FEATURE ARTICLE
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Ionic liquids and deep eutectics as a transformative platform for the synthesis of nanomaterials
Ionic liquids and deep eutectics as a transformative platform for the synthesis of nanomaterials†

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Ionic liquids (ILs) are becoming a revolutionary synthesis medium for inorganic nanomaterials, permitting more efficient, safer and environmentally benign preparation of high quality products. A smart combination of ILs and unconventional synthesis methods allows added value to be drawn from the broad matrix of available property combinations. Mixed systems such as Deep Eutectic Solvents (DES) offer a similarly broad combinatorial playground, which is also beginning to translate into applications. Approached holistically, these liquids therefore enable new universal manufacturing techniques that provide solutions to the existing problems of nanomanufacturing, and beyond that will open completely new horizons and possibilities for controlling the growth and assembly of nanostructures. Examples that illustrate the power of ILs in the improved manufacturing of nanomaterials are explored, such as the synthesis of light phosphors with exceptional quantum yields, record-figure-of-merit thermoelectrics, and efficient photocatalysts, alongside developments in DES nanostructure and deep eutectic-solvothermal and ionothermal techniques.

Introduction

Manufacturing of materials on the nanoscale has moved into the focus of chemists, materials scientists and engineers in the last few decades. The chemical and physical properties of materials can change significantly when brought to the nanoscale. These size-dependent properties make nanostructured materials extraordinarily valuable functional materials in many applications ranging from sunscreen, cosmetics, food products, packaging, clothing, disinfectants, household appliances, energy conversion and storage, and construction materials.¹ With growing societal demand for a high standard of living, products and devices that rely on nanomaterials are becoming increasingly important. Consequently, the development of economically-viable and value-added sustainable processes for manufacturing nanomaterials, which are less intensive in terms of energy and the environment, is becoming an important task. Therefore, ILs are a transformative tool because of their unique properties and property combinations. ILs are frequently discussed as “green” alternatives in organic synthesis and catalysis.²⁻⁴ ILs are not “green” per se, but they can be a vehicle to design processes in compliance with the 12 principles of Green Chemistry⁵ and Engineering.⁵ Let us scrutinize the benefits of task-specific ILs in this context, such as the possibility to tune phase, size, morphology, and nanostructure without using additional chemical additives. These become particularly potent as green synthesis tools once combined with the advantages of unconventional synthesis techniques, such as microwave, ultrasound, physical deposition and sputtering techniques (vide infra). Therefore, unprecedented green and efficient nanomanufacturing processes leading to superior materials can be realised by approaching problems holistically with ILs.

ILs are generally defined as salts with melting points below 100 °C whose melts are composed of discrete ions. ILs are not necessarily new materials; examples have been known for over 100 years,⁶ but attention was renewed over the last 20 years. Research interest intensified particularly from 1995 onwards, first due to efforts by the United States Air Force in applying ILs as electrolytes, and subsequently from suggestions that ILs have special properties as replacements for conventional solvents.⁷ However, it is now being realised that it is rather the unique combination of properties, unattainable by molecular compounds or crystalline salts alone, that render ILs exceptional materials and brings them under more intense scrutiny.⁸ The non-volatility, high thermal stability, and wide liquid windows achievable by many ILs has proven an important driver for research⁹ and a technology enabler, supporting numerous advances beyond the initial investigations of ILs as liquid electrolytes.¹⁰ To name but a few applications, ILs have been used as replacement solvents for thermal fluids,³ lubricants,¹¹ processing of cellulose,¹² biphasic chemical processes (e.g., BASF’s BASIL® process; BASF was the...
first to show eco-efficient chemical processes by implementing ILs, photovoltaics, fuel cell electrolytes, and in separation science, chemical synthesis and catalysis, and as non-volatile highly energy-dense materials (aka. “energetic liquids”).

