical process steps in the recovery of metals from leach solutions using solvent 667 668 extraction is presented in Figure 8.

669



671 **Figure 8.** Schematic of the typical process steps in the recovery of metals from leach solutions using

solvent extraction.

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674 This section is divided according to the application of the IL-based solvent, with a division being 675 made between traditional solvent extraction processes and more recent advances. First, the use of 676 undiluted IL acting as both the extractant and hydrophobic phase in traditional solvent extraction is 677 presented. Secondly, the application of IL as hydrophobic diluting phase in conjunction with neutral and acidic extractant is discussed. Finally, the application of TSIL in solvent extraction for the recovery 678 679 of metals from WEEE is addressed. Following this review of more 'traditional' solvent extraction 680 processes, the use of IL-based aqueous biphasic system applied to WEEE valorisation is discussed as 681 well as the developing field of hydrophobic DES and triphasic IL-IL systems for the simultaneous separation of different metal groups. Because of the large volume of publications on the topic, the 682 discussion will be limited to a selected number of extraction systems directly addressing the recovery 683 of metal from WEEE or the separation of critical metal groups relevant to recycling of WEEE. The 684 685 major studies involving the use of ILs in solvent extraction processes for the recovery of metals from 686 WEEE are summarised in Table 5.

Table 5. Reported IL used in solvent extraction processes for the recovery of metals from WEEE and
 ores (NMHB – nickel metal hydride battery, FLP – fluorescent lamp phosphor).

Waste substrate	Elements recovered	Extractant	Diluent	Reference
Synthetic NMHB leach solution	Co, Ni, Fe, Mn, Zn, K, Mg, REE	–	P ₆₆₆₁₄]Cl N ₁₈₈₈]Cl (3) [N ₁₈₈₈][NO ₃]	Larsson & Binnemans (2014)
		(4) [N ₁₈₈₈][SCN]	(4) [N ₁₈₈₈]Cl	
Synthetic NMHB leach solution	Co, Ni, Fe, Mn, Zn, REE	Cyanex 923	[N ₁₈₈₈][NO ₃]	Larson & Binnemans (2015a)
Synthetic NdFeB and SmCo magnet leach solution	Nd, Sm	[(CnCOOH	Chen et al. (2015d)	
NdFeB magnet	Pr, Nd, Dy	TBP	[N ₁₈₈₈][NO ₃]	Kikuchi et al. (2014)
NdFeB magnet	Nd, Dy, Co, Cu, Fe, Mn	[P ₆₆₆₁₄]Cl		Vander Hoogerstraete et al. (2014)
NdFeB magnet	Nd, Sm, Fe, Co	[P ₆₆₆₁₄][NO ₃]		Riaño & Binnemans (2015)
NdFeB magnet	Nd, Sm, Pr, Gd, Fe, Co, B	(1) [N ₁₈₈₈][SCN] (2) Cyanex 923	Toluene Toluene	Riaño et al. (2017)

Simulated FLP solution	REE	[N ₁₈₈₈][PC-88A] & [N ₁₈₈₈][DEHPA]		Yang et al. (2012)
FLP	La, Ce, Eu, Gd, Tb, Y	DODGAA	$[C_4C_1im][NTf_2]$	Yang et al. (2013)
Commercial HALO phosphor	Sb, Ca	[N ₁ :	Dupont & Binnemans (2016)	
Bastnäsite leach solution	Ce, F	DEHEHP	[C ₈ C ₁ im][PF ₆]	Zuo et al. (2009)
Bastnäsite and monazite	La, Ce, Nd, Th, Ca, Mn, Fe, Mg	[N ₄₄₄₄][DEHPA]	[EtNH ₃][NO ₃] / [C ₂ C ₁ im][SCN] / [C ₄ C ₁ im][BF ₄].	Azizi & Larachi (2018)
Salt Lake brine	Li, Mg	TBP	[C ₄ C ₁ im][NTf ₂]	Shi et al. (2016)
Bauxite residue	Sc, Fe, Ti, Si, Ca, Al	[N ₁₁₁₍₂₍₀₎	Onghena et al. (2017a)	

691 6.1. Undiluted ILs

692 Recent research on the application of ILs in solvent extraction has focused on simple IL extraction 693 systems without added complexing ligand. Hydrophobic ILs based on [NTf₂]⁻ and cation imidazolium, 694 pyrrolidinium or pyridinium have been shown to extract aqueous metal cations without the need for 695 additional ligands (Kozonoi & Ikeda, 2007). Regarding metals found in WEEE, these ILs were found 696 to be efficient at extracting precious metals such as Au(III), Pt(IV) or Ir(IV) from acidic aqueous 697 solutions containing large amounts of HCl (Génand-Pinaz et al., 2013; Papaiconomou et al., 2012 & 698 2014; Yang et al., 2014; Tong et al., 2015; Kubota et al., 2017). Furthermore, separation of Au(III) from 699 Pt(IV) and from Pt(IV) and Pd(II) is achieved using $[C_4C_1im][NTf_2]$ and $[C_8C_1im][NTf_2]$, respectively 700 (Papaiconomou et al., 2012; 2014).

701 Linear quaternary ammonium ILs and quaternary phosphonium ILs have also been extensively 702 studied and used in complete recycling processes of WEEE. ILs based on these cations and a wide 703 variety of anions including chloride, bromide, dicyanamide (Svecova 2015, Svecova 2016), nitrate, 704 dioctyl diglycolamate (Rout & Binnemans, 2014a), N,N,N',N'-tetra(2-ethylhexyl)malonate, N,N,N',N'-705 tetra(2-ethylhexyl)malonamide (Rout & Binnemans, 2014b) or oleate (Parmentier et al., 2015a) were 706 studied. The hydrophobic fatty acid based IL tetraoctylphosphonium oleate ([P₈₈₈₈][C₁₇COO]) displays 707 a high affinity for first row transition metals in chloride media (Parmentier et al., 2015a) and was 708 employed in a continuous selective metal extraction process for the recovery of cobalt from two mixed 709 salt solutions (Co/Na, Ca/Co/K) (Parmentier et al., 2015b). The ammonium alternative to 710 [P₈₈₈₈][C₁₇COO], tetraoctylammonium oleate ([N₈₈₈₈][C₁₇COO]), was used in the solvent extraction of 711 Co(II), Ni(II) and Zn(II) with success. Comparison of Co(II), Ni(II) and Zn(II) extraction with the IL 712 as well as the IL precursors (sodium oleate and tetraoctylammonium chloride) indicates that Co(II) and 713 Ni(II) at low chloride concentration are extracted via the IL anion whilst Zn(II) can be extracted both 714 via the oleate anion and via the tetraoctylammonium cation of the IL. Slope analysis and EXAFS 715 measurements reveal that Ni(II) and Co(II) are extracted via the formation of $[N_{8888}^+][M(C_{17}COO)_3^-]$ 716 complexes with M being Co(II) or Ni(II). In contrast, Zn(II) was extracted via a different extraction 717 mechanism, in which the Zn(II) ion was extracted by two IL moieties as $[N_{8888}^+]_2[MCl_2(C_{17}COO)_2^{2-}]$ 718 (Parmentier et al., 2016). This example highlights the need to determine the dominant extraction 719 mechanisms as this can shed light on the higher distribution ratio for one metal compared to another. 720 Other interesting phosphonium- and ammonium-based ILs for the extraction of transition metals include 721 those containing a thiosalycilate or thiocyanate anion (Leyma et al., 2016; Rout & Binnemans, 2016). 722 The latter was used for the separation of transition metals from rare earth elements, a separation relevant for the recycling of permanent magnets, nickel metal hydride batteries and lamp phosphors. The 723 724 extraction of metal ions from a chloride or nitrate solution with a thiocyanate IL is an example of "split-725 anion extraction", where different anions are present in the aqueous and IL phase. As per the Hofmeister 726 series, the $[SCN]^-$ prefers the IL phase compared to the Cl⁻ and $[NO_3]^-$ ions and does not exchange to 727 the aqueous phase, allowing it to preferentially coordinate the 3d transition metals.

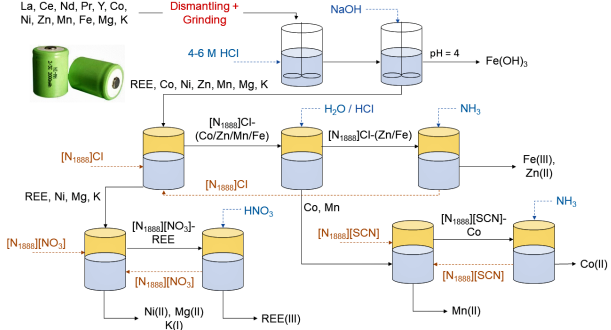
728 Trihexyl(tetradecyl)phosphonium chloride ($[P_{66614}]Cl$) is a commercially available, non-fluorinated 729 IL consisting of an organophosphorous-based cation with a chloride anion - cf. Figure 5. Due to the four 730 long alkyl chains, [P₆₆₆₁₄]Cl has no reported surfactant properties, in contrast to some of the shorter alkyl chain quartenary phosphonium salts alternatives, thereby enhancing phase separation (Bradaric et 731 732 al., 2003). [P₆₆₆₁₄]Cl functions by an anion exchange mechanism; and it can be used undiluted, 733 eliminating the need for molecular solvents. The extraction experiments have primarily been conducted 734 from aqueous chloride medium. [P₆₆₆₁₄]Cl IL shows great promise for the extraction of metal ions and 735 has been studied for the extraction of Zn(II), Fe(II/III), Co(II) as well as REE(III), Au(III), Hg(II), 736 Pd(II), Pt(IV), Bi(III), Re(IV) and U(IV). For example [P₆₆₆₁₄]Cl, or [P₆₆₆₁₄]Br, extract almost 737 quantitatively Pd(II) and Pt(IV) from aqueous solutions of HCl (Papaiconomou et al., 2015; Kubota et 738 al., 2017). Distribution coefficients as high as 200 for Rh(III) are obtained in aqueous solution containing 1 mol.L⁻¹ HCl, while Rh(III) remains preferentially in the aqueous phases when the 739 concentration of HCl exceeds 6 mol.L⁻¹ (Svecoca et al., 2016). Aliquat 336 ([N₁₈₈₈]Cl) is a well-740 741 established quaternary ammonium extractant capable of extracting a range of metal ions through anion exchange or formation of a neutral extraction complex (Černá et al., 1992). 742

