Halometallate ionic liquids: thermal properties, decomposition pathways, and life cycle considerations†

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Halometallate ionic liquids provide new opportunities for industrial catalytic processes because of their unique blend of physical and chemical properties. Tunability underpins the success of ionic liquids because small structural changes can have drastic effects on either property. Catalysis can be optimised by adjusting structures to target properties such as Lewis basicity and acidity, but the structural changes have simultaneous impacts on physical properties. In this work, we provide a thorough, methodical, and reliable list of thermal parameters to help define temperature limits to prevent catalyst poisoning and limit the need to replace costly and environmentally demanding solvents. Mechanistic insights show that decomposition is particularly detrimental for halometallate ionic liquids, and life cycle analysis highlights that lower levels of organic cations are better for economic and environmental sustainability.

1. Introduction

Ionic liquids (ILs) can dissolve large amounts of metals while retaining the free-flowing properties of liquids. The resulting metal ionic liquids are known to be highly tuneable as speciation of the metal can be controlled by the concentration of the metal, the identity of the anions, and the identity of the cations.1 Catalysis is by far the most popular application of ionic liquids currently being adopted by industry,2 in part because of excellent tunability of metal centres, but also because the properties of the ionic liquid medium opens up new avenues for novel and established catalysts. Examples of beneficial properties include: low vapour pressures which eliminate solvent loss and enable gas/liquid phase processes, diverse structures that improve separations, and high chemical and thermal stabilities which reduce decomposition of the liquid phase and provides access to conditions unsuitable for traditional molecular solvents.3 However, not all ionic liquids are equal because of the vast diversity in their chemical structures, which is why studying and understanding ionic liquid physical properties is key to their success. Furthermore, altering part of an ionic liquid structure to achieve a specific task can have subsequent effects on other physical or chemical properties, making design a difficult process and further warranting the study of properties critical to industrial processes.4

Thermal stability is a particularly important property for halometallate ionic liquids because exceeding the upper temperature limit produces decomposition products that can poison catalysts, react with reagents and products, and change the physical properties of the medium. Degraded solvents also need to be replaced or replenished, which adds a further environmental burden because ionic liquid production involves many steps, each with energy requirements and emissions of their own.5 Recycling is often suggested as a way to offset the environmental burden of ionic liquids, but reuse depends on preventing decomposition to maintain the performance of the catalysts – waste prevention often being the leading principle of green chemistry. These issues are especially pertinent in industrial settings, where scales are larger and prolonged lifetimes make ionic liquids more economically feasible. Furthermore, highly corrosive and harmful decomposition products (such as HF) can evolve from overheated ionic liquids, which adds a serious safety concern and provides further justification for studying decomposition pathways.6,7

Many emerging applications of ionic liquids exploit the unique interactions between cations, anions, and metals centres. For example, new inorganic Zn nanomaterials can be prepared directly from metallic Zn using highly coordinating ionic liquids to give a more cost-effective and environmentally

† Electronic supplementary information (ESI) available: Synthesis and characterisation, thermal analysis data, supporting NMR and MS data, DFT and LCA data. See DOI: https://doi.org/10.1039/d2gc01983c
friendly production process, a type of “controlled corrosion”. Novel soft materials can also be prepared directly from Lewis acidic metal ionic liquids, which are able to influence polymer architectures to provide highly flexible gels with unique surface compositions. Both examples utilise opposite ends of a donating/accepting scale but intermediate properties can be accessed by design. Halozincate-based ionic liquids are of particular interest in this work because they are air and moisture stable, speciation is well known, and the Lewis basicity/ acidity can be controlled by changing the ratio of the organic ionic liquid and metal salt. For example, mixing $[C_6C_1Im]Cl$ and ZnCl$_2$ at a ratio of 2:1 gives the tetrachlorozincate(II) anion $[ZnCl_4]^{2-}$, but with each successive addition of ZnCl$_2$ polyanionic species grow as linear chains (Fig. 1), which subsequently increases the Lewis acidity of the metal centres. These and other halometallate systems have previously been quantified in terms of Lewis acidity through Gutmann acceptor numbers (ANs) and photoelectron spectroscopy. Some studies have also started to report thermophysical properties such as viscosity, but a systematic/methodical analysis of thermal properties is absent from the published literature.

Here, MX$_x$ (where M = Zn and X = Cl$^-$ or Br$^-$) was dissolved in ionic liquids with identical or mixed anions to give $[C_6C_1Im]X_M X_x$, where the mole fraction ($y$) was varied between 0–1. Other metal centres (Co, Ni, Pt, Ag) were also studied for $\chi = 0.33$ (i.e., one third of the solution was MX$_x$ or AgX) to understand how thermal properties change as a function of the metal. We present a particular focus on high temperature stability because many new applications utilise elevated temperatures for catalytic processes or material applications (e.g., 150–250 °C). We also report solid–liquid transitions, which are important because they present a lower temperature limit, and together with thermal stability, define an operating range for the liquid medium. We report other industrially-relevant thermal parameters, such as long term thermal stabilities and heat capacities ($C_p$), which are used to estimate energy requirements for heating. Using hyphenated and hybridised TGA techniques and ex situ analysis, we also present a mechanistic insight to halometallate ionic liquid thermal decomposition and show why overheating metal containing ionic liquids is particularly detrimental, making thermal studies of metal ionic liquids a critical aspect for their adoption to industry. Finally, we present life cycle analysis (LCA) data for the production of chlorozincate ionic liquids as a function of mole fraction and discuss the environmental and economic impacts in the context of solvent and catalyst stability.

2. Experimental

2.1 General procedures

Details of synthesis and purity are presented in the ESI.$^\dagger$ All halometallate ionic liquids prepared in this work were found to be room temperature ionic liquids (RTILs), with the exception of $[C_6C_1Im]Cl_{0.3}PtCl_2$ which was isolated as a brown solid (note: the $[PtCl_4]^{2-}$ anion is square planar). Although all ionic liquids were prepared under an inert atmosphere in a glovebox, the samples were momentarily exposed to air before analysis. All ionic liquids studied in this work have previously been found to be air and moisture stable from loading samples onto sample bars in air before analysing with a range of X-ray spectroscopies that show speciation is unchanged (i.e., no oxidation/decomposition). Any residual water absorbed from the air before thermal analysis was removed from the sample by in situ drying (details below).