While they will be introduced comprehensively in the latter half of this review, we will briefly touch upon Deep Eutectic Solvents (DES) here. DES are a relatively recent class of systems, which are frequently regarded to be in the same category as ILs as solvents. They are eutectic mixtures of two or more compounds, where the melting point is particularly depressed, making the liquid state accessible. Usually, DES contain salts such as choline chloride mixed with an organic hydrogen-bonding species, giving them an ‘IL-like’ character. DES have high ionic strength (albeit lower than a pure salt due to the molecular component), and potentially similar physical and chemical properties depending on the mixture. It is worth noting here that DES are not always ionic-molecular mixtures, just that the most popular systems are. Given the significant overlap, it is therefore relevant to address ILs alongside DES in the same review. Since there are differences between review coverage for ILs (more recent and extensive) and DES (significantly uncovered) for nanomaterial synthesis, the two parts of this review will be addressed in different styles; firstly, ILs will be discussed and addressed critically to demonstrate their effectiveness as materials synthesis media. Secondly, recent progress and developments in applications of DES in materials synthesis will be comprehensively discussed to address the current gap in the review knowledge.

Ionic liquids and their general benefits as synthetic media in the manufacturing of inorganic nanomaterials

The high energy density of research within the field of ILs has caused an explosion in the number of new ion classes known to support IL behaviour. These range from the well-recognized systems such as azoliums (e.g., imidazolium, triazolium), phosphoniums, pyridiniums, pyrrolidiniums, alkylammoniums, to

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many new cation classes. Anions include both a wide variety of inorganic anions such as halides, nitrates, perchlorates, sulfates, nitrites, hexafluorophosphates, tetrafluoroborates, azides, and a continuously expanding number of organic anions like triflates, benzoates, sulfacetamides, alkylsulfates, alkylcarbonates, carboxylates, and more. Indeed, an enormous number of potential IL cations and anions can be imagined and their combination gives rise to a massive number of ILs (10^{18}) possible cation–anion combinations for potential ILs have been suggested, each with a unique set of chemical and physical properties. The notion is that desirable properties and unique property combinations can be ‘designed-in’, by choosing the right cation and anion to tune the overall degree of hydrophilicity, solubility, solvent structure through H-bonding, viscosity, and preferential interaction with the formed nanomaterial. ILs are therefore intriguing as nanosynthesis media, because IL-dependent IL-specific properties such as electrostatic interactions, but particularly secondary bonding interactions such as H-bonding, π-interactions, van der Waals forces, and ion size which subtly influence the nanomaterial formation, can be tuned. For the most frequently used, archetypical 1-alkyl-3-methylimidazolium [C₅mim]⁺ cation systems, the different secondary bonding interaction possibilities are shown in Fig. 1.

The imidazolium cation contains an aromatic core which can participate in π-system interactions. This electron-accepting region is likely responsible for the electrostatic attraction with polar moieties on the surface of particles. In contrast, 1-alkyl-2,3-dimethylimidazolium cations, [C₄dimim]⁺ (see bottom left of Fig. 1), also contain an imidazolium core like [C₅mim]⁺. In this case, the acidic 2-H proton is replaced by a methyl group, reducing its H-bonding capability and acidity. Pyrrolidinium cations, [C₅pyr]⁺, do not possess any acidic protons, but retain aromaticity. Pyrrolidinium [pyr]⁺ cations belong to the group of quaternary ammonium cations which possess neither acidic protons nor an aromatic ring system (Fig. 1, bottom right). In addition, the alkyl side chain may be endowed with groups that can preferentially interact with the metal NP, for example thiol groups which stabilise gold particles through chalcophilic interactions.

The Lewis acidity or electron donor power of the IL anion is an important property of the IL anion with respect to the synthesis of nanomaterials. This factor dictates its hydrophilicity, and hence, its capacity to engage in intermolecular H-bonding, especially when considered alongside ion geometry. Fig. 2 illustrates the probable interactions of a prominent IL anion, bis(trifluoromethanesulfonyl)amide (Tf₂N⁻). Here, electrostatic, H-bonding, fluorophilic, and van der Waals interactions are observed. These possibilities are expressed to different extents within the set of known IL anions. Halides are generally strongly coordinating and hydrophilic, whereas hexafluorophosphate and (tris(perfluoroalkyl)-trifluorophosphate) are weakly coordinating and hydrophobic. Anions such as thiocyanate are comparatively weakly coordinating, yet hydrophilic.