 P_{66614}]Cl and its nitrate derivative trihexyl(tetradecyl)phosphonium nitrate were employed in separations relevant to the recycling of rare earths from permanent magnets and Co from Ni metal hydride batteries. Wellens et al. (2012) demonstrated that Co(II) could be selectively separated from

- 746 Ni(II), Ca(II) and Mg(II) in 8 M HCl aqueous chloride concentration using [P₆₆₆₁₄]Cl with a separation 747 factor of ≥ 104 . Co was extracted as the tetrahedral tetrachlorocobaltate complex, $[CoCl_4]^{2-}$. Co(II) was 748 separated from the co-extracted Mn(II) by four stripping cycles with water. [P₆₆₆₁₄]Cl was found to be 749 the best option as the IL phase compared to [N₁₈₈₈]Cl, [P₈₈₈₈]Br, [P₆₆₆₁₄]Br and [P₄₄₄₁₄]Cl as it is a compromise between commercial availability, separation characteristics and ease to handle the IL. This 750 751 process has been up-scaled to provide a proof-of-principle for the continuous IL separation process of 752 Co(II) from Ni(II) (Wellens et al., 2013). A similarly high Co/Ni separation ratio was achieved using [P₆₆₆₁₄]Cl from sulphate media with the extraction proceeding via a split-anion mechanism. Co(II) 753 754 present in the IL phase was stripped using water and the IL regenerated by contacting with CaCl₂ 755 (Onghena et al., 2017b).
- 756 [P₆₆₆₁₄]Cl was also employed for the removal of Co(II) and Fe(III) from Sm(III) and Nd(III) 757 respectively in chloride solutions (Vander Hoogerstraete et al., 2013). Separation factors of 5.0×10^6 758 and 8.0×10^5 were obtained for Nd/Fe and Sm/Co respectively at HCl concentrations of 8.5-9 M. The REE(III) remain in the aqueous phase while Co(II) and Fe(III) are extracted as anionic tetrachloro-759 760 complexes in the IL. The distribution of Co(II), Mn(II) and Fe(III) to the IL increased with increasing 761 HCl concentration until 8.5-9 M while Zn(II) and Cu(II) decrease with increasing HCl concentrations. However, Al(III), Cr(III), Ni(II), Mg(II), and Ca(II) could not be separated from REEs. After extraction, 762 763 Co was easily stripped from the IL with water although Fe could only be stripped by forming a water-764 complex with ethylenediaminetetraacetic acid (EDTA). Using the soluble iron IL 765 trihexyl(tetradecyl)phosphonium nitrate, prepared from [P₆₆₆₁₄]Cl, REEs were separated from Co(II) 766 and Ni(II) (Vander Hoogerstraete & Binnemans, 2014). The separation of Sm(III) from Co(II) and 767 La(III) from Ni(II) were studied. Extraction of the REEs is facilitated by an inner salting-out effect of 768 a highly concentrated metal nitrate aqueous phase. After a scrubbing step using 7.5 M of NH₄NO₃, the 769 purity of REEs in the loaded IL phase was 99.9%. Complete stripping and regeneration of the IL was 770 achieved using pure water. Extraction of REE by [P₆₆₆₁₄][NO₃] from nitrate solution occurs by formation 771 of a neutral complex $Ln(NO_3)_{3+n}(P_{66614})_n$, while La and Sm exist as the anionic $[La(NO_3)_6]^{3-}$ and [Sm(NO₃)₆]³⁻ complexes in the IL. The high negative charge of the extracted complexes and the 772 773 resulting electrostatic interactions have a pronounced effect on the viscosity of the IL phase.
- 774 [P₆₆₆₁₄]Cl was used for the recovery of REEs from waste NdFeB permanent magnets in a seven-step 775 process (Vander Hoogerstraete et al., 2014). This included (1) crushing and milling the magnet into 776 coarse powder, (2) roasting to transform the metals into the corresponding oxides, (3) selective leaching 777 of the REEs with HCl leaving Fe in the solid residue by raising the pH of the leach solution above 2, 778 (4) extracting Co(II), Cu(II) and Mn(II) into [P₆₆₆₁₄]Cl 101 leaving the REEs in the aqueous phase, (5) 779 precipitating the REEs by the addition of oxalic acid, (6) recovering the precipitate by filtration and (7) 780 calcining the oxalate precipitate to yield rare-earth oxides. Roasting the waste magnet prior to 781 dissolution was found to facilitate dissolution and reduces the HCl consumption, with \geq 90 wt.% of

 Nd_2O_3 and Dy_2O_3 extracted compared to ≤ 25 wt.% of Fe₂O₃. A 3.5 M NH₄Cl was added to the leachate 782 783 as a salting-out agent to prevent the extraction of the REEs into the IL. An alternative process for the 784 extraction and separation of RE and other valuable elements from used NdFeB permanent magnets was 785 proposed by Riaño & Binnemans (2015) using [P₆₆₆₁₄][NO₃]. After roasting, the used magnets were dissolved in HNO₃. At the end of the leaching process, the solution pH was raised to 4 to remove Fe by 786 precipitation. The remaining Nd(III) and Sm(III) were separated from Co(II) using [P₆₆₆₁₄][NO₃]. 787 Evaluation of the extraction parameters indicated that pH had little influence on the process selectivity, 788 Co(II) extraction was supressed at high metal loading concentrations and addition of NH₄NO₃ salted in 789 the REEs. After scrubbing Co(II) using NH4NO3, the loaded Nd(III) and Sm(III) were separated by 790 791 stripping the IL phase using an EDTA and 10 M NH₄NO₃ solution. The preferential interaction of 792 EDTA with Dy allowed for extraction to the aqueous phase whilst a high NH₄NO₃ concentration 793 suppresses stripping of Nd. At 70 °C, 0.03 M EDTA and 10 M NH NH₄NO₃, 79.8% and 4.0% of Dy 794 and Nd were stripped. Addition of concentrated HNO₃ to the stripping solution dissociated the weaker 795 [Nd(EDTA)]⁻ complex and facilitated its back extraction into the IL yielding a final Dy solution. All 796 elements were precipitated as oxalate salts and calcined. Although this process is complicated due to the many steps involved, over 99% of Co, Nd and Dy are recovered as oxide with >99.5 % purity. 797

Larsson and Binnemans (2014) attempted the selective recovery of metals from a synthetic nickel metal hydride battery leach solution using four IL systems: $[P_{66614}]Cl$, $[N_{1888}]Cl$, 10% v/v Cyanex 923 (neutral extractant) in $[N_{1888}][NO_3]$ and 30% v/v $[N_{1888}][SCN]$ in $[N_{1888}]Cl$. A simplified process flowsheet is presented in Figure 9. This process clearly shows the potential of ILs to recover metals from complex aqueous solutions using careful selection of the ILs and understanding their influence on the metal extracted complex (chloride vs. nitrate, etc).



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 807 Figure 9. Flow-sheet describing recovery of metals from waste nickel metal hydride batteries using ILs (adapted from Larsson & Binnemans, 2014).

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810 The feasibility of an efficient solvent extraction process for the purification of the critical element 811 indium from a chloride medium was demonstrated using [P₆₆₆₁₄]Cl and [N₁₈₈₈]Cl. Results from single-812 element extraction studies indicate that the phosphonium IL [P₆₆₆₁₄]Cl outperformed [N₁₈₈₈]Cl with 813 higher distribution ratio obtained across the entire HCl concentration range due to the greater 814 hydrophobicity of [P₆₆₆₁₄]Cl. Maximum extraction occurred at 5 M HCl for both ILs, with the extraction 815 mechanism assigned to the evolution of indium speciation as a function of chloride concentration and 816 the formation of anionic chlorindate complexes at high HCl concentration. A percentage stripping over 817 99% was reached for the 0.5 M HCl-[P₆₆₆₁₄]Cl system if 4.5 equivalents of NaOH were used. Two 818 successive extraction/stripping cycles indicate that the IL can be recycled for reuse in liquid/liquid 819 extraction of indium(III) without any loss of activity and with little loss of [P₆₆₆₁₄]Cl. Multi-element 820 extraction studies show [N₁₈₈₈]Cl is the preferred IL for the separation of In(III) from As(III), Mn(II) 821 and Ni(II), taking advantage of the low affinity of these metals for the IL phase at lower HCl 822 concentrations. However, In(III) could not be separated from cadmium(II), iron(III), lead(II), tin(IV) 823 and zinc(II) in a single step and additional scrubbing and stripping steps appear necessary (Deferm et 824 al., 2016).