2.2 Thermal analysis

Differential scanning calorimetry (DSC) experiments were conducted on a TA Instruments Discovery DSC2500 equipped with an RCS-90 chiller using 1–5 mg of sample. The DSC was calibrated before use by baseline conditioning, temperature calibration, cell constant, reversing heat capacity calibration, and MDSC calibration with indium and sapphire calibration standards. Dry nitrogen gas was used for all experiments at a flow rate of 50 mL min$^{-1}$, unless stated otherwise. Samples were prepared in sealed aluminium Tzero hermetic pans with pin hole lids to allow an initial drying step (120 °C for 45 min) to drive water from the samples (i.e., in situ drying) and to erase the thermal history of the samples. Glass transition temperatures ($T_g$) were measured by the midpoint method at half height on heating cycles at 10 °C min$^{-1}$.

Heat capacities ($C_p$) were measured with the same DSC2500 instrument in sealed aluminium Tzero pans with pin hole lids with 15 mg of sample. The same pre-drying step (120 °C for 45 min) was used before the instrument was switched to quasi-isothermal modulated DSC (QIMDSC) mode (temperature amplitude = 1.00 °C and period = 120 s) and $C_p$ values were recorded in 20 °C increments from 20 °C to 140 °C. Data was recorded at each temperature step over a 10 minutes iso-thermal window and the average $C_p$ value was reported, along with the standard deviation as a measure of error. To assess reproducibility, three separate sample pans containing $[C_6C_1Im][NTf_2]$ were analysed and the $C_p$ (25 °C) was 714.0 ± 0.8 J mol$^{-1} \degree$C$^{-1}$ and the $\Delta C_p$ (140–25 °C) was calculated to be 98.1 ± 1.2 J mol$^{-1} \degree$C$^{-1}$ (note: the error is the standard devi-

Fig. 1 Structures of (a) $[MC]Cl_2$– anions and polyatomic ($Zn,CiCl_2x,y$)$^{2-}$ anions with mole fractions of metals and abbreviations, (b) 1-alkyl-3-methylimidazolium and trihexyltetradecyl-phosphonium cations.
Thermogravimetric analysis (TGA) was measured on a TA Instruments Discovery TGA 550 with high temperature (HT) platinum pans and a gas flow rate of 50 mL min\(^{-1}\). A two-point temperature calibration with Zn and Alumel was carried out before analysis. For organic samples, Pt pans were cleaned by heating up to 1000 °C for 1 hour in 100 mL min\(^{-1}\) air, which also ensured the TGA furnace was regularly cleaned of decomposition residues (*note*: regular exhaust cleaning is necessary for consistent sample purge rates). For metal-containing samples, Pt pans were chipped free of carbon residues and submerged in 37% HCl at 60 °C and left to stir for 24–72 h before being mechanically exfoliated in a sand bath. After washing with acetone and thoroughly drying, any remaining residue was burned away using a blowtorch for 2–3 min.

TGA-mass spectrometry (TGA-MS) was measured on a TGA/DSC 11E/UMX Mettler Toledo Instrument (Al, Zn, and In temperature calibrated) combined with a HR20-QIC Hidden Analytical mass spectrometer. Platinum pans were dried with an inert \(\text{N}_2\) purge at 50 mL min\(^{-1}\) and samples were dried isothermally at 100 °C for 45 min before ramping at 10 °C min\(^{-1}\). Simultaneous thermal analysis (STA) was measured on a NETZSCH Jupiter 449F5 instrument calibrated with Zn and In, and a 50 mL min\(^{-1}\) \(\text{N}_2\) purge was used along with Pt crucibles with a heating rate of 10 °C min\(^{-1}\).

Kinetic analysis was carried out through either the Flynn–Wall isoconversion method,\(^{24}\) modulated TGA (MTGA) method,\(^{25}\) or an isothermal method.\(^{26-28}\) Both Flynn–Wall and MTGA employ variable heating, but MTGA uses temperature modulation in a single experiment as opposed to several separate experiments. For MTGA experiments, a modulation temperature of 5 °C and a period of 200 s were used with Hi-Res ramping at 2 °C min\(^{-1}\), a resolution of 6.00, and sensitivity of 1.00. Activation energies \(E_a\) were calculated over a 20 min window centred around 10% mass loss to ensure enough cycles had passed: the average and standard deviation of the continuous \(E_a\) signals are reported. Previously, we calculated the MTGA \(T_d\) of commercial \([\text{C}_2\text{C}_1\text{Im}]^+\text{NTf}_2^-\) \((127.2 \pm 2.1 \text{ kJ mol}^{-1})\) to be only 1.7 kJ mol\(^{-1}\) lower than a published value calculated from the isoconversional method \((131 \text{ kJ mol}^{-1})\),\(^{26}\) which showed that MTGA was a rapid alternative when studying ionic liquid decomposition kinetics. The Flynn–Wall and MTGA methods are both model free and assume first order kinetics, while the isothermal method assumes zero order kinetics.

Because of the difficulty in identifying decomposition pathways, we have analysed several samples in this work by multiple kinetic analyses. A recent review by Xu et al. gives a good overview of the different isoconversional and Arrhenius kinetics analysis methods.\(^{29}\)

### 2.3 DFT calculations

DFT calculations were carried out using the Gaussian 16 (version C.01) suite of programs.\(^{30}\) Three functionals (B3LYP,\(^{31,32}\) PBE\(^{33}\) and TPSS\(^{34}\)) combined with Grimme’s-D3 dispersion correction with the Becke and Johnson damping (BJ-damping) function (hereon referred to as B3LYP-D3BJ, PBE-D3BJ and TPSS-D3BJ) were used in conjunction with the def2-TZVPP basis set to investigate the stability of several zinc chloride species.\(^{35}\) The difference in results between the three tested functionals were within the error limit for DFT (~10 kJ mol\(^{-1}\)). Consequently, further calculations of transition states and those including continuum solvent environments were carried out using B3LYP-D3BJ/def2-TZVPP only.

The CPCM (conductor-like polarisable continuum model) was used to provide a generalised solvation environment.\(^{36,37}\) In this work we employed DCM and 1-hexanol to provide a continuum solvent environment with dielectric constants \((\epsilon)\) of 8.93 and 12.51 respectively. Recently Rowe et al. have employed the SMD-GIL solvent model to describe the solvation environment of bismuth halometallates.\(^{38}\) Structures were fully optimised under no symmetry constraints and confirmed as minima or transition states (a single imaginary frequency) using vibrational analysis. Optimisation convergence criteria were set to \(10^{-9}\) on the density matrix and \(10^{-7}\) on the energy matrix, and the numerical grid was improved from the default to a pruned (optimised) grid of 99 radial shells and 590 angular points per shell. Vibrational frequencies and zero-point vibrational energy corrections (ZPE) were attained using the harmonic approximation.