These interaction capabilities are not only important for specific interactions during the formation of nanomaterials in ILs, but also for their own self-organisation, which is an important aspect in nanomaterials synthesis. It is now well-established that ILs are highly structured liquids, as comprehensively reviewed by Hayes et al. Akin to surfactant self-assembly processes, the ensemble of interionic interactions between constituents of ILs makes them heterogeneous, with a certain degree of molecular self-organisation. The presence of ionic charges imposes the first structural constraint, which is electroneutrality. In a perfect ionic melt, ions are typically surrounded by a first coordination sphere of 4–6 counterions, and then by successively less ordered concentric layers of alternating charge. These local order effects imposed by Coulomb forces have a longer range than what is generally found in molecular liquids, and cause an unexpected long-range force decay which has been observed in interfacial (AFM (atomic force microscopy) and SFA (surface force apparatus)) measurements. The second structural constraint is imposed by the presence of non-polar moieties such as alkyl chains, which can reach significant lengths. The resulting self-assembly therefore depends on the relative contribution of Coulombic, hydrophilic and solvophobic interactions. For example, imidazolium ILs bearing longer alkyl chains display more prominent domain separation (nanostructure) due to a greater contribution from van der Waals interactions in the apolar regions, which increases with the chain length, and alters the balance of...
interactions with the charged, polar parts.\textsuperscript{37} ILs are also capable of having little to no nanostructure, if there is no real amphiphilic character, such as comparing ethylammonium nitrate (extended network) to ethanolammonium nitrate (small ion clusters).\textsuperscript{38} Consequently, as reaction media, a more strongly nanostructured IL may provide an additional reaction control route, \textit{via} the ordered domains which exist on large length- and timescales, in which reactions could occur (\textit{vide infra}).

For alkyl imidazolium ILs, molecular dynamics (MD) simulations\textsuperscript{39–41} and coarse-grained modelling\textsuperscript{42} suggested structural segregation into hydrophilic (charged) and hydrophobic (non-charged) domains: They are composed of rigid ionic channels, built from the H-bonded network of anions and cation head-groups, and segregated lipophilic domains formed by the alkyl tails of the cation, as shown in Fig. 4.

This structural picture has been experimentally validated by X-ray\textsuperscript{44,45} and neutron diffraction studies.\textsuperscript{46–48} Similar studies have been carried out for tetra-alkylammonium- and tetraalkylphosphonium-based ILs,\textsuperscript{49} and a set of pyrrolidinium ILs.\textsuperscript{50} 2D NMR (nuclear magnetic resonance) techniques have also been employed for structure analysis.\textsuperscript{51} This segregation of hydrophilic (charged) and hydrophobic (uncharged) domains is also observed in the crystalline solid state of many ILs.\textsuperscript{52–56} These supramolecular interactions are the origin of the thermotropic mesophase behaviour of many ILs, leading to the formation of liquid crystalline phases.\textsuperscript{57,58}

Due to their unique structure and structural variety, ILs have proven to be excellent media for the preparation and stabilization of NPs, making features available that conventional reaction media such as classical organic solvents are not able to provide. Firstly, ILs are excellent solvents for a large variety of NP starting materials, ranging from highly ionic inorganic salts to molecular metal–organic precursors, but over that they provide additional benefits:

**High nucleation rates facilitating NP formation**

Although polar, ILs can have low interfacial tension.\textsuperscript{59} Since low interfacial tension values favour high nucleation rates, very small particles can be generated.

**Electrosteric stabilization through the IL preventing particle coalescence and growth**

Once formed, the particles are prevented from undergoing Ostwald ripening due to the formation of an IL solvation corona akin to that seen in aqueous surfactant solutions,\textsuperscript{60} which effectively screens electrostatics, and thus agglomeration of NPs. Many ILs resemble surfactants, and indeed surfactant (surface-active) ionic liquids (SAILs), have become an interesting research area in their own right.\textsuperscript{61–63} Thus, it is no surprise that ILs play the same role as surfactants in stabilising NPs through coordination \textit{via} the cation or anion. Cations and anions with long or bulky alkyl chains can both electrostatically and sterically stabilise nanoparticles in solution, giving either long-term or indefinite colloidal stability.\textsuperscript{64}