 $[N_{1888}]$ -based ILs were used for the recovery of antimony and REEs from simulated waste phosphor from fluorescent lamps. Phosphor powder used in fluorescent lamps is a mixture of individual phosphors, namely HALO (Ca_{4.86}Mn_{0.10}Sb_{0.04}Sr_{0.004}(PO₄)₃Cl_{0.10}F_{0.90}), YOX (Y_{1.92}Eu_{0.08}O₃) and other constituents. Dupont and Binnemans (2016) considered the leaching and recovery of Sb from HALO phosphor only. Commercial HALO phosphor was completely leached using dilute HCl and extracted

- 830 as the anionic tetrachloroantimonate(III) complex (SbCl₄⁻) using $[N_{1888}]$ Cl by an anion exchange
- 831 mechanism. Quantitative extraction of Sb(III) was achieved from the leach solution containing Sr(II),
- 832 Ca(II) and Mn(II) even in the absence of addition chloride ions. Mn(II) was only extracted at very high
- 833 chloride concentrations (\geq 4 M NaCl), whilst Sr(II) and Ca(II) do not form anionic chloro-complexes
- under the tested experimental conditions and were therefore not extracted. The loaded IL phases was
- scrubbed with 1 M NaCl prior to Sb(III) stripping using a NaOH solution at pH 7. The residual Ca in
- the leach solution was precipitated by addition of NaOH to yield a valuable calcium phosphate apatite
- 837 precipitate.
- 838 Yang and co-workers (2012) used "bifunctional IL extractants" (Bif-ILEs) prepared from [N1888]Cl 839 and the commercial organophosphorus acid extractants, di-(2-ethylhexyl)phosphoric acid (DEHPA) and di(2-ethylhexyl)orthophosphinate (PC-88A), to extract REEs from a simulated solution of a 840 841 fluorescent powder containing a high concentration of Al(NO₃)₃. [N₁₈₈₈][DEHPA] and [N₁₈₈₈][P507] 842 possess binary functional groups, one each on the cation and anion, through which complexation can 843 occur. The extraction performance of the synthesized [N₁₈₈₈][DEHPA] and [N₁₈₈₈][P507] Bif-ILEs was 844 compared to that of neutral organophosphorus extractants di(1-methylhepthl)methyl-phosphonate, TBP 845 and Cyanex 923 in *n*-heptane. It was found that the sequence of extractability for REE(III) is Cyanex 923 $> [N_{1888}][DEHPA] = [N_{1888}][PC-88A] > di(1-methylhepthl)methyl-phosphonate > TBP. The extraction$ 846 using [N₁₈₈₈][DEHPA] and [N₁₈₈₈][PC-88A] was thought to occur through the unshared electron pair of 847 848 P=O in the [PC-88A]⁻ or [DEHPA]⁻ groups and the N lone electron pair of the $[N_{1888}]^+$ group to form 849 macromolecular extracted complexes (Figure 10).
- 850

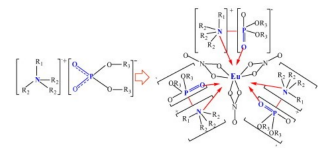


Figure 10. The proposed structure of [N₁₈₈₈][DEHPA] and its coordination environment with Eu(III)
 (Sun et al., 2010).

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The coexisting Al_2O_3 in the fluorescent powder was changed to a salting-out agent ($Al(NO_3)_3$) in the extraction process and this increased the extraction efficiency of REEs. Using a counter-current extraction process at a 4:1 organic to aqueous phase ratio and pH = 0.56, the REE recovery of 95.2 wt.% was achieved in stages 5-7. However, the use of Bif-ILEs had some drawbacks, namely the addition of 10% isopropanol modifier to prevent the formation of a third-phase on extraction and the 860 important viscosity of the IL phase after extraction (Yang et al., 2012). Another study using the same

Bif-ILEs systems found that $[N_{1888}]$ [PC-88A] and $[N_{1888}]$ [P204] were suitable for the separation of

heavy REEs in nitrate medium and the separation of light REEs in chloride medium (Guo et al., 2014).

- 863 EXAFS studies attributed the synergistic effect of bifunctional IL extractant $[N_{1888}]$ [DEHPA] for REE
- 864 extraction to the greater stability and hydrophobicity of the metal complex formed. In addition, 865 extraction was shown to proceed via co-extraction of the aqueous phase counter anion to produce a
- 866
- 867

868 **6.2. Neutral extractants**

neutral complex in the IL phase (Sun et al., 2010).

869 The use of neutral extractant in conjunction with an IL diluent system for the recovery of aqueous 870 metal complexes is the best studied extraction phenomenon involving ILs (Dai et al., 1999). Extraction 871 using neutral extractant proceeds via an ion-exchange mechanism whereby to compensate for incoming 872 positively charged metal cation into the IL, an IL component is released to the aqueous phase to maintain charge neutrality. This was reported for a wide range of neutral ligands including crown ethers 873 874 (Dietz & Dzielawa, 2001; Jensen et al., 2002; Dietz et al., 2003; Jensen et al., 2003; Dietz & Stepinski, 2005; Garvey et al., 2012; Hawkins et al., 2012), phosphine oxide ligands (Kubota et al., 2009; Petrova, 875 876 2016), phosphate ligands (Billard et al., 2011; Zhu et al., 2015), diglycolamide ligands (Shimojo et al., 877 2008; Shen et al., 2011), calixarene ligands (Luo et al., 2004) and others (Figure 5).

878 Systematic studies on the extraction of alkaline and alkali earth cation with DCH18C6 crown ether extractant in [CnC1im][NTf2] show that the predominant extraction mechanisms, ion-exchange or 879 880 neutral complex extraction, depend on the hydrophobicity of the IL cation, the Lewis acidity of the 881 metal ion and the aqueous phase nitrate anion concentration (Hawkins et al., 2012; Wankowski & Dietz, 882 2016). Jensen et al. (2002) reported that for such systems, the cation exchange extraction mechanism is 883 not driven by the strength of metal-extractant complexation but by the energetically favourable 884 solvation energy obtained from the release of the IL cation to the aqueous phase. Although the IL 885 components can be recovered from the aqueous solution by 'salting-in', this adds an extra step to the 886 recovery process and decreases its economic viability. The toxicity of low concentrations of the cation 887 in waste water must also be considered. Increasing the cationic alkyl chain or incorporating fluorinated 888 groups does not fully prevent IL loss (Dietz & Stepinski, 2005) but increases the toxicity of the released 889 IL ion (Pham et al., 2010). In addition, the stripping of loaded neutral extractants in IL is problematic 890 and requires aggressive stripping solutions (Regel-Rosocka & Wisniewski). These factors have limited 891 the application of IL containing neutral extractants for the recovery of value from aqueous streams.

One system, the neutral extractant Cyanex 923 dissolved in [N₁₈₈₈][NO₃], has been applied to recover of metals (Co, Ni, Fe, Mn, Zn, REE) from a synthetic Ni metal hydride battery leach solution (Larson & Binnemans, 2015a). Cyanex 923 is a mixture of trialkylphosphine oxides with octyl and hexyl side 895 groups. Addition of 15 % Cyanex 923 (v/v) in $[N_{1888}][NO_3]$ increased the distribution ratio of the test 896 metals and allowed higher extraction efficiencies and Co-Ni separation factors compared to direct 897 application of [N₁₈₈₈][NO₃] alone. Over 98 wt.% of extracted Ni was scrubbed using a MgCl₂ solution 898 prior to stripping Co and Mn from the IL phase using a 3M NaNO₃ solution. In further stripping steps, 899 REE were removed using a 1 M HCl strip before the remaining Fe and Zn were stripped using a 1 M 900 HNO₃ solution. The separation of Co from Mn in the strip solution using $[N_{1888}]$ [SCN] dissolved in 901 [N₁₈₈₈][NO₃] (40% v/v) was also investigated. This research represents an example of "split-anion 902 extraction", whereby the aqueous and organic phases possess different anions, allowing the use of a complexing anion in the organic phase that forms an extractable metal complex. This removes the 903 904 necessity to add a significant concentration of complexing anion in the aqueous phase and facilitates 905 metal stripping (Larson & Binnemans, 2015b). Similarly, Kikuchi and co-workers (2014) extracted 906 REE derived from NdFeB magnets from aqueous nitrate solution using TBP diluted in [N₁₈₈₈][NO3]. Aqueous nitrate added as NaNO₃, was found to salt-in the REE and increase the distribution of the latter 907 908 to the IL phase. The presence of Fe(III) in solution has a negative effect on the overall REE extraction 909 efficiency and selectivity. After removal of Fe from the system, the Nd(III) concentration in the IL phase reached 160 g.L⁻¹ after three loading cycles (Kikuchi et al., 2014). 910

The IL 1,3-dihexylimidazolium nitrate $[C_6C_6im][NO_3]$ was shown to preferentially extract light rare earth elements over 1st row transition metals and this was applied with moderate success to the separation of Sm(III)/Co(II) and La(III)/Ni(II) pairs from nitrate media, a separation relevant to the recycle of these metals from WEEE. Due to the high co-extraction of Co (30%) and to a lesser extent Ni (15%), successive scrubbing steps using 6M NaNO₃ solution were required. Interestingly, $[C_6C_6im][NO_3]$ presents an UCST phase behaviour, with the IL being fully miscible with water at temperature above 90 °C (Depuydt et al., 2017).

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919 6.3. Acidic extractant

920 Acidic extractants are Brønsted acidic molecules containing a protonated functional group such as 921 an alkylcarboxylic acid, alkylphosphoric acid or alkylsulfuric acid group. Coordination of an acidic 922 extractant with a metal cation usually proceeds via deprotonation of the ligand and formation of a charge 923 neutral complex. The pKa of the extractant dictates the ability to deprotonate, thereby influencing the 924 extraction behaviour. By altering the pH of the aqueous solution, selectivity of metal ion extraction can 925 be controlled. In comparison to neutral extractants, extraction behaviour using acidic extractant is easier 926 to predict and back-extraction of the loaded organic phase can be achieved by contacting with an acidic solution and regenerating the extractant (Cocalia et al., 2005; Free, 2013). The full potential of ILs to 927 928 solvate charged species is properly utilised if neutral complexes are extracted. Acidic extractants 929 commonly used in conjunction with ILs are shown in Figure 5.