### 2.4 Life cycle analysis

In order to estimate the cost and environmental impacts of ionic liquids from a life cycle perspective, synthesis throughout the entire supply chain was considered. This included extraction of raw materials, manufacturing of intermediate chemicals, and production \(\text{(i.e., cradle-to-gate)}\). For most chemicals, data was obtained from the ecoinvent database;\(^{39}\) however, for ionic liquids and some precursors, data was unavailable. For these processes, Aspen HYSYS V11 was used to model scaled-up production processes. Physical properties were obtained from published sources or estimated using semi-empirical models or the property constant estimation system (PCES) built into Aspen-HYSYS v11. Full details are provided in the ESL.\(^{†}\)

The total annualized cost (TAC), which estimates the cost of the product over the lifetime of the plant, is used following the approach of Towler and Sinnott.\(^{40}\) In short, it consists of the annualized capital expenses (CAPEX) and the annual operating expenses (OPEX), with a 10-year plant life operating 330 days \((7920 \text{ hours})\) a year. The environmental impacts are quantified using the life cycle assessment (LCA) approach following the ISO 14040 principles and phases: (i) goal and scope, (ii) inventory analysis, (iii) impact assessment and (iv) interpretation. Here, SimaPro was used as the LCA modeling tool, and the functional unit was 1 kg of ionic liquid—production was located in Europe. The life cycle inventory (LCI), which consists of the process inputs and outputs, was obtained from both ecoinvent when available or process simulation as explained earlier for missing data. Finally, in the impact...
assessment phase, these inventories were translated into environmental impacts using ReCiPe 2016 as the characterization method, which converts data to 17 environmental indicators (midpoints) such as global warming potential (GWP) and ozone depletion, and finally aggregates into three damage areas (endpoints): human health, ecosystem quality and resources.

3. Results and discussion

3.1 Thermal stability

We first decided to investigate the halozincate series of ionic liquids to explore how changing the amount of metal affected thermal properties. This was particularly important given the previously identified change in Lewis acidity across the series, which could significantly affect how cations and anions interacted with each other and decomposition products. TGA ramping experiments in an N₂ atmosphere showed that larger quantities of ZnCl₂ caused a considerable increase in thermal stability for the [C₈C₇Im]ClₓZnCl₂ ionic liquids (Fig. 2a). For example, T₁% and T_onset values increased by 159.5 °C and 170.4 °C respectively from xZnCl₂ = 0 to 0.67. Thermal stability was effectively found to obey a sigmoidal relationship between metal free [C₈C₇Im]Cl and pure ZnCl₂ (Fig. 2b). This observation was also supported by bromide analogues, but the increase in T₁% from xZnBr₂ = 0 to 0.67 was smaller at 123.8 °C. For [C₈C₇Im]ClₓZnCl₂, the TGA thermogram showed an initial profile that matched metal-free [C₈C₇Im]Cl, while a small portion of the sample (<30%) mirrored the mass losses observed for [C₈C₇Im]ClₓZnCl₂. The composition of the [C₈C₇Im]ClₓZnCl₂ ionic liquid is known to be predominantly [C₈C₇Im]Cl with smaller amounts of [C₈C₇Im]₂[ZnCl₄]₄. Consequently, thermal parameters have captured the decomposition of the less thermally stable organic component. However, decomposition of the organic cation may affect speciation of the residual [ZnCl₄]²⁻ anion—this will be discussed later in this work—so we have reported these values as an upper limit to prevent significant decomposition of the liquid phase and poisoning of the metal complex.

Halide containing ionic liquids have been demonstrated to decompose via reverse Menshutkin chemistry,⁴² where the

![Fig. 2 TGA data and derived parameters of halometallate ionic liquids in N₂, showing: (a) 10 °C min⁻¹ ramping experiments for [C₈C₇Im]ClₓZnCl₂; (b) plot of T₁% as a function of ZnCl₂ mole fraction for [C₈C₇Im]ClₓZnCl₂ with bromide analogues; (c) 10 °C min⁻¹ ramping experiments for [Cation] X₀.₃₃ZnX₂ with metal-free analogues; (d) 10 °C min⁻¹ ramping experiments for [C₈C₇Im]ClₓMCl₂ for four tetrahedral metals; (e) long term thermal stability parameter T₀.₀₁/¹₀ as a function of mole fraction for [C₈C₇Im]ClₓZnCl₂; (f) Ea of thermal decomposition for [C₈C₇Im]ClₓZnCl₂ from isothermal and MTGA.](image)
halide anion acts as a nucleophile to produce neutral chloroalkanes and alkylimidazolides.\textsuperscript{43} For [C\textsubscript{6}C\textsubscript{5}Im\textsubscript{3}][ZnX\textsubscript{X}'], thermal stability significantly increased after all free halide anions had been bonded by ZnX\textsubscript{2} in the [ZnX\textsubscript{X}']\textsuperscript{2−} complex. This suggested that the reverse Menshutkin decomposition pathway is either suppressed or regulated by the breaking of Zn–X bonds. The identity of the halide anion had very little effect on stability of [ZnX\textsubscript{X}']\textsuperscript{2−}imidazolium ionic liquids because T\textsubscript{1%} showed a 0.1 °C difference for CI\textsuperscript{−} vs. Br\textsuperscript{−} (Table 1). However, analysis of the phosphonium analogue [P\textsubscript{66614}]Cl\textsubscript{0.33}ZnCl\textsubscript{2}, which also contains the anionic complex [ZnCl\textsubscript{4}]\textsuperscript{2−}, showed that the stability of [ZnCl\textsubscript{4}]\textsuperscript{2−} could exceed the upper limit of the imidazolium analogues. Unfortunately, a large deviation between T\textsubscript{1%} and T\textsubscript{onset} highlighted that small amounts of weight loss were occurring at lower temperatures for [P\textsubscript{66614}]Cl\textsubscript{0.33}ZnCl\textsubscript{2}, which was likely caused by low level haloalkane impurities from the commercially sourced ionic liquid.\textsuperscript{44} Regardless, bulk decomposition did occur at higher temperatures (T\textsubscript{onset} was 71.6 °C higher than [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}ZnCl\textsubscript{2}], which indicated that steric hindrance was reducing the rate of the reverse Menshutkin process at lower temperatures (supported by the similarity of the TGA profiles) and therefore raising thermal stability.