** Templating reaction medium enabling NP shape and morphology control**

The impact that the IL can have on the morphology of nanomaterials is illustrated in Fig. 3, which shows SEM images of NiO prepared from an aqueous solution and in 1-butyl-3-methylimidazolium bis(trifluoromethanesulfonyl)amide, $[\text{C}_4\text{mim}][\text{TF}_2\text{N}]$.\textsuperscript{43} Sheet-like morphologies are highly desired for NiO used as an electrode material in supercapacitors. The surface energy has a high dispersion force component enhancing the differences between the interfacial energies of different crystal faces, and thus enforcing morphology. Simply by varying the IL it is therefore possible to obtain ZnO as nano-dots, rods or sheets.\textsuperscript{65}

As stated, ILs are not homogeneous media but present self-organisation on the molecular level. This special quality has been exploited for synthesis of extended ordered nanoscale structures, or for size and morphology control. It has been demonstrated that in the synthesis of metal nanoparticles (NPs) from hydrophobic metal organic precursors, the size of the apolar domains in ionic liquids directly influences the resulting metal particle size.\textsuperscript{66,67} Again, MD simulations provided structural support for this idea, as depicted in Fig. 5.\textsuperscript{68}

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**Fig. 3** NiO prepared in the IL 1-butyl-3-methylimidazolium bis(trifluoromethanesulfonyl)amide (left) and in aqueous solution (right). Reproduced with permission from RSC.\textsuperscript{43}

**Fig. 4** Structural heterogeneity of the ionic liquid 1-hexyl-3-methylimidazolium bis(trifluoromethanesulfonyl)amide ($[\text{C}_7\text{mim}][\text{TF}_2\text{N}]$) illustrated visually from a configuration obtained by molecular simulation. Left: Coloured with respect to the different atom types. Right: Charged atoms (the imidazolium ring of cation and the anion) coloured in red and in green the atoms considered nonpolar (those of the alkyl chain of the cation sufficiently far from the positive head-group). Reproduced with permission from RSC.\textsuperscript{39}
additional capping agents. Due to their subtle interaction with crystal surfaces, the particle size, morphology, and phase can be influenced by the IL. Finally, a well-chosen IL can direct the NP assembly towards hierarchical structures.

Ionic liquids and green nanomanufacturing

ILs are often intrinsically assumed to be “Green Solvents” and are strongly connected with the field of Green Chemistry. These claims hold mainly for ILs which have negligible vapour pressures. Therefore, they are non-flammable, cannot be inhaled, and the solvent cannot pollute the atmosphere when an IL is used for an industrial process. Certainly, these aspects render ILs safer and environmentally more benign than conventional solvents, which are often VOCs (volatile organic compounds). However, it is often neglected that in the synthesis of certain ILs, large amounts of VOCs are used whereas other ILs have manufacturing routes where VOCs can be omitted or largely reduced. Some ILs are highly toxic, yet others are FDA approved.

With approximately $10^{18}$ different ion combinations which each result in an IL with a different set of properties, one must not overgeneralise: ILs are not intrinsically green, but they can be made green. Their use in specific processes may lead to improvements that comply with the 12 principles of Green Chemistry laid out by Anastas and Warner, namely: 1. prevention, 2. atom economy, 3. less hazardous chemical syntheses, 4. designing safer chemicals, 5. safer solvents and auxiliaries, 6. design for energy efficiency, 7. use of renewable feedstocks, 8. reduce derivatives, 9. catalysis, 10. design for degradation, 11. real-time analysis for pollution prevention, and 12. inherently safer chemistry for accident prevention. Green Engineering has been defined by the US Environmental Protection Agency (US EPA) as “The design, commercialization and use of processes and products that are feasible and economical while reducing the generation of pollution at the source and minimizing the risk to human health and the environment.” In this context, ILs act as technology enablers. By looking at the arguments made in the previous section, it becomes evident that employing ILs in the production of nanomaterials renders their manufacturing greener: in NP synthesis the IL acts first as the solvent and reaction medium, then also as an agent for templating, stabilisation, and size, phase and morphology control. This multirerole character means that other additives such as stabilizing agents or surfactants can be omitted in the synthesis. This de-intensifies the process by reducing the number of chemicals needed for the synthesis, especially hazardous additives, with the knock-on effect that a simpler mixture minimises the formation of undesirable by-products and thus waste. Safety is also improved by lower vapour pressures, which additionally facilitate recycling of the IL.