930 Although widely tested on synthetic aqueous solutions, few studies have used acidic extractants 931 diluted in ILs for the recovery of metals from WEEE. The only example consists of the recovery of 932 lanthanides present in waste fluorescent lamps phosphor using N,N-dioctyldiglycolamic acid 933 (DODGAA) in [C₄C₁im][NTf₂] (Yang et al., 2013). DODGAA, is an anionic extractant possessing a 934 carboxyl and carbamoyl group linked together by an ether chain reported to exhibit high selectivity for 935 REEs, especially the lighter lanthanides (La(III) to Gd(III)) (Shimojo et al, 2014). In a 1st processing 936 step, Y and Eu oxides were leached using a 5 M H₂SO₄ solution at 100 °C. The solid residue was leached 937 a second time using 5 M HNO₃ solution for 6 hours at 100 °C to recover the difficult to extract RE 938 phosphate (La, Ce and Tb). The extraction efficiency of DODGAA in [C₄C₁im][NTf₂] was compared 939 to that of the carboxylic extractant PC-88A in dodecane for the recovery of the REE present in both 940 leach solutions. DODGAA in $[C_4C_1im][NT_{f_2}]$ provided greater selectivity and extraction efficiencies 941 for the REEs than PC-88A at similar pH. Y, Eu, La and Ce were separated from the metal impurities 942 Fe, Al and Zn, although some Ca was co-extracted. Stripping was easily achieved using a dilute acidic 943 solution, and UV spectroscopy of the aqueous phase confirmed minimal loss of the IL cation. The 944 extractant-IL system was recycled and reused five times with no appreciable loss in efficiency (Yang 945 et al., 2013). DODGAA has been shown by other researchers to extract REEs by deprotonation of three 946 ligands and the formation of a charge neutral complex (Shimojo et al, 2014).

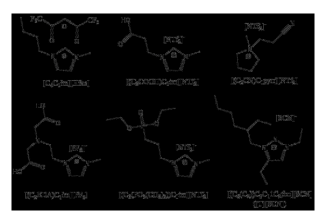
947 Non-task-specific protic ILs can also on occasion extract metals without the need for additional 948 complexing agents. Reyna-González and co-workers (2010; 2012) studied the extraction Cu(II) from 949 an aqueous phase to the protic ILs 2-butylthiolonium bis(trifluoromethanesulfonyl)amide 950 ([(C₄S)C₁im][NTf₂]) and 3-butylpyridinium bis(trifluoromethanesulfonyl)imide ([3-C₄pyr][NTf₂]). 951 During the extraction process the H atom attached to the N atom of the cationic ring is released to yield 952 neutral amine capable of interacting with Cu(II) and water molecules to form a labile copper(II) system. 953 The presence of water in the IL is important to promote incorporation of hydrated Cu(II) complexes. 954 Due to the release of protons to the aqueous phase as well as the hydrophobicity of the cation, release 955 of [3-C₄pyr][NTf₂] to the aqueous phase was not recorded during extraction (Reyna-González et al., 956 2012). High selectivity of Cu(II) compared to Co(II), Ni(II), and Fe(II) was achieved using 957 [(C₄S)C₁mim][NTf₂] (Reyna-González et al., 2010).

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959 6.4. Functionalized ILs

Functionalized ILs, or TSILs, represent ILs incorporating a functional group on their anionic or cationic components that acts as metal complexation site, allowing ILs to function as a solvent and complexing agent. The cation is most often the functionalized moiety due to easier synthesis, which simply requires a quaternization reaction between a 'functionalised' alkyl halide and a nucleophile (tertiary amine, phosphine, etc.). The structure of the TSILs in this section are shown in Figure 11. TSILs with functionalized anions most often include strongly coordinating species such as
alkylphosphate (Rout et al., 2013), alkyldiglycolamate (Rout et al., 2012), β-diketonate (Mehdi et al.,
2010), salicylate, thiosalicylate or benzoate (Stojanovic et al., 2010) anions. TSILs are covered in
greater detail in the reviews by Lee (2006) and Chiappe and Silvio (2014).

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970

971 Figure 11. Selected examples of TSILs used in metal ion extraction.

972

973 The use of ILs as a simultaneous solvent and extraction media limits the ion-exchange extraction 974 mechanism and prevents the unwanted partitioning of the IL components or extractant to the aqueous 975 phase compared to extraction systems where the IL is solely used as a diluent. However, this is only 976 valid for ILs with functionalized cations, as extraction with some IL containing functionalised anions proceeds on occasion via cation-exchange (Medhi et al., 2010). The principal factors limiting the 977 978 applicability of TSILs are their high viscosity, cost, complicated synthesis and unknown toxicity 979 compared to conventional ILs (Visser et al., 2001; 2002). In addition, further understanding of the 980 extraction mechanisms and characterisation of the extracted complex are required, with crystal structure 981 studies being a popular analysis method for TSIL-metal complexes (Harjani et al., 2006; Mehdi et al., 982 2010; Nockemann et al., 2010a; Nockemann et al., 2010b).

983 TSILs incorporating a 'hard' base oxygen derived functional group such as a carboxylate, ketone or 984 phosphonate are used in metal extraction and separation processes. TSILs have been used to recover 985 REE(III) from fluorescent lamps as well as REE(III) and Co(II) from waste permanent magnets. The use of [N_{111(2(0)OH)}][NTf₂] for such applications was partially described in Section 5.1. In addition, 986 [N_{111(2(O)OH)}][NTf₂] was employed for the solvent extraction of scandium. A primary study showed that 987 988 Sc(III) and Fe(III) could be selectively separated from La(III), Ce(III), Nd(III), Dy(III), Y(III), Al(III), 989 Ti(IV), Ca(II) and Na(I) in chloride solutions using [N_{111(2(O)OH)}][NTf₂] as the IL displays a poor affinity 990 towards these elements (Onghena & Binnemans, 2015). The findings from this study were subsequently 991 applied by the same authors to the recovery of scandium from 'sulfation-roasted' leachates of bauxite 992 residue (Onghena et al., 2017a). The distribution ratio of Sc(III) found in this study was much lower

993 than those reported from chloride media (Onghena & Binnemans, 2015). This was assigned to the high 994 sulphate concentration in the leachate and the competing coordination between the sulphate and betaine 995 ligands to Sc(III). As expected, the extraction of Sc(III) by $[N_{111(2(0)OH)}][NTf_2]$ is heavily pH dependent. 996 At the following extraction conditions: pH = 2.4 and O/A = 1.5, 84 wt% of Sc(III) was extracted 997 compared to 23 wt% for Al(III), 4.3 wt.% Fe(III) and 2.6 wt.% Ca(II). The co-extracted metals were 998 scrubbed using dilute HCl solution prior to contacting the IL phase with 3 M H₂SO₄. The composition 999 of the resulting strip solution is 90% Sc(III), 8% Al(III), 0.7% Fe(III) and the lanthanides with 0.5%. 1000 Finally, scandium was precipitated from the strip solution by addition of oxalic acid and calcined for a final oxide product with a purity of 98% (Onghena et al., 2017a). 1001

1002 Chen et al. (2015d) assessed the recovery of Nd(III) from Fe(III) and Sm(III) from Co(II) from 1003 synthetic NdFeB and SmCo permanent magnet leach solutions using 1-alkylcarboxylic acid-3-1004 methylimidazolium bis(trifluoromethylsulfonyl)imide IL ($[(C_nCOOH)C_1im][NTf_2]$ (n = 3, 5, 7), 1005 (Figure 11). Separation of the REE from Fe(III) and Co(II) was found to be primarily pH dependent 1006 and the extraction efficiency of all the investigated metal ions increased with increasing aqueous phase 1007 pH, reaching as high as 99%. In addition, the extraction efficiency increased with a decrease in alkyl 1008 chain length at lower pH. By careful tuning of the aqueous solution pH, Sm(III) and Nd(III) could be 1009 selectively separated from Co(II) and Fe(III) respectively, with separation factors in the range $10^4 - 10^5$. Like the carboxy-functionalised [N_{111(2(O)OH)}][NTf₂], REE extraction proceeds via de-protonation of the 1010 1011 carboxylic group and formation of a 1:3 REE:IL charge neutral complex. The IL was easily stripped in 1012 a single step by dilute aqueous HCl or oxalic acid (Chen et al., 2015d).

1013 Novel 1,2,3-triazolium ILs with [SCN]⁻ or iodide anions (Figure 11) were synthesized for the 1014 extraction of transition metals and REEs from HCl medium (Raiguel et al., 2017). These ILs exhibited 1015 better properties compared to imidazolium ILs, namely low viscosity, low solubility in water and high base stability. The ILs incorporating the 3,5-diethyl-1-(2-ethylhexyl)-4-methyl-1,2,3-triazolium ([T]) 1016 1017 cation with an iodide anion displayed a high extraction efficiency for Cu(II), moderate extraction 1018 efficiencies towards Fe(III), Co(II), Ni(II) and La(III), Nd(III) and Dy(III). After substitution of the 1019 iodide anion for thiocyanate, close to quantitative extraction is obtained for the transition metals and 1020 low to medium extraction of the REE(III). EXAFS analysis of Cu(II) speciation in [T][I] after loading 1021 with CuCl₂ indicates its presence as the [CuCl₂]⁻. Upon extraction, the redox-active Cu(II) is reduced to 1022 Cu(I) whilst I⁻ is oxidized to I₂ which in turn reacts with remaining I⁻ to form I₃⁻. The high base stability 1023 of [T][I] was used in the separation of Cu(II), Zn(II) and Fe(III). Addition of Na₂CO₃ precipitated 98% 1024 of Zn(II) and Fe(III), leaving Cu(II) in the IL.