Following analysis of the halozincate series, we next looked into the effects of different metal centres on thermal stabilities. From T\textsubscript{1%} values, the order of stability was Ni > Co > Zn > Pt for [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}MCl\textsubscript{2}], which was likely caused by low level haloalkane impurities.\textsuperscript{44} In most cases, the identity of the metal centre had very little effect on the initial decompositions of the samples. For example, there was a small 18.8 °C difference between [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}NiCl\textsubscript{2}] and [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}ZnCl\textsubscript{2}]. However, the [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}PtCl\textsubscript{2}] sample was found to undergo a series of decomposition steps from 171.7 °C, which was ≈90 °C lower than that of the [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}NiCl\textsubscript{2}] ionic liquid. Unlike all other metals, the [PtCl\textsubscript{4}]\textsuperscript{2−} complex is square planar and capable of forming large cluster [Pt\textsubscript{6}Cl\textsubscript{m}\textsuperscript{−} anions under certain conditions.\textsuperscript{46,47} Thermal decomposition of [NH\textsubscript{4}][PtCl\textsubscript{6}] has previously been reported and the authors have suggested that decomposition proceeds through Cl\textsuperscript{−} loss to give [PtCl\textsubscript{4}]\textsuperscript{2−} which then decomposes to volatile products and involatile Pt nanoparticles.\textsuperscript{48} Similar decomposition mechanisms have been reported for [NH\textsubscript{4}][Cl\textsubscript{2}][ZnCl\textsubscript{2}], which passes through the [NH\textsubscript{4}][ZnCl\textsubscript{2}] intermediate to give volatile ZnCl\textsubscript{2} and [NH\textsubscript{4}][Cl\textsubscript{2}].\textsuperscript{49} One publication has also reported that heating [C\textsubscript{6}C\textsubscript{5}Im][NiCl\textsubscript{4}] to 700 °C for 1 hour (far beyond T\textsubscript{1%} or T\textsubscript{onset}) gives mixtures of NiCl\textsubscript{2} and metallic Ni.\textsuperscript{50} Altogether, these reports suggest that M–Cl dissociation is likely to be the initial decomposition step of halometallate anions, with the potential to form larger bridged clusters as Cl\textsuperscript{−} is lost. Weakening of the Fe–Cl bonds as a function of temperature has even been observed experimentally during in situ XAFS measurements of [FeCl\textsubscript{4}]\textsuperscript{−} ionic liquids.\textsuperscript{51} However, very little mechanistic insight has been presented to date. Interestingly, the difference between T\textsubscript{1%} values for [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}ZnCl\textsubscript{2}] and [P\textsubscript{66614}][Cl\textsubscript{0.33}ZnCl\textsubscript{2}] observed in this work indicated the decomposition mechanism is caton dependant and thus more complex.

Long term thermal stability was evaluated using the T\textsubscript{0.01/10} parameter (the temperature at which 1% mass loss occurs over 10 hours) to provide more accurate upper temperature limits over extended periods (Fig. 2e and Table 1). As expected, T\textsubscript{0.01/10} values were consistent for metal free halide ionic liquids such as [C\textsubscript{6}C\textsubscript{5}Im][Cl] and [C\textsubscript{6}C\textsubscript{5}Im][Br], deviating by only 2.2 °C. For the [C\textsubscript{6}C\textsubscript{5}Im][Cl][ZnCl\textsubscript{2}] series, the presence of nucleophilic Cl\textsuperscript{−} anions limited stability as observed with T\textsubscript{1%} data, but once all free chloride was sequestered in the [ZnCl\textsubscript{4}]\textsuperscript{2−} coordination complex, T\textsubscript{0.01/10} values significantly increased and reached a maximum at 337.5 °C for [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.67}ZnCl\textsubscript{2}]. This value is approximately 138 °C higher than ionic liquids with [NT\textsubscript{f\textsubscript{2}}]\textsuperscript{−} anions\textsuperscript{26} and marginally exceeds the values of thermally robust dicaticionic ionic liquids.\textsuperscript{27} Furthermore, the 158.3 °C increase in T\textsubscript{0.01/10} from metal free [C\textsubscript{6}C\textsubscript{5}Im][Cl] to metal-rich [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.67}ZnCl\textsubscript{2}] covers a wide temperature range than spans all other published data. E\textsubscript{g} values were within the expected ranges for ionic liquids\textsuperscript{28} and both stepwise and MTGA derived values were reasonably similar for both techniques, but values diverged for γ = 0.33. This was most likely due to a change in reaction mechanism which invalidated one of the assumptions of a zero order (stepwise) or first order (MTGA) mechanism.

Alongside high temperature thermal analysis, we have also reported low temperature thermal transition and heat capacities measured using DSC (Table 1) Previous publications have reported some DSC data for halometallate ionic liquids, but large swings in melting points the presence of multiple glass transitions suggests the data was influenced by experimental factors.\textsuperscript{10,52} We have therefore aimed to provide high quality and reliable DSC data by avoiding the use of a Glovebox, which is subject to pressure swings and therefore difficult to get accurate masses for heat capacity measurements. Instead, we have prepared samples in air using an analytical balance with pinhole pans, followed by drying procedures (see Experimental) to remove residual water from momentary exposure to air. Organic and aqueous solvents have large impacts on heat capacities,\textsuperscript{29} so thorough drying is a necessity for accurate data. All ionic liquids (except [C\textsubscript{6}C\textsubscript{5}Im][Cl\textsubscript{0.33}PtCl\textsubscript{2}]) were found to have low temperature glass transitions, and small variations in T\textsubscript{g} values were noted for changes in composition. Using the measured T\textsubscript{g} values from DSC and T\textsubscript{1%} values from TGA, we have calculated and reported liquid ranges for most ionic liquids. The measured C\textsubscript{p} values of metal free ionic liquids were in the expected ranges.\textsuperscript{24} However, metal ionic liquids had significantly higher C\textsubscript{p} values that generally increased with higher mole fractions as molecular weight increased. This highlighted that the energy requirements for heating bulk ionic liquids would be significantly higher if the metal ionic liquids were used—further justification for the drive towards thin films and solid supports.