Therefore, fastidious design of nanomaterial manufacturing processes using an IL offers advantages, and can easily satisfy...
half of the principles of Green Chemistry. There is still plenty of room for improvement however, especially in terms of energy and materials (atom) and economic efficiency of the processes. To close this technological gap and fully evaluate the potential of ILs in the context of green, eco-efficient nanomanufacturing, new synthetic procedures have been established that satisfy the principles of Green Chemistry and Engineering. We will now explore combinations of IL technology with unconventional synthetic conversion methods that draw advantages from IL properties.

Conversion methods

Bottom-up, chemical routes to nanomaterials in ionic liquids

In the bottom-up chemical approach for the preparation of inorganic nanomaterials, starting materials such as simple salts, complex compounds, or organometallic precursor molecules are converted within the IL to the desired product (Fig. 7).

For instance, a neutral organometallic precursor would dissolve preferentially within the lipophilic (apolar) domains of the IL. If the experimental conditions impose low levels of mass transport (i.e. low temperature, no stirring), the size of the resulting metal nanoparticles, such as Ru-NPs, is dictated by the number of available zero-valent atoms in the non-polar domain, which is in turn related to its size and hence the length of the alkyl chain. This suggests that the phenomena of nucleation and growth occur in these non-polar IL domains, and are controlled by the local concentration of the metal, as depicted in Fig. 8. Here, the IL functions as both reaction medium and nanoparticle stabilising agent. The preparation of noble metal nanomaterials such as Au, Pd, Pt, Ru, Ag, Ir and Rh have been studied via this route, thanks to their ease of reduction, relative insensitivity to oxygen and high catalytic value. Nonetheless, the protecting nature of the IL means that even more oxophilic metals such as Cu, Ni, W, Cr and Mo can also be produced and stabilized as zero-valent NPs, given that suitable, chemically stable ions are chosen. Analogously to solvothermal synthesis, when ILs are used as the reaction medium in this manner the term ionothermal synthesis has been coined. As most ILs have a negligible vapour pressure, the maximal achievable temperature therefore becomes a function of thermal stability, not reactor pressure. Typical reaction temperatures can reach about 250 °C which, for example, has been used for the conversion of metal carbonyl to NPs.

Microwave synthesis

ILs are excellent media for absorbing microwaves as they are composed of large ions with high polarizability and conductivity, leading to high heating rates which results in a high formation rate of nuclei, favourable for NP formation. High heating rates translate into much shorter heating times and therefore faster conversion than obtainable via conventional heating. This is in addition to the discussed benefits such as the IL acting as an easily-removed NP stabilising agent, thus conferring numerous advantages over conventional solvents. However, in the majority of microwave reactions to produce NPs, so far, ILs have been used only as additives to classical VOCs to enhance the microwave uptake rather than as pure synthesis media. This versatile class of reaction has been termed MAIL (microwave-assisted ionic liquid). Many nanomaterials such as oxides and phosphates, fluorides (hydr)oxyfluorides, and higher chalcogenides such as sulphides, selenides, and tellurides can be produced in this manner. However, adding dilute ILs to a reaction mixture is analogous to adding a simple salt to enhance the microwave uptake; it does not take advantage of the full range of available benefits. When the IL is only the minority component in the reaction mixture, its templating and NP stabilising influence can easily be lost, as the concentration is too low (cf. the critical micelle concentration, or CMC).

Surprisingly, reactions in neat ILs or in systems where the IL is the majority component have scarcely been investigated. To fully utilise the potential of microwave irradiation of ILs to produce NPs, neat ILs (or at least systems where the IL is both the solvent and majority component) have to be employed. There is a broad possibility to optimise several parameters in microwave synthesis such as the irradiation time, reaction temperature (it is even possible to irradiate a sample with microwaves, and simultaneously cryogenically cool with e.g. liquid nitrogen), the heating ramp, the power input rate, the cooling ramp after completion of reaction, size of the reaction
container, quantity of IL, and the precursor concentration and identity. This latter parameter is multifaceted because it not only determines solubility, but also the precursor decomposition pathway and associated kinetics. The choice of IL is crucial because certain systems such as those based on certain imidazolium ILs can yield colloidal stable dispersions.\textsuperscript{111} Similarly, certain cations such as [cholinium]\textsuperscript{+} yield extremely small ZnS nanoparticles, due to strong surface coordination which blocks crystal growth.\textsuperscript{112} This method therefore allows for the fast and facile preparation of metals, metal chalcogenides as well as halide materials, and even convoluted multinary compounds such as ternary and complex oxides.\textsuperscript{98–101}