1026 6.5. Aqueous biphasic systems for metal extraction

1027 An alternative to traditional solvent extraction for the extraction metals is the use of ILs in 1028 aqueous biphasic system (ABS). ABS are mainly composed of water, and can be formed, when 1029 combining in the right proportions, an inorganic salt and a hydrophilic IL, which creates two immiscible 1030 - but reversible - aqueous phases (a salt-rich and an IL-rich phases). Since such systems are obtained 1031 with hydrophilic ILs, the range of potential compounds that can be explored for the extraction is 1032 significantly enlarged and unlocks the full potential of ILs as designer solvents. As the IL-rich phase 1033 consists of an aqueous solution of IL, a much lower amount of IL is required, thus reducing the overall 1034 process cost. In addition, the environmental impact of the use of ILs is diminished as less amounts are 1035 used and hydrophilic ILs are significantly less toxic and more biodegradable than their hydrophobic 1036 counterparts. Finally, ABS overcome the high viscosity of many ILs due to the presence of water 1037 thereby increasing the reaction kinetics and transport phenomena. The extraction of metals using 1038 polyethylene glycol-based ABS is well established (Rogers et al., 1993). Aqueous solutions of ILs and 1039 their combination with ABS have been extensively used with success to extract and purify biomolecules 1040 from biomass (Freire et al., 2012). Despite this, there are few reports on the use of IL-based ABS for 1041 the extraction of metals. One of the main limitations of ABS for metal extraction is the difficult handling 1042 of metal ions with high sensitivity to hydrolysis and/or precipitation as most reported ABS experiments 1043 were carried out using slightly acidic or alkaline solutions, with pH typically ranging from 5 to 13. A 1044 full review of metal extraction using ABS is available elsewhere (Billard, 2016).

1045 An important development in the applicability of ABS extraction to the recovery of metals from 1046 acidic WEEE leachates is the recent proof-of-concept for acidic ABS (AcABS) in which the inorganic 1047 acid itself, in this case HCl, replaces the salt as the driver of ABS formation (Gras et al., 2018). The use 1048 of the leachate solution itself as a salting-out agent allows for the simultaneous leaching and separation 1049 of critical metals, thereby simplifying the extraction and stripping processes. These systems display 1050 several advantages over other extracting solvents. Unlike classical systems based on an organic solvent 1051 and an extracting agent, the IL-based AcABS [P44414]Cl-HCl-H2O exhibits a thermomorphic behavior 1052 with a well pronounced LCST, i.e. the biphasic mixture becomes biphasic by heating above a certain 1053 temperature for a given HCl concentration. This system was applied to the extraction of Fe(III), Pt(IV) 1054 and Co(II) from concentrated HCl solutions and to the separation of the critical transition metal pair 1055 Co(II) from Ni(II). Extraction of Co(II), Fe(III) and Pt(IV) proceeds as in the traditional two-phased 1056 systems based on $[P_{66614}]Cl$ via anion exchange with the IL anion. An additional benefit of this AcABS is the poor miscibility of the IL in the HCl-rich phase, ensuring quantitative recovery of the IL after 1057 1058 extraction. The traditional ABS also based on [P₄₄₄₁₄]Cl with NaCl was shown to selectively separate 1059 Co(II) and Ni(II) in chloride media (Onghena et al., 2015). [P₄₄₄₁₄]Cl exhibits a large biphasic region, 1060 with just 5 wt% NaCl capable of inducing an ABS in a solution containing 40 wt.% IL. The [P₄₄₄₁₄]Cl-NaCl-H₂O system also displays a LCST phase behavior in water. 1061

1062 An interesting IL-surfactant-water microemulsion system based on the IL [P₆₆₆₁₄]Cl and the 1063 surfactant Triton 100 (TX100) was developed extraction and separation of Au(III), Pd(II), and Pt(IV) 1064 from acidic chloride media (Tu Nguyen et al., 2017). A number of factors were assessed including the 1065 microemulsion structure, temperature, surfactant concentration, IL concentration, extraction time, 1066 chloride concentration, and pH. Metal extraction was found to proceed through an anion exchange 1067 mechanism between the anionic chlorometallate complexes and the IL anion. A five-step methodology 1068 was elaborated for precious metal recovery consisting of (i) mixing the feed solution with the IL-based 1069 microemulsion, (ii) inducing TX100-[P₆₆₆₁₄]Cl / aqueous phase splitting by heating to 90 °C, (iii) phase 1070 separation, (iv) stripping and (v) regeneration. Following this simple protocol, 99.9% of precious metals 1071 were selectively recovered at optimized condition. A two-step stripping process was successfully 1072 performed based on stripping of Au(III) using an aqueous solution of sodium thiosulfate followed by 1073 the recovery of Pd(II) and Pt(IV) with a thiourea-HCl solution. The performance of the system after 1074 five extraction-stripping cycles was found to decrease by approximately 9% for all three metals tested.

Similarly to the results reported using the hydrophobic [P₆₆₆₁₄][NO₃] (Vander Hoogerstraete & 1075 1076 Binnemans, 2014), Nd(III) was selectively separated from Co(II) and Ni(II) using an ABS comprised 1077 of [P₄₄₄₄][NO₃] and NaNO₃ as phase former (Chen et al., 2018) based on the ability of lanthanides to 1078 form anionic nitrate complexes. The extraction efficiency was primarily influenced by the aqueous 1079 nitrate concentration, with the Nd(III) extraction efficiency increasing from less than 15 % in aqueous 1080 solution containing no NaNO₃ to quantitative Nd(III) extraction for NaNO₃ greater than 0.3 g.L⁻¹. An 1081 increase in the NaNO₃ content of the aqueous phase also resulted in increased separation factors 10³ 1082 and above when [NaNO₃]=0.6 g.L⁻¹. In contrast, IL concentration had little effect on the extraction 1083 yield: for a 1 mL feed solution containing 1 g.L⁻¹ of Nd(NO₃)₃ quantitative extraction of Nd(III) was 1084 obtained for all IL concentration tested (0.2-0.6 g). After separation of the IL rich phase, extracted 1085 Nd(III) was recovered by contacting with an aqueous solution of sodium oxalate such that the Nd(III) 1086 to oxalate molar ratio was of 1.5. However, a sharp drop in extraction efficiency over repeated cycles 1087 was observed with only 85.4 % of extraction efficiency achieved after only three cycles. NaNO₃ was 1088 found to be a better salting out agent than NH₄NO₃ whilst [N₄₄₄₄][NO₃] has a smaller biphasic region 1089 compared to [P₄₄₄₄][NO₃], in accordance with the more hydrophobic nature of phosphonium cations 1090 over ammonium ones. As with the [P₄₄₄₁₄]Cl system previously described, a strong LCST-type 1091 temperature dependent behavior of the binodal curve was observed as the temperature was increased 1092 from 15 °C to 35 °C.

1093 The ABS composed of the carboxyl-functionalized phosphonium IL $[P_{444(C100H)}]Cl$ (80 wt.%) 1094 and a 16 wt.% NaCl solution was employed as the salting-out agent for the selective extraction of Sc(III) 1095 (Depuydt et al., 2015). Whilst the viscosity of the pure IL was over 20000 cP at room temperature, the 1096 viscosity of the IL-rich phase in the employed ABS was drastically decreased to 400 cP. REEs do not 1097 form anionic chloride complexes, the extraction of Sc(III) was shown to proceed via deprotonation of 1098 three IL-cations (pKa= 2.32) to form zwitterionic compounds that can coordinate scandium(III) ions in

- 1099a 3:1 ratio. The pH was varied from 3.33 to 0 in the aqueous feed solution containing 5 mmol kg⁻¹ of1100Sc and 16 wt % of NaCl to determine the influence of the pH on the solubility of the IL cation in the1101water-rich phase. ¹H NMR analysis of the aqueous phase indicates that the solubility of the IL cation1102increases with decreasing pH from approximately 5.8 wt.% at pH=6.8 to 10.5 wt.% at pH=0. The1103zwitteronic form of the $[P_{444(C100H)}]^+$ is more hydrophobic than its protonated counterpart, thus, a lower
- solubility is observed at higher pH values since low pH values prevent deprotonation of the cation.
- 1105 Quantitative stripping of IL-rich phase was achieved after addition of oxalic acid at a molar ratio of 1.5.
- 1106

1107 **6.6. Hydrophobic deep eutectic solvents (DES)**

Whilst the use of hydrophilic DES is well documented at the laboratory scale for many applications,
the use of hydrophobic DES for the liquid-liquid extraction of metals is relatively recent and opens the
possibility for the substitution of expensive hydrophobic ILs.

1111 A hydrophobic DES composed of a x:1 (x=2-4) molar ratio of decanoic acid and lidocaine was 1112 investigated for the extraction of the transition and alkaline metals Co(II), Zn(II), Ni(II), Cu(II), Fe(II), 1113 Na(I), Li(I) and K(I) from chloride media (Van Osch et al., 2016). The physical properties of the DES 1114 are described, and single element extraction studies show that most transition metal cations are extracted 1115 with high efficiencies although some Fe(II) was found to precipitate and the DES phase forms a gel 1116 after extraction of Mn(II). The loading capacity of the DES is higher than that of corresponding ILs 1117 composed of fatty acids and quaternary ammonium salts. The higher the decanoic acid to lidocaine 1118 ratio, the higher the decrease in the distribution ratio because of the greater hydrophobicity of the DES. 1119 Contrary to what might be expected, the majority of the chloride remains in the aqueous phase. This is 1120 reasoned through the interaction between the protonated lidocaine and the chloride anion: it is likely 1121 that lidocaine hydrochloride is formed, which has a higher water solubility in comparison to lidocaine 1122 itself. Thus, metal extraction proceeds via an ion-exchange reaction in which upon extraction of the 1123 metal ion to the DES phase, protonated lidocaine is transferred simultaneously to the water phase. The 1124 loaded DES phase was stripped using sodium oxalate, and the regenerated DES was recycled. Due to 1125 the loss of lidocaine upon metal extraction, the DES cannot be recycled indefinitely. Although the 3:1 1126 and 4:1 molar ratio decanoic acid to lidocaine DES could be reused a second time without any drop in 1127 Co(II) distribution coefficient, a drastic drop in D for the 2:1 DES was observed. In addition, the water 1128 phase became turbid for all DES ratio.