3.2 Decomposition mechanism

After our initial TGA experiments, we were motivated to further investigate decomposition mechanisms to understand how ionic liquid structures decompose in the presence of...
| Cation | Anion | Metal | $\chi$ | MW | $T_m$ | $T_g$ | $T_{1%}$ | $T_{onset}$ | Liquid range | MTGA | Flynn–Wall | Isothermal | Stepwise | $T_{0.01/3}^{oc}$ | $C_p/J \cdot mol^{-1} \cdot ^{oc}^{-1}$ |
|--------|-------|-------|-------|----|-------|-------|----------|----------|-------------|------|------------|------------|----------|---------------|------------------|
| [C$_6$C$_{1}$Im]$^+$ | Cl$^-$ | ZnCl$_2$ | 0 | 230.8 | −59.91 | 194.5 | 238.6 | 254.4 | 135.6 ± 1.6 | 140.8 | 129.9$^{d}$ | 133.1 ± 5.7 | 179.2 | 399.3 ± 0.4 | 64.2 ± 0.4 |
| 0.1 | −49.74 | 201.6 | 232.1 | 251.3 | 120.4 ± 0.9 | 148.4 ± 1.1 | 177.1 |
| 0.33 | −43.55 | 243.3 | 286.8 | 286.9 | 161.1 ± 7.5 | 170.4 | 161.7 ± 4.9 | 252.1 | 780.6 ± 0.5 | 117.9 ± 0.5 |
| 0.4 | −43.56 | 286.2 | 309.8 | 329.7 | |
| 0.43 | −46.45 | 282.5 | 305.3 | 329.0 | |
| 0.5 | 734.1 | −47.35 | 327.0 | 373.8 | 374.3 | 153.2 ± 0.8 | 154.5/46.1 | 301.9 | 115.6 ± 0.7 | 167.3 ± 0.8 |
| 0.6 | 870.4 | −52.81 | 348.9 | 392.8 | 401.7 | 144.7 ± 0.4 | 155.3 ± 4.2 | 323.6 | 1038.6 ± 0.9 | 118.9 ± 1.1 |
| 0.67 | 1006.7 | −38.39 | 354.0 | 409.0 | 392.4 | 129.3 ± 0.8 | 164.2 | 158.2 ± 4.1 | 337.5 | 1337.0 ± 0.8 | 154.3 ± 1.4 |
| 1 | 136.3 | | 369.8 | 473.5 | |
| [C$_6$C$_1$Im]$^+$ | Br$^-$ | ZnBr$_2$ | 0 | 275.2 | −64.98 | 208.2 | 277.4 | 273.1 | 121.8 ± 0.5 | 129.3 ± 3.2 | 181.4 | 411.2 ± 0.2 | 57.7 ± 0.2 |
| 0.1 | −49.68 | 206.8 | 247.6 | 256.5 | 124.7 ± 0.6 | |
| 0.33 | −44.21 | 243.4 | 300.6 | 287.6 | 152.0 ± 1.0 | |
| 0.5 | 1000.8 | −54.73 | 304.9 | 380.5 | 359.7 | 157.0 ± 0.2 | |
| 0.6 | 1226.0 | −45.37 | 330.7 | 396.2 | 376.1 | 150.6 ± 1.3 | |
| 0.67 | 1451.2 | −35.96 | 332.0 | 401.4 | 368.0 | 140.5 ± 0.8 | 151.0 ± 1.2 | 331.3 | 1254.9 ± 0.7 | 894.1 ± 1.3 |
| [P$_{66614}$$^+$] | Cl$^-$ | ZnCl$_2$ | 0 | 519.3 | −68.96 | 216.1 | 350.3 | 285.0 | 155.5 ± 2.1 | 118.8 | 114.0 | 1117.3 ± 0.7 | 184.3 ± 1.7 |
| 0.33 | 1174.9 | < −90 | 225.8 | 358.4 | < 35.58 | 176.1 ± 0.8 | 185.6 | 117.4 ± 2.5/32.2 ± 4 | 314.0 | 2337.7 ± 1.3 | 496.2 ± 2.2 |
| [C$_6$C$_1$Im]$^+$ | Cl$^-$ | NiCl$_2$ | 0.33 | 591.2 | −43.75 | 262.0 | 313.5 | 305.8 | 146.0 ± 0.6 | 143.8 | 244.7 | 866.3 ± 0.4 | 109.4 ± 0.5 |
| 0.33 | 591.4 | −42.21 | 253.6 | 295.4 | 295.8 | 150.9 ± 0.6 | |
| 0.5 | 451.9 | −47.6 | 351.7 | 422.1 | 399.3 | 138.6 ± 0.3 | |
| PtCl$_2$ | 0.33 | 727.5 | 73.00$^b$ | −26.28$^b$ | 171.7 | 179.5 | 98.6 | 143.8 ± 0.5 | |
| [C$_6$C$_1$Im]$^+$ | Cl$^-$ | AgCl | 0.33 | 604.9 | 211.0 | 236.3 | 213.3 | 133.7 ± 2.7 | 133.6 ± 2.6 | 197.2 |
| 0.33 | 738.2 | 232.6 | 265.9 | 127.9 ± 5.0 | |
| [P$_{66614}$$^+$] | Br$^-$ | AgBr | 0.33 | 1174.9 | −71.54 | 208.1 | 329.7 | 279.6 | 151.5 ± 0.5 | |
| [C$_4$C$_1$Im]$^+$ | Cl$^-$ | ZnCl$_2$ | 0 | 147.4 | −62.73$^d$ | 202.0 | 245.2 | 264.7 | 152.5 ± 1.5 | 132.2$^d$ | 133.5 ± 1.3 | 180.7 | 732.3 ± 0.3 | 79.6 ± 0.6 |
| 0.5 | 621.9 | −49.23 | 320.2 | 370.8 | 369.4 | 143.4 ± 1.1 | |
| [C$_3$C$_1$Im]$^+$ | Cl$^-$ | ZnCl$_2$ | 0.33 | 146.6 | 89.00$^f$ | 429.5 | 97.47$^f$ | 565.8 | 45.15$^f$ | |
| 0.33 | 295.9 | 97.47$^f$ | |
| [C$_8$C$_1$Im]$^+$ | Br$^-$ | ZnCl$_2$ | 0.33 | 686.7 | −43.91 | 247.4 | 279.5 | 291.3 | 147.8 ± 0.6 | 150.6 ± 0.6 | 135.5 ± 0.6 | 896.7 ± 0.6 | 135.5 ± 0.6 |
| 0.5 | 823.0 | −44.68 | 293.7 | 363.5 | 358.4 | 145.1 ± 0.5 | 282.2 | 1025.0 ± 0.6 | 128.2 ± 0.7 |
| [C$_6$C$_1$Im]$^+$ | Cl$^-$ | ZnBr$_2$ | 0.5 | 911.9 | −52.64 | 312.8 | 382.6 | 365.5 | 155.1 ± 0.3 | 142.5 ± 4.1/100.5 ± 1.3 | 295.3 | 1020.4 ± 0.2 | 132.7 ± 0.5 |

$^a$From ref. 26. $^b$From ref. 10. $^c$From ref. 45.
metals. Insights such as this can help in the design of more robust solvents and catalysts or help to identify harmful decomposition products to improve safety on large scales.