### Ultrasound assisted synthesis

Sonochemistry, albeit mostly in conventional and volatile solvents, has attracted increasing interest over the past decade for the synthesis of inorganic nanostructured materials.\textsuperscript{113} The effects of ultrasound on chemical conversions originate from acoustic cavitation, which occurs in several stages involving the steps of nucleation, growth, and collapse of bubbles in the liquid. The collapse of the bubbles provokes locally extreme conditions: high temperatures of 5000 K and high pressures of about 500 atm can be reached. Yet, sonochemical synthesis is often considered to be a mild method, because the temperature and pressure in the bulk solvent barely change. As many ILs have an extremely low or negligible vapour pressure they are uniquely suited for sonochemistry, as they do not interfere with the chemistry within the caviton unlike traditional volatile solvents.\textsuperscript{114} In ILs, the situation is best described by a two-site model. Here, the reaction cavity is composed of the bubble’s gas phase interior, which originates from the reactants, and can reach 5000 K, whereas the immediately surrounding shell is composed of the (ionic) liquid, reaching local temperatures of 850–2000 K, and then surrounded further by the bulk.\textsuperscript{113} 3500 K has been reported as the temperature for an impoling cavity in 1-ethyl-3-methylimidazolium ethylsulfate under sonication.\textsuperscript{115} As with the microwave method, the lower heat capacity of most ILs permits faster heating under ultrasound irradiation compared to aqueous solutions.\textsuperscript{116–118} Thus, nanomaterial synthesis time is again reduced by choosing ILs as the reaction medium, and augmented further by faster heating of small volumes, and reduced mass transport (often supported by the relatively high viscosity of ILs), which favours the formation of NPs. Another rarely-mentioned beneficial factor is that ultrasonic cavitation generates high shear which helps to break particle agglomerates into singly dispersed particles. However, a serious consideration is that ILs may decompose significantly under intense sonication.\textsuperscript{114} It is noted that ultrasound horns provide rather harsh conditions, whereas simple ultrasonic cleaning baths (\textit{i.e.} 45 kHz and 60 W) are well suited for experimental nanosynthesis in ILs.\textsuperscript{43,65,73,119–122}

In summary, ultrasound-assisted synthesis in ILs is a simple, convenient, elegant, rapid and cost-effective method to synthesize NPs. It complies with the requirements of green chemistry by reducing energy intensity in production obviating the use of environmentally- as well as economically-hazardous substances. The final product is dictated by a combination of sonication time, power, and frequency, container and sample volume, and the precursor identity.

### Top-down, physical routes to nanomaterials in ionic liquids

Due to the unique properties which can be realized for ILs, namely the vapour pressure and stability under the requisite experimental conditions, top-down chemical routes to nanomaterials are also viable, such as physical vapour deposition, sputtering and laser ablation. Amongst these, laser ablation is the least well-studied, potentially due to laser ablation of a target compound generally leading to relatively large, polydisperse particles.\textsuperscript{123} After initial works,\textsuperscript{124} this synthetic method has largely been neglected, with the exception of a few recent studies.\textsuperscript{125} To date, physical vapour deposition and sputtering are the more investigated methods. Other exciting and versatile top-down synthesis methods also exist which are noteworthy here, especially electrochemical and plasma syntheses,\textsuperscript{126–128} but this review will focus only upon physical and chemical routes, with electrochemistry falling out of scope.

#### Physical vapour deposition

The negligible vapour pressure and flammability of ILs lends them well to handling under high vacuum conditions at elevated temperatures. In this unique approach to obtain nanomaterials, metals, pre-prepared intermetallic compounds and alloys as well as ceramic materials and metal salts are evaporated onto an IL, by employing physical vapour deposition (PVD) methods. This technique is useful even for the preparation of thermodynamically unstable compounds. Our group has pioneered high temperature evaporation for the fabrication of nanoparticles in ILs by modifying the SMAD (solvated metal atom dispersion) technique,\textsuperscript{130} originally developed by Klambek for conventional VOCs.\textsuperscript{131} Substrate vapours and a solvent or stabilising agent are co-condensed under high vacuum onto a target that is maintained at cryogenic (77 K) temperatures (see Fig. 9). In this methodology, instead of using conventional volatile organic solvents and stabilising agents with significant vapour pressures, which require cooling of the substrate to maintain the vacuum during evaporation, ILs offer the possibility to work at ambient temperature with a liquid substrate. With this technique, universal synthetic procedures were developed for metal and metal–metal oxide colloids with long-term stability, as well as fluoride nanophosphors by thermal evaporation or electron-gun evaporation.\textsuperscript{64,130,132}