1129 Quaternary tetraheptylammonium- and menthol-based hydrophobic DES containing carboxylic 1130 acids, lauric acid, decanoic acid, oleic acid and ibuprofen, were investigated for the recovery of indium 1131 from chloride and oxalate media (Tereshatov et al., 2016). DL-menthol:lauric acid based DES were 1132 found to be ineffective extracting agents. However, all $[N_{7777}]^+$ based DES were successful in extracting 1133 In(III), with the $[N_{7777}]$:ibuprofen (7:3 molar ratio) DES displaying the highest distribution ratio. 1134 Furthermore, all $[N_{7777}]$ based DES display the same extraction behaviour in HCl and oxalate solution

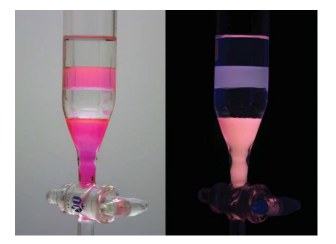
- 1135 respectively, suggesting that the fatty acid alkyl chain does not influence In(III) extraction. In both
- 1136 media, In(III) extraction is dictated by its speciation. Similarly to In(III) extraction in the IL [P₆₆₆₁₄]Cl
- 1137 (Deferm et al., 2016), the DES preferentially extracts the anionic complexes $[InCl_4(H_2O)_2]^-$ and
- 1138 $[In(C_2O_4)_2]^-$ in chloride and oxalate solution respectively (Tereshatov et al., 2016) through ion-pairing
- 1139 with the $[N_{7777}]^+$ cation as deprotonation of the carboxylic acid is assumed to be suppressed at the
- 1140 experimental pH used. Diethylene triamine pentaacetic acid was used to strip the loaded DES.
- 1141

1142 6.7. Ionic liquid – ionic liquid solvent extraction

1143 Systems composed of two different immiscible ILs allow for the development of new separation 1144 processes and the potential for the formation of ternary or quaternary IL systems for extraction as 1145 exemplified in Figure 12.

1146 Arce et al. (2006) and Neves et al. (2017) demonstrated that if large enough structural or chemical 1147 differences exists between two ILs, then this ion mixture may yield a biphasic mixture at room 1148 temperature. For example, mixtures of $[P_{66614}]Cl$ and $[C_nC_1im]Cl$ were found to be 'immiscible' in the 1149 temperature range 298 to 458 K, where the *n* was smaller or equal to 5, with only approximately 10 1150 mol% of imidazolium ion salt dissolved in the phosphonium phase for $[C_2C_1 \text{ im}]Cl$ (Arce et al., 2006). 1151 These findings were expanded upon by Neves et al. (2017) for systems with 4 ions showing that six 1152 pairs of cholinium- and phosphonium-based aprotic ILs ([N_{111(20H)}][NTf₂] and [P₆₆₆₁₄][X] where 1153 X=[Cyanex272], $[C_9COO]^2$, $[NTf_2]^2$, $[N(CN)_2]^2$, Br⁻ and Cl⁻) were able to form biphasic systems. The 1154 formation of an IL liquid-liquid system is cation driven: only mixtures of two ILs with significantly 1155 structurally different IL cations or of highly different hydrogen-bond acidity can undergo liquid-liquid 1156 demixing. The IL anion does not have a significant impact on the creation of a binary systems but plays 1157 an important role in defining the mutual solubility magnitude, which was suggested to linearly depend 1158 on the volume of the IL anion. In all cases except for systems formed by ILs with a common anion, 1159 significant ion exchange occurs, the extent of which correlates with the ILs cohesive energy (Neves et 1160 al., 2017).

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1163

Figure 12. A stable tetraphasic mixture of (from top to bottom): pentane, $[P_{66614}][NTf_2]$, water and [C₂C₁im][NTf₂]. The pink colouration is due to rhodamine B dye, which is soluble in the ionic layers. The figure on the left was taken in daylight and the figure on the right was taken under illumination from UV light (254 nm). Reproduced from Arce et al. (2006) with permission from The Royal Society of Chemistry.

1169

1170 This principle was successfully used to separate cobalt from nickel and REEs from nickel 1171 respectively using the biphasic system $[C_2C_1im]Cl$ and $[P_{66614}]$ -bis(2,4,4-trimethylpentyl)phosphinate 1172 (Wellens et al., 2013; out et al., 2014). A solubility of 9.2 wt% (or 31 mol%) of $[C_2C_1im]Cl$ in 1173 $[P_{66614}][P_{RROO}]$ was determined from ¹H-NMR peak analysis peaks. The solubility of $[P_{66614}][P_{RROO}]$ in 1174 $[C_2C_1im]Cl$ was found to be below the detection limit (Wellens et al., 2013).

1175 Vander Hoogerstraete and co-workers (2015) demonstrated a proof of principle for the separation of tin(II), scandium(III) and yttrium(III) using the ternary IL system composed of [N_{111(2(O)OH)}][NTf₂]-1176 H₂O-[P₆₆₆₁₄][NTf₂]. A ternary system was required for this separation as it is known that tin(II) and 1177 scandium(III) extraction by carboxylic ligands display the same pH dependent behaviour. 1178 $[N_{111(2(O)OH)}][NTf_2]$ was selected over $[N_{111(2OH)}][NTf_2]$ as one of the phases due to its significantly larger 1179 1180 triphasic region. Initial biphasic tests showed that Sc(III) and Sn(II) could be separated from Y(III) in 1181 the [N_{111(2(O)OH)}][NTf₂]-H₂O system at pH 1.0 whilst 80% of Sn(II) was extracted to the IL phase in the [P₆₆₆₁₄][NTf₂]-H₂O in the presence of the neutral tributylphosphate (TBP) ligand whilst the REEs 1182 1183 remained mainly in the aqueous phase. A pH below 1, along with the addition of 1M MgCl₂ to the 1184 aqueous phase, was required to suppress tin hydrolysis and prevent [P₆₆₆₁₄][NTf₂]-TBP emulsions. In 1185 the ternary system, the best separation was obtained at low chloride concentrations without pH 1186 adjustment and at high TBP concentration in the [P₆₆₆₁₄][NTf₂] phase. Under these conditions (20 wt.% 1187 TBP, 0.25 M MgCl₂), 89 % of tin was extracted to the [N_{111(2(O)OH)}][NTf₂] phase, 89% of Y(III) remained 1188 in the aqueous phase and 85% of Sc(III) was extracted to the $[P_{66614}][NTf_2]$ phase. Although not 1189 conclusive, these encouraging results underline the versatility of ILs and the possibilities available for 1190 IL-IL liquid-liquid extraction.

1191 The IL-IL mineral separation of rare-earth minerals (bastnäsite and monazite) from Ca, Mn, Fe and 1192 Mg rich samples (ankerite, dolomite and calcite) was attempted (Azizi & Larachi, 2018). Specifically, 1193 [N4444][DEHPA] was added as a droplet phase in three different ILs used as a continuous phase, 1194 ethylammonium nitrate ([EtNH₃][NO₃]), [C₂C₁im][SCN] and [C₄C₁im][BF₄]. These ILs were selected 1195 as i) [N₄₄₄₄][DEHPA] displays a far superior affinity for the REEs compared to the other ILs, ii) the 1196 droplet phase viscosity is lesser than that of the continuous phases and iii) the droplet phase and 1197 continuous phase must display minimal solubility with one-another. The optimal system of 1198 [N₄₄₄₄][DEHPA] dispersed in [C₂C₁im][SCN] was shown to outperform froth flotation and traditional 1199 liquid-liquid mineral separation for the recovery of RE minerals. DFT analysis found that anionic 1200 moiety of $[N_{4444}]$ [DEHPA] directly interacts with the metal cations on the mineral surfaces through 1201 covalent bonds resulting in very favourable interaction energies compared to the other ILs.