Firstly, we chose to analyse decomposition vapours of the $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ $\chi$ $\text{ZnCl}_2$ series using hyphenated TGA-MS (Fig. 3a–c). Importantly, TGA-MS favours volatile species because low volatility decomposition products can condense in transfer lines or exhaust ports. In this work, we observed mainly hydrocarbon fragments and the reverse Menshutkin product $\text{MeCl}$, as expected from previous reports on metal free ionic liquids.\textsuperscript{27,55} Interestingly, $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$, $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{ZnCl}_2$, and $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.67}\text{ZnCl}_2$ were all found to release $\text{MeCl}$ on heating. This supported the hypothesis that free chloride was produced as a decomposition product in metal containing $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ $\chi$ $\text{ZnCl}_2$ ionic liquids when they were heated to high temperatures. Liberated Cl$^-$ is expected to immediately participate in the reverse Menshutkin decomposition mechanism at high temperatures, releasing chloroalkanes and alkylimidazoles. Alkylimidazole by-products have significantly lower volatility than chloroalkanes, hence why they are not observed in TGA-MS but are observed in vacuum based thermal decomposition studies.\textsuperscript{56} MeCl was also observed during decomposition of $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{NiCl}_2$ (ESI, Fig. S133a\textsuperscript{†}), which highlighted that the same process was occurring for different metals. Unlike the imidazolium ionic liquids, $[\text{P}_{66614}]\text{Cl}_{0.33}\text{ZnCl}_2$ did not produce MeCl because it does not contain a methyl group (ESI, Fig. S133c\textsuperscript{†}). Larger chloroalkanes were not observed, most likely because of low volatility and/or steric hinderance around the phosphonium centre.

STA data for $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ showed a sharp endothermic signal during mass loss (Fig. 3d), which indicated that a single decomposition process dominated. For $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{ZnCl}_2$, the two-step decomposition gave two endothermic signals—the first signal matched that of $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ while the second signal matched that of $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.67}\text{ZnCl}_2$. Broadening of the first signal indicated that the decomposition process was not as well defined as $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$, because Zn–Cl bond dissociation was restricting the decomposition process. Interestingly, this data suggested that decomposition did proceed through reverse Menshutkin chemistry, and as volatile products vaporised, the remaining ZnCl$_2$ concentrated in the ionic liquid so that subsequent decompositions matched that of high ZnCl$_2$ content ionic liquids. This process would ultimately give ZnCl$_2$ which has a higher $T_{1\%}$ than any of the ionic liquids measured in this work. $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{ZnCl}_2$ also produced a similar heat flow profiles to $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{ZnCl}_2$ on thermal decomposition, but $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{AgCl}$ and $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}_{0.33}\text{NiCl}_2$ gave less well defined profiles suggesting competing processes could be occurring during decomposition (ESI, Fig. S134†).

Fig. 3 (a–c) TGA-MS data for $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ $\chi$ $\text{ZnCl}_2$ where $\chi$ = 0 (a), 0.33 (b), and 0.67 (c). (d) STA data for the same $[\text{C}_8\text{C}_1\text{Im}]\text{Cl}$ $\chi$ $\text{ZnCl}_2$ ionic liquids. (e) Example TGA abort thermograms for ex situ analysis. (f) TGA data for selected ionic liquids spiked with 15% $\text{C}_8\text{Im}$ (coloured, solid) and the pure ionic liquids without $\text{C}_8\text{Im}$ for comparison (grey, dashed).
Following hybridised and hyphenated TGA techniques, we
decided to analyse decomposition residues to broaden our
analytical tool belt and shed insight on the involatile products
that do not immediately contribute to mass loss. We pro-
grammed an abort method to stop TGA experiments at pre-
determined mass losses, and then analysed the residues
ex situ with 1H NMR spectroscopy (ESI, Fig. S63–69†) and +ve
mode ESI-MS (Table 2 and ESI, Fig. S62†). These methods
probe the organic cationic components; difficulties in analys-
ing chlorozincate anions with techniques such as Raman
(vibrational frequencies can be difficult to assign) and −ve
mode MS (dianions dissociate to monoanionic species) make
probing the inorganic component challenging. Nevertheless, the 1H NMR and MS data presented in this study
has proven to be very insightful for understanding decom-
position mechanisms.

NMR analysis of the TGA pan residue after 60% thermal
decomposition of [C₈C₇Im]Cl primarily showed the presence
of four compounds (ESI, Fig. S63†) which were identified by
high-resolution mass spectrometry as the intact [C₈C₇Im]⁺
cation (m/z 195), the mixed alkyl chain products [C₁C₇Im]⁺
(m/z 97) and [C₈C₈Im]⁺ (m/z 293), and neutral N-octylimidazole
C₈Im. For the latter, the protonated product was identified in
MS (m/z 181; ESI, Fig. S62†), but the molecule was confirmed
to be neutral by comparing the 1H NMR chemical shifts with
those of synthesised C₈Im (ESI, Fig. S63 and S65†) and HC₈Im
(ESI, Fig. S66†). The intact [C₈C₇Im]⁺ cation was the major
component of the residue (Table 2) with the decomposition
products present in smaller amounts (generally <15%), but the
quantities depended upon the degree of thermal decom-
position. Presence of the mixed chain products suggested that
neutral decomposition products C₈Im and C₈Cl were reacting
in the ionic liquid phase to form new ionic liquid cations—
which is also indirect evidence of the reverse Menshutkin
mechanism. Analysis of decomposed [C₈C₇Im]Cl₀.₃₃ZnCl₂ (ESI,
Fig. S67–69†) and the bromide analogue showed that the
equivalent decomposition products were produced, but larger
quantities of neutral C₈Im were present, along with neutral
C₈Im (m/z 83; protonated salt). The same result was also found
for [C₈C₇Im]Cl₀.₆₃ZnCl₂ but with significantly increased quan-
tities of C₈Im. None of these impurities were observed below the decomposition parameter Tₙₐₓ.