Such experimental PVD setups are shown in Fig. 9, and consist primarily of a rotating reaction flask in which there is an axially mounted evaporation source, which can be resistive or an electron beam. The system is maintained under high vacuum by a high-speed pump assembly. A quantity of the respective IL is introduced into the rotating reaction vessel, thereby creating a liquid film upon the internal wall of the flask which is constantly regenerated through flask rotation. After the system has been evacuated, a metal, an intermetallic compound or a metal salt is evaporated and condensed into...
the IL film. This gives the advantage that the particles can diffuse immediately upon contact with the surface of the substrate into the bulk. As a consequence, the formation of larger aggregates is suppressed. The rotating flask constantly renews the substrate surface with a fresh film of the IL. The formation of a closed surface can be further disturbed by stirring the IL in the bottom of the flask with a magnetic stirrer. Resistive heating is used for evaporation in the temperature regime of 25–2000 °C whereas electron beam evaporation is best suited for the high temperature region between 1000 and 2500 °C. Evaporation via an electron beam has the advantage of being a containerless method. The electron beam can be focused on the centre portion of the sample, and only the inner portion of the sample is first melted and then evaporated. This method can be used for large scale vaporisation (kg h⁻¹) of metals and their alloys. Fortunately, many physico-chemical and physical data such as evaporation conditions, boiling points under vacuum or gas phase composition are well known from matrix-isolation spectroscopy studies. A quartz crystal microbalance can be used to monitor the progress of evaporation. As the IL is present in excess, formation of larger particles is prevented by blocking growth. The concentration can be varied by changing the evaporation time. For multinary compounds, it is possible to either evaporate from dual sources, or from one source employing a mixture of the starting materials or a pre-prepared compound. Overall, this method has proven versatile and it allows the easy casting of metal, composite, and ceramic materials as NPs, even in large quantities. Of course, the morphology and size of the particles produced can still be controlled by the IL used for synthesis in this method. ⁶⁴

Sputtering

The thermal evaporation rates for each source material are different due to thermodynamic restrictions. This limitation can be circumvented by sputtering. It was observed that the NP size can be tuned via the IL identity, especially by tuning viscosity, as well as the temperature. ¹³³–¹³⁷ The sputtering technique is not restricted to pure metals, and is also applicable to binary alloy systems like Ag–Au. ¹³⁸ Au–Pt alloy NPs were accessible by co-sputtering of Au and Pt into N,N,N-trimethyl-N-propylammonium bis(trifluoromethyl-sulfonyl)amide, even though bulk Au and Pt are immiscible. ¹³⁹ By subsequent oxidation of the metal it is even possible to access small oxide nanoparticles. ¹⁴⁰ This method enables fast establishment of materials libraries with composition variation. By combinatorial sputtering, a large number of NPs with different composition can be manufactured (and tested for a given application). Two or more targets can be used, or alternatively a single target containing a pre-prepared compound. In addition, the sputter power (discharge current) and sputter rate can be tuned as well as the Ar partial pressure and sputter time. For example, Cu and Au were simultaneously sputtered onto a wafer with small cavities containing the IL 1-butyl-3-methylimidazolium bis(trifluoromethane-sulfonyl)amide, yielding alloyed Au–Cu NPs. ¹⁴¹ The sputtering process does not rely on the different reactivity of starting materials, thus providing the benefit of favouring alloy particle production, rather than core–shell particles. This method therefore offers good potential to synthesize multicomponent NPs with the exact size composition required, simply by varying the sputtering conditions and stoichiometry. ¹⁴²–¹⁴⁷ Variation in the products can further be induced by mixing a strongly-stabilizing IL with a poor one, or diluting the IL with, for example, an organic solvent such as anisole. ¹⁴⁸