1202 Although not technically an IL-IL extraction system, the solvent extraction of Co, Fe and B from 1203 waste NdFeB magnets in DES [N_{111(20H)}]Cl:lactic acid using [N₁₈₈₈][SCN] was successfully achieved and scaled-up over multiple mixer-settlers (Riaño et al., 2017). The dissolution of Co, Fe, B, Nd, Dy, 1204 1205 Pr and Gd oxides from crushed and roasted rare-earth permanent magnets (NdFeB) was attempted using 1206 three different DES: [N_{111(20H)}]Cl:urea (1:2), [N_{111(20H)}]Cl:EG (1:2) and [N_{111(20H)}]Cl:lactic acid (1:2). At a liquid to solid ratio of 20 mL.g⁻¹, 70 °C leaching temperature and 24 hr leaching time, over 80 1207 1208 wt.% of all the target elements were lixiviated using $[N_{111(20H)}]$ Cl:lactic acid and little to no dissolution 1209 occurred in [N_{111(20H)}]Cl:urea and [N_{111(20H)}]Cl:EG. This contrast in leaching efficiency was assigned to 1210 the ability of the lactic acid moiety to deprotonate and form the metal complex M(CH₃CHOHCOO)_{n,DES} 1211 (where n is the metal oxidation state). Interestingly, whilst the water content had no influence on the 1212 leach efficiency of the non-roasted magnet, it does have an important influence on the leaching of the 1213 roasted oxides with a decrease in leaching efficiency with increasing water content. Almost no Fe and Co are recovered at 80 wt.% (L/S=12.5 mg.L⁻¹, 70 °C, and 24 hr) water content due to their hydrolysis. 1214 1215 Recovery of the target metals from the DES [N_{111(20H)}]Cl:lactic acid (1:2) without water is achieved 1216 using a two-stage solvent extraction procedure. In this procedure, the DES phase was employed as the 1217 more polar phase for solvent extraction and the IL $[N_{1888}]$ [SCN] diluted in toluene as the less polar 1218 phase. Co(II), Fe(III) and B(I) were separated from the REE(III) in an initial solvent extraction stage using $[N_{1888}]$ [SCN]. EXAFS analysis indicates that cobalt is extracted as the complex $[Co(SCN)_4]^{2-}$ in 1219 1220 $[N_{1888}]$ [SCN] whilst iron is extracted as $[N_{1888}]$ [FeCl4]. In a second stage, Nd was separated from Dy 1221 using the acidic extractant Cyanex 923 in toluene. Cyanex 923 was selected ahead of D2EHPA due to 1222 the difficulty in stripping Dy(III). The less polar phases were recycled and reused a second time with 1223 little loss in efficiency although boron extraction declined due to presence of chloride anion in 1224 [N₁₈₈₈][SCN] resulting from anion-exchange.

1226 7. Critical perspective and recommendations – ILs as specialty solvents

1227 **7.1. Ionic liquids – the case for application-specific solvents**

1228 The publications reviewed here demonstrate the growing application of ILs for the treatment of 1229 WEEE. Despite these promising results, few studies have made the jump from laboratory to industrial 1230 scale. In this section, the major obstacles currently preventing the use of ILs in metal extraction 1231 processes are highlighted. The selection of ILs is usually justified based on their green credentials, 1232 namely their tunability, low vapour pressure, non-flammability and high stability. However, ILs also 1233 suffer from some important disadvantages preventing their industrial application such as their high 1234 viscosity, corrosiveness, hygroscopicity, toxicity and their higher cost compared to molecular solvents. 1235 As such, a more nuanced perspective is required when considering the use of an IL and the following 1236 issues must be addressed prior to any successful application. The discussion is summarised in Figure 1237 13. All data here presented is qualitative and should be used just as a guideline. Based on the guidelines presented in Figure 13 and the literature reviewed in the previous Sections, optimised process 1238 1239 flowsheets are proposed for the valorisation of the inherent metallic value in fluorescent lamp phosphor 1240 waste (Figure 15) and nickel metal hydride battery waste (Figure 16).

1241

	hla						Dueferred
Undesir	ible						Preferred
		Toxici	ty – conventi	ional so	lvents		
Dichloromethane Hexane, Benzene,	•	Xylene, Toluene, Heptane, Cyclohexane, Isooctane			Water, Supercritical CO _{2,} Ethanol, Acetone, Ethyl acetate,		
Toxicity – IL anion							
		[BI	[BF ₄] ⁻ , [SCN] ⁻ , [N(CN) ₂] ⁻ ,			[Cl] ⁻ , [NO ₃] ⁻ , [C ₁ SO ₄] [,] [C ₁ COO] ⁻ ,	
[NTf ₂] ⁻ , [PF ₆] ⁻		[C ₇ H ₇ O ₃] ⁻ , [C ₈ PO ₄] ⁻			[C ₃ H ₆ O ₃] ⁻ , [C ₁₂ SO ₄] ⁻ , [C ₁₇ COO] ⁻		
Toxicity – IL anion							
$[C_n C_1 im]^+$ (n \ge 12)	$[P_{66616}]^+$	$[\mathrm{C_4C_1im}]^+$	[N ₁₈₈₈]	+	[C ₄ pyr] ⁺	[C ₃ CNpyr]	$[N_{111(2OH)}]^+$ $[N_{111(2(0)OH)}]^+$
Cost - IL							
Cation	А	nion [C ₄ C	C1im][SCN]	$[C_4C$	1im][NTf2]	[C ₁ C ₁ mim][HS0	O ₄] [N _{111(2OH)}]Cl:EG
functionalised TSII	nctionalised TSIL functionalised TSIL		$[P_{66616}]Cl$ $[C_4C_1]$		im]Cl [N ₁₈₈₈]Cl		
Metal Prices							
Al Pb Zn La ₂ O ₂	, Mn Cu Y	₂ O ₃ Sb Ni Sn	Gd ₂ O ₃ Nd ₂ O	O ₃ Co	Se Ta Eu ₂	O ₃ Dy ₂ O ₃ Tb ₂ O ₃	3 Ag In Au PGM

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• How to design ILs more environmentally benign than currently used molecular solvents?

Figure 13. Qualitative toxicity and cost scale for the judicious selection of ILs for metal extraction

1246 The structural modifications that confer hydrophobicity to ILs are the same that increase their 1247 toxicity. Thuy Pham and co-workers (2010) reviewed the toxicity of common ILs across multiple 1248 trophic levels and found that toxicity seemed to be determined mainly by the cationic component and 1249 that it correlates directly with the length of the cationic alkyl-chain(s). In addition, ILs incorporating 1250 polar functional groups in the alkyl side chain such as ether, hydroxyl and nitrile moieties displayed 1251 lower cytotoxicity compared to those with simple alkyl side chain. Regarding the anionic effects, 1252 fluorinated anions, particularly $[NTf_2]^-$, are generally more toxic than their halogenated counterparts 1253 (Viboud et al., 2012). In addition, some fluorinated anions have been shown to hydrolyse and can 1254 release the toxic product HF as a result of decomposition (Swatloski et al., 2003; Freire et al., 2010). 1255 The authors show that $[C_4C_1im][PF_6]$ and $[C_4C_1im][BF_4]$, two of the most widely used ILs, decompose 1256 in contact with water. The decomposition reaction leads to the formation of toxic HF.

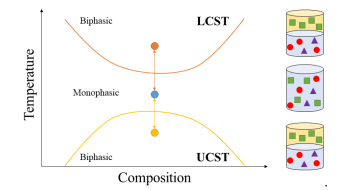
1257 The "greenness" of molecular solvents is well documented according to multiple criteria. An 1258 established guide to green solvent selection is the 'Pfizer solvent selection guide' which classifies 1259 common organic diluents like hexane, benzene or petroleum spirits as undesirable (Byrne et al., 2016). 1260 In this context, it makes little sense to label the substitution of hexane in the organic phase by 1261 [C₄C₁im][NTf₂] as environmentally friendly. 'Greener' alternatives include the substitution of 1262 imidazolium ILs to pyridinium-based ILs, the use of large fatty acid as anions over fluorinated anions 1263 to confer the IL hydrophobic properties and – if possible – the application of DES. Water-miscible ILs 1264 with shorter cationic alkyl chains can be salted-out to form a reversible biphasic system suitable for 1265 metal extraction. It is also worth considering that many ILs act both as the hydrophobic phase and 1266 extractant, thereby potentially reducing their over whole toxicological impact. Aliquat 336 ([N₁₈₈₈]Cl) stands out as such an example. However, IL systems in which metal extraction proceeds via a cation-1267 1268 exchange mechanism should be avoided.

1269

Can ILs be designed to offer an improvement compared to existing industrial practices?

In other words, is there any advantage in using IL over existing solvents, either through increased 1270 1271 selectivity, more favourable kinetics and thermodynamics or through process simplification? E-waste 1272 represents a variable and heterogeneous waste stream, with product composition changing between 1273 manufacturers and product iterations. Current industrial hydrometallurgical methods for the recovery 1274 of metals from e-waste borrow heavily from tried and tested processes established in the mining industry 1275 for the extraction of metals from ores (Cui & Zhang, 2008). Any proposed IL-based approach needs to 1276 be robust and flexible to adapt to such variable conditions. Many of the studies reviewed here report 1277 distribution and separation factors superior to those obtained in conventional organic systems. 1278 Problematically, most of these IL studies only focused on simulated systems. Such an approach risks 1279 missing interactions that occur only in complex systems such as in leachates from WEEE from a waste 1280 treatment facility and ignores the role of trace contaminants. Physical separation processes, the 1281 preliminary step in all waste treatment facilities (Figure 1), are never 100% efficient due to the overlap

- between different physical properties, with impurities being entrained in the output stream. An example of a problematic trace contaminants is the extraction of Fe(III) from chloride solutions by phosphoniumbased ILs. Fe(III) is strongly and efficiently extracted from even dilute chloride solutions and its stripping can prove problematic (Cui et al., 2015). This in turn can complicate the regeneration of the IL and lower the extraction efficiency of target metals.
- 1287 The high viscosity of ILs and especially DES is a well identified barrier to their application due to the 1288 increased cost of pumping and slow mixing processes and kinetics of mass transfer. Certain ILs, 1289 particularly those that operate by an anion-exchange mechanism, are known to significantly increase in 1290 viscosity after metal extraction due to strong intermolecular interactions. For example, the viscosity of 1291 water saturated [P₆₆₆₁₄]Cl at 60 °C increases from 95 cP to approximately 800 cP after contacting with 1292 a feed solution containing 40 g.L⁻¹ Co(II) (Vander Hoogerstraete et al., 2013). The problem of viscosity 1293 can be partially overcome by saturating the IL with water and operating at higher temperature and/or 1294 lower feed concentrations. ABS and thermomorphic ILs fully overcome the issues associated with 1295 viscosity by tuning their water miscibility through varying the salt concentration and temperature 1296 respectively.
- This tuneable and reversible water miscibility is extremely desirable in homogeneous liquid-liquid extraction. Performing the extraction in a monophasic solution prior to separation of the phases through a change in temperature reduces the viscosity of the solution and accelerates mass transfer, resulting in higher extraction efficiencies. This is schematically illustrated in Figure 14; ILs can either display LCST or an UCST.
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1303

Figure 14. Schematic illustration of LCST or UCST behaviour and its application to homogeneous
 liquid-liquid extraction.