The decomposition product C₈Im is capable of coordi-
ating metals through the nitrogen lone pair. Given the Lewis
acidity of the chlorozincate ions, we next decided to spike
[C₈C₇Im]Cl[ZnCl₂ ionic liquids with 15 wt% C₈Im to probe
potential interactions. Solution phase 1H NMR showed that
chemical shifts of C₈Im were shifted downfield in the presence
of Lewis acidic [C₈C₇Im]Cl[ZnCl₂ (ESI, Fig. S70 and Table S1†),
supporting that coordination was occurring. Variable tempera-
ture NMR also suggested that this coordi-
nation was reversible and C₈Im could dissociate from the
coordination complex (ESI, Table S2†). TGA analysis of the
spiked C₈Im ionic liquid mixtures also proved to be insightful
(Fig. 3f). The neutral C₈Im readily vapourised out of [C₈C₇Im]Cl
but was restricted from vapourising out of [C₈C₇Im]Cl[z₃ZnCl₂;
however, it appeared that most of the C₈Im was lost before
decomposition. Interestingly, [C₈C₇Im]Cl[ZnCl₂ did not
undergo any mass loss until complete thermal decomposi-
tion. This suggested that C₈Im was tightly bound to the Lewis acidic
anion and consequently held in solution well beyond the
boiling point of the neutral alkylimidazole. The C₈Im impurity
was also observed to compromised thermal stability because
decomposition started at a lower temperature than the pure
ionic liquid.

Altogether, ex situ and in situ techniques demonstrated that
imidazolium halometallate complexes were decomposing
through similar mechanisms—chloride dissociation, followed
by the reverse Menshutkin process, with the neutral
N-alkylimidazolium products either vapourising or coordinat-
ing to Lewis acidic metals (Fig. 4). Previous work on
N-methylimidazolide complexes of palladium has shown that
loss of methylimidazole during thermal decomposition pro-
duces three coordinate complexes that dimerise to give

### Table 2 Positive mode ESI-MS data from ex situ analysis of TGA abort experiments

| Cation        | Anion   | Metal   | Temperature/°C | Mass decomposed/% | Peak intensity |
|---------------|---------|---------|----------------|--------------------|---------------|
| [C₈C₇Im]⁺     | Cl⁻     |         | 180            | 0                  | 83            |
|               |         |         | 240            | 20                 | 97            |
|               |         |         | 263            | 60                 | 181           |
| [C₈C₇Im]⁺     | Cl⁻     | ZnCl₂   | 0.33           | 230                | 195           |
|               |         |         | 316            | 20                 | 293           |
|               |         |         | 401            | 60                 | 60            |
| [C₈C₇Im]⁺     | Cl⁻     | ZnCl₂   | 0.6            | 334                | 83            |
|               |         |         | 416            | 20                 | 71            |
|               |         |         | 471            | 60                 | 181           |
| [C₈C₇Im]⁺     | Br⁻     | ZnBr₂   | 0.33           | 396                | 195           |
|               |         |         | 60             | 1.7                | 293           |
| [C₈C₇Im]⁺     | Cl⁻     | NiCl₂   | 0.33           | 375                | 5807          |
|               |         |         | 60             | 2                  |               |
increasingly longer palladium chloride chains. Dimerisation would explain the TGA profiles for the [C₄C₁Im]Cl₂ZnCl₂ series and support the observations of increasingly larger chlorozincate complexes at elevated temperatures. However, we cannot completely eliminate the formation of neutral complexes in competitive decomposition pathways.

To further support our experimentally measured data, we chose to calculate dissociation energies for chlorozincate complexes with DFT. In the gas phase, dissociation energies for loss of a single chloride anion from [Zn₃Cl₇]²⁻ increased with larger amounts of Zn (Tables S3 and Fig. S137†), and thus calculations correlated with experimentally observed thermal stabilities. However, inclusion of a solvent continuum slightly affected this order, most notably for [Zn₃Cl₈]²⁻ which readily dissociated through loss of Cl⁻ or loss of ZnCl₂ to give either [Zn₂Cl₄]⁻ or [Zn₃Cl₅]⁻. While it was not clear why this particular chlorozincate complex was prone to dissociation in our calculations, experimental evidence supports the presence of [Zn₃Cl₈]²⁻ in ionic liquids. However, this discrepancy, along with the existence of other more unusual clusters observed in our calculations (Fig. S138†) in both the gas and solution phase, suggested that exotic structures may be accessed at elevated temperatures. Importantly, in all cases, [Zn₂Cl₆]²⁻ was noted to be the most stable chlorozincate complex, which supported the experimentally measured high thermal stability of the ionic liquid. Interestingly, for [ZnCl₄]²⁻ and [ZnCl₂]²⁻, the monoanionic dissociation products [ZnCl₃]⁻ and [Zn₂Cl₄]⁻ were more stable than the dicatonic species in the gas phase, which supports previously observed mass spectrometry data for chlorozincate anions. However, this trend is reversed with the addition of a continuum solvent environment, with [ZnCl₄]⁻ and [Zn₃Cl₅]⁻ being ~30 and 70 kJ mol⁻¹ more stable than the monoanionic species.

To further probe the effects of different metal centres, activation energies (Eₐ) for thermal decomposition of the tetrahedral complexes (i.e., [ZnCl₄]²⁻, [CoCl₄]²⁻ and [NiCl₄]²⁻) with 1-alkyl-3-methylimidazolium cations were calculated using DFT. This method has previously been shown to provide qualitative estimates of the thermal decomposition temperatures of several ionic liquids. Our results confirmed that the first decomposition step proceeds via a concerted SN2 reaction, similar to that found for [C₄C₁Im]BF₄ ionic liquids. At elevated temperatures, the volatile chloromethane product will rapidly vaporise from the solution, leaving the Lewis basic C₄Im and Lewis acidic MCl₃ which most likely form a Lewis pair. The Eₐ values calculated by DFT follow the experimentally observed TGA data: Ni (149.4 kJ mol⁻¹) > Co (148.8 kJ mol⁻¹) > Zn (147.8 kJ mol⁻¹). Eₐ values were comparable to those experimentally measured by TGA kinetics, but it must be considered that TGA Eₐ values are derived from the rate of mass loss which can capture multiple decomposition processes, rather than a single reaction. Despite this, DFT calculations supported the experimentally observed thermal stability trends and confirmed the hypothesised reaction mechanism.

3.3 Life cycle analysis

To further investigate the economic and environmental impacts of halometallate ionic liquid production, we decided to use a “cradle-to-gate” LCA approach. Production of ionic liquids is known to involve many steps because of their structural complexities, and this was identified as a hotspot early on when compared to molecular solvents. Thus, any benefits gained from ionic liquid use is offset by their production—burden shifting—which emphasizes that synthesis of ionic liquids is particularly problematic when scaling up. For this reason, protic ionic liquids have been identified as more sustainable alternatives because they are produced by acid-base neutralisation, which avoids problematic steps such as alkylation and ion metathesis. Although halometallate ionic liquids still require an alkylation reaction, their synthesis involves direct mixing of two components, as opposed to salt metathesis in solution. Halides are particularly problematic when looking at environmental impacts of ionic liquids. However, with halometallate ionic liquids, halides are an integral part of the end product, and as the ratio of MCl₃ increases so does the halide content. Therefore, LCA analysis of halometallate ionic liquid can provide a valuable insight to production of the most industrially relevant ionic liquids.
Here, we present LCA data for the chlorozincate $[C_8C_1Im]Cl\chi ZnCl_2$ series as a case study to assess how the ratios of components affects environmental and economic impacts (Fig. 5).