Bottom-up vs. top-down methods – a comparison

Chemical bottom-up synthesis of NPs involves the reaction (decomposition, thermolysis or metathesis) of simple salts or organometallic precursor molecules, corresponding with each component (ion) that is involved in the final product. When the kinetic reaction rates for the decomposition of different precursors vary for each source compound (which is often the case due to their different reactivities), synthesising multinary compounds becomes challenging. For alloy nanoparticles where one metal is much more noble than the other, like Cu and Au, it is almost impossible to tune the final alloy composition over a wider range of stoichiometric ratios and quite often core–shell materials are obtained. This may be desired or undesired, depending on the application. For catalysts, for example, it could be desirable to have a particle core composed of a cheap and abundant metal and a shell of the catalytically active noble (expensive) material. Here, bottom-up methods have a clear advantage. However, wet chemical approaches are also often conducive to the formation of by-products, which may adhere to the NP surface, influencing the reaction outcome and potentially the materials performance of the product. In that sense, top-down physical methods are cleaner processes which are independent of the different reactivity of the starting materials. Overall, neither synthetic approach can be declared a hands-down ‘winner’; both chemical and physical techniques are pertinent and their relative merit depends on the intended nanomaterial, and its properties and desired application. Some of the respective advantages of the different approaches are shown in Scheme 1. ¹⁴⁹
IL synthesis showcases

A set of examples will now be explored where IL-based nano-manufacturing methods, both bottom-up and top-down, have been used for the synthesis of nanomaterials for energy related applications such as catalysis, energy-efficient lighting and thermolectric power conversion. The examples were chosen primarily to illustrate the universal power of the presented synthetic methodologies, but also to demonstrate that nanomaterial synthesis in ILs can have strong green aspects when designed mindfully. The scrutinised materials are of high relevance for energy applications, and the products from ILs in each case showed an improved performance in comparison to what could be obtained using classical nanosynthesis methods.

Photocatalysts

New tools for clean energy are becoming increasingly necessary to meet the increasing global energy demand, driven by decreasing access barriers to energy-consuming technologies such as computation and transport. While it is not a clean fuel source when produced from hydrocarbons, hydrogen can be used as a green vector for solar energy by using incident radiation to split water.\textsuperscript{150,151} Sunlight could also be used as an energy source for cleaning organic pollutants from air and water. Both of these require viable photocatalytic systems, which are often more powerful when nanostructured. A suitable photocatalyst must have a sufficiently large band gap (\(\geq 1.23\) eV for water splitting) and appropriately-positioned valence and conduction bands, with the band gap efficiently utilising the available solar spectrum.

Suitable semiconducting materials with a viable band gap that can absorb light in the near UV and in the visible region include perovskites such as SrTiO\(_3\) and BaTiO\(_3\), and related materials.\textsuperscript{152} Currently, research to find the optimal photocatalyst material is mainly driven by band gap engineering to match the solar spectrum, with new computational screening approaches beginning to show merit.\textsuperscript{153} However, the choice of synthesis method dictates a number of other important factors essential for a good photocatalyst. Firstly, it is important to maximise the accessible surface area of catalytically active surface sites. However, surface area is not the only important factor. The identity of the crystal facets which are found terminating the nanostructures at the interface also dictate the catalytic performance, alongside the exposed atom identities, the nature and quantity of any defects, the extant electron/hole conduction pathways, and the ease of charge carrier separation and recombination. These factors are all influenced by the materials morphology, size and phase, and hence its nanostructure; it is not unusual for precisely nanostructured materials to vastly exceed the specific catalytic activity of nonspecific materials with high surface area alone. Therefore, it is crucial to develop and understand controlled, robust, and reliable synthetic methodologies to optimise properties on the nanoscale.

A one-step room-temperature ultrasound synthesis for photocatalytically active MTiO\(_3\) (M = Ca, Sr, Ba) nanoparticles in ionic liquids was recently developed in our group.\textsuperscript{121} For this, a solution of M(CH\(_3\)COO)\(_2\)\(\times\)H\(_2\)O, Ti(OiPr)\(_4\) (OiPr = isopropoxide) and finely ground NaOH powder was irradiated with ultrasound in a commercial ultrasound bath (