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1307 The synthesis of ILs exhibiting thermomorphic behaviour requires a balance between the 1308 hydrophilicity and hydrophobicity of the ions. The appearance of an LCST for [P₄₄₄₄]- and [N₄₄₄₄]-based 1309 ILs was shown to vary with anion selection (Khono & Ohno, 2012). Depuydt and co-workers (2017b)

recently synthesised a range of phosphonium IL with dioctyl sulfosuccinate (DOSS) as anion. Whilst 1310 1311 [P₄₄₄₄][DOSS] is hydrophobic, incorporation of three ethylene glycol moieties to the cation lowers the 1312 hydrophobicity of [P_{444(EG3)}][DOSS]. [P_{444(EG3)}][DOSS] has an LCST of 19 °C in water and was shown 1313 to efficiently extract both divalent and trivalent metals. Furthermore, the presence of additional ions in 1314 solution can alter the LCST or UCST temperatures due to the strong polar/electrostatic interactions between the water, IL and the ions. The UCST and LCST of [N_{111(2(0)OH)}][NTf₂] and [P₄₄₄₁₄]Cl 1315 1316 significantly decrease after loading with REEs (Schaeffer et al., 2016) and after addition of NaCl to the 1317 aqueous phase respectively (Onghena et al., 2015). The ability to separate an aqueous IL solution into 1318 two phases by varying the temperature or by addition of salting-out salts allows for the potential 1319 development of metal recycling processes combining leaching and extraction of metal ions.

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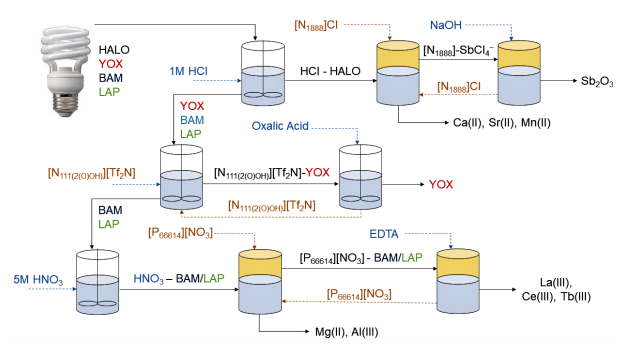
Are IL-based processes economically viable?

1321 The most cited of all barriers to IL application is their high initial cost; ILs normally fall in the range 1322 of 5 –20 times more expensive than molecular solvents (Plechkova & Seddon, 2008). Although prices 1323 for small quantities should not be used as a guide to commercial utility, at the time of writing the retail 1324 price on the Sigma-Aldrich website for $[C_4C_1im][SCN]$, $[C_4C_1im][NTf_2]$, $[C_4C_1im][C_1COO]$ and $[C_4C_1im]Cl$ is €2880 kg⁻¹, €1505 kg⁻¹, €776 kg⁻¹ and €288 kg⁻¹ (Sigma Aldrich, 2017). The economic 1325 1326 sustainability of ILs is greatly improved if the IL is recycled. Due to their low vapour pressure and 1327 thermal and chemical stability, ILs are theoretically recyclable over numerous application cycles. It is 1328 estimated that 10 to 20 recycle and application cycles gives them the same cost per cycle as conventional organic solvents, and over 50 recycles makes them significantly cheaper (Plechkova & Seddon, 2008). 1329 1330 As such, all research should aim to demonstrate the stability and recyclability of their ILs over multiple 1331 cycles. In addition, the solubility of well-known hydrophobic fluorinated ILs notably increases in the 1332 presence of inorganic acids such as HNO₃, HClO₄ and HCl (Fu et al., 2016, Mazan et al., 2016). This 1333 can lead to the irreversible loss of the IL to the aqueous phase during solvent extraction, lowering the 1334 environmental and financial sustainability of the process. A mass balance should be performed at each 1335 stage of the process to quantify overall IL losses and to determine which operation(s) primarily 1336 contribute.

1337 However not all ILs are expensive, especially at an industrial scale. Chen and co-workers (2014) demonstrated through process modelling that the price of the Brønsted acidic IL triethylammonium 1338 1339 hydrogen sulphate ($[N_{222}][HSO_4]$) and 1-methylimidazolium hydrogen sulphate ($[HC_1im][HSO_4]$) on an industrial scale was of \$1.24 kg⁻¹ and \$2.96–5.88 kg⁻¹, making these significantly cheaper than 1340 1341 fluorinated ILs. In addition, DES along with tetraalkylammonium and tetralkylphosphonium salts are 1342 intrinsically cheaper than imidazolium-based ILs (Plechkova & Seddon, 2008). [N1888]Cl and [N_{111(20H)}]Cl (the starting component of most DES) currently retail at €92 kg⁻¹ and €58 kg⁻¹ respectively 1343 1344 (Sigma Aldrich, 2017). As in all cases, a compromise must be made between the IL/DES selection 1345 (cation and anion selection, functionalisation, etc.), its intended application (leaching, extractant,

hydrophobic diluent, medium for electrodeposition, etc.) and the price of the target metal. Whilst the use of more expensive IL can be economically justified for the recovery of In, Ag, Au, PGM and some rare earth elements, the application of fluorinated ILs for the leaching of La_2O_3 makes little economic sense.

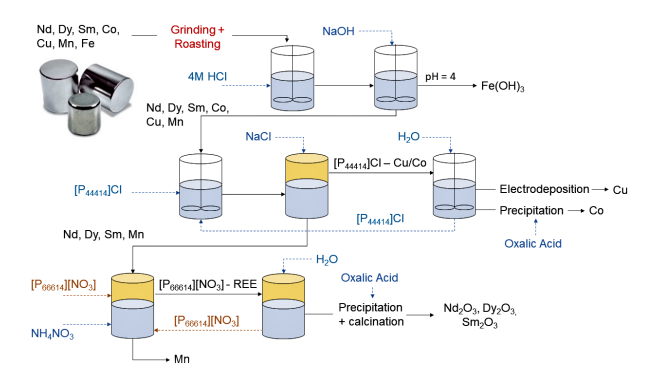
1350 The IL-based solvents covered in this review are for the most part limited to the recovery of value 1351 from PCBs, waste fluorescent phosphor, nickel metal hydride batteries and permanent magnets. To 1352 highlight the great strides made over the last years on the potential of IL-derived solvents for WEEE 1353 treatment, optimised process flowsheets are proposed for the extraction of value in fluorescent lamp 1354 phosphor waste (Figure 15) and nickel metal hydride battery waste (Figure 16). The versatility of 1355 quaternary phosphonium- and ammonium-based ILs for metal extraction based on the selective 1356 formation of anionic metal complexes in various leaching solution (chloride, nitrate, etc.) is fully 1357 exploited. When possible, solvent extraction processes were substituted for their ABS counterpart based 1358 on similar IL cations to further reduce the environmental impact and take advantage of the 1359 thermomorphic properties of phosphonium ILs. The substitution by ABS also simplifies the metal 1360 recovery process as electrodeposition of easily reduced metals or precipitation are possible directly from 1361 the IL-rich phase. The use of the more expensive $[N_{111(2(O)OH)}][NTf_2]$ is justified based on its great 1362 selectivity for YOX phosphor dissolution and the high price of the recovered Eu(III). Through careful analysis of the waste matrix, selective sequential leaching steps can be designed in order to solubilise 1363 1364 elements of interest at each stage and simplify their purification. Furthermore, the addition of pre-1365 treatment steps such as mechanical activation, conversion of metals to oxides and/or the hydrolysis of 1366 Fe(III) greatly increases the recovery yields and simplifies the downstream purification of the other anionic chlorometallates. In all cases, closed-loop systems are suggested to recycle the IL, thereby 1367 1368 improving the ecological and economic aspects of the proposed processes.



- 1371 Figure 15. Suggested simplified process flow diagram for the recovery of antimony and REE from
- 1372 waste phosphor.

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1375 Figure 16. Suggested simplified process flow diagram for the recovery of cobalt, copper and REE from

1376 waste NdFeB magnets.

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1378 **8.** Conclusions

1379 WEEE is a global problem issue but also a potentially opportunity due to its high content of valuable 1380 metals. The physico-chemical characteristics of ILs mean they have potential applications in the 1381 recovery of metals from WEEE. Many experimental and theoretical studies have identified appropriate 1382 IL systems for the selective extraction and purification of metals from WEEE. The extraction 1383 performance of many IL systems has been characterised, allowing selection of a particular IL for a 1384 specific metal possible. Quaternary ammonium ILs, DESs and ILs-based ABS were highlighted as the 1385 most promising solutions due to their low environmental impact, competitive price, compositional 1386 flexibility and operational facility. However, significant barriers remain to commercial exploitation of 1387 ILs for metal recovery from WEEE. There is a lack of IL application to actual WEEE waste streams, as 1388 many researchers have worked on simulated systems and this risk missing interactions that occur with 1389 trace components present in WEEE. A further barrier is the high initial cost of many ILs and particularly 1390 fluorinated ILs. Despite high recyclability, the capital investment required has limited the use of ILs to 1391 extraction of high value metals from WEEE and use in systems were a high metal purity is achieved. 1392 Improved upstream collection and sorting to minimise the volume of commingled WEEE would allow 1393 more selective use of ILs.

1394

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1399 1400

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