The lifecycle tree (Fig. 5a) highlights steps that were estimated with flow sheeting using Aspen HYSYS. Inspection of the tree shows that production of organic $[C_8C_1Im]Cl$ requires significantly more steps than production of inorganic $ZnCl_2$. Increasing quantities of $ZnCl_2$ therefore resulted in a significant decrease in cost and impact categories (Fig. 5b). For example, the cost of $[C_8C_1Im]Cl_{0.67}ZnCl_2$ was $1.11$ per kg lower than $[C_8C_1Im]Cl$, which is a 36% reduction. This is mainly due to the high cost of $[C_8C_1Im]Cl$ ($3.04$ per kg) compared to $ZnCl_2$ ($0.97$ per kg); $[C_8C_1Im]Cl$ is produced from 26 wt% 1-chlorooctane ($2.84$ per kg) and 74 wt% 1-methylimidazole ($1.84$ per kg), but $ZnCl_2$ is produced from 47 wt% $Zn$ ($1.66$ per kg) and 53 wt% $HCl$ ($0.20$ per kg).

The environmental impacts also decreased by similar percentages (EQ: 34%, RD: 41%) with the exception of human health (HH), which was reduced by 19% from that of $[C_8C_1Im]Cl$. This deviation reflects that the precursors for $ZnCl_2$ production are particularly problematic for human health (e.g., $Cl_2$ and $HCl$), but overall, the decrease does support that fewer steps are inherently better. The decrease in resources depletion (RD) is primarily due to $[C_8C_1Im]Cl$; production of 1-chlorooctane and 1-methylimidazole are the main contributors at 68% and 28%, respectively. However, for $ZnCl_2$, the diethyl ether solvent used in production is the main contributor by 71%. Organic solvents can be replaced by better alternatives but starting materials cannot, so process development could further decrease the impact of high $ZnCl_2$ ionic liquids relative to non-metal counterparts. In the human health category, the main damage contributors for $[C_8C_1Im]Cl$ are again 1-chlorooctane and 1-methylimidazole by 69% and 21%, respectively, while $Zn$ and $ZnCl_2$ are the main contributors for $ZnCl_2$ with 40% and 27% contributions, respectively. The contribution of $ZnCl_2$ is mainly due to air emissions of $Zn$ and $CO_2$ from its production. Finally, the impact on ecosystem quality by $[C_8C_1Im]Cl$ is almost double that of $ZnCl_2$. For $[C_8C_1Im]Cl$, 1-chlorooctane and 1-methylimidazole contribute the most by 78% and 14%, respectively, while the main contributors in $ZnCl_2$ are $ZnCl_2$, $Zn$, diethyl ether and $HCl$ (26%, 26%, 23% and 20% respectively). The results above show that only the precursors 1-chlorooctane and 1-methylimidazole are responsible for damages in all impact categories in $[C_8C_1Im]Cl$ while it varies by the damage area in the case of $ZnCl_2$ and it extends beyond precursors to include process emissions and solvents.

4. Conclusions

We have reported a wide range of industrially relevant thermal parameters for 26 halometallate ionic liquids, along with their metal-free analogues. Together, these parameters help define liquid ranges, provide a basis for estimating heating costs, and present upper temperature limits for processes. In terms of thermal stability, the identity of the metal centre had negli-
gible effects when compared to the identity of the ionic liquid cation or the mole fraction of the metal salt. These observations suggested that chloride dissociation was occurring during thermal decomposition, which was immediately followed by reverse Menshutkin chemistry in a concerted mechanism. TGA-MS and STA supported these observations, and ex situ analysis of decomposition residues showed that the neutral alkylimidazole by-products were forming Lewis adducts with halometallate anions. This was particularly problematic for highly Lewis acidic halometallate ionic liquids such as [C₈C₅Im]ClₓZnCl₂ because thermal stability was compromised by the change in speciation. Calculated M–Cl dissociation energies supported the observed stability trend for chlorozincate ionic liquids, but the presence of unusual clusters suggested that the inorganic ions could form more exotic structures at elevated temperatures.

Overall, our mechanistic insight provides a strong basis for designing more thermally robust halometallate ionic liquids to avoiding decomposition which will ultimately poison catalysts. Replacing or topping-up degraded solvents is environmentally damaging, especially when many steps are needed to meet the structural complexity inherent in ionic liquids. However, LCA calculations show that larger mole fractions of metals are better in terms of cost and environmental impact. This is largely due to the complexity of the organic cations, which has significantly more steps for production—simplicity is key. Theoretically, small amounts of [C₈C₅Im]Cl could be used to lower melting points of molten salts into lower temperature regimes, bridging the gap between ionic liquids and molten salts. This would simultaneously reduce the economic and environmental impact of the medium, but physical and chemical properties must also be taken into consideration. This highlights that further work is needed to explore structure–property relationships in halometallate ionic liquids if they are to be successfully used on larger scales.

Author contributions
CJC conceived the project and carried out most of the experimental analysis. YC acquired TGA-MS and STA data, RPM carried out DFT calculations, and HB carried out life cycle analysis. The manuscript was written by CJC, RPM and HB and proof-read by all authors.

Conflicts of interest
There are no conflicts to declare.

Acknowledgements
CJC would like to thank Thomas Clayton (University of Nottingham), Istvan Matyi (TA Instruments), and Philip Davies (TA Instruments) for regular help and support with thermal analysis. JPH and CJC would like to thank EPSRC (EP/K005138/1 and EP/P013341/1) for financial support during the early stages of the project. PL and CJC would also like to thank EPSRC (EP/K005138/1 and EP/P013341/1) for support during the later stages of the project. KRJL acknowledges a Royal Society University Research Fellowship (URF/FR150353) for financial support, and RPM acknowledges support from a Daphne Jackson Fellowship—jointly funded by the Royal Society of Chemistry and the Royal Academy of Engineering. RPM also thanks the Imperial College Research Computing Service (https://doi.org/10.14469/hpc/2232) for computational resources.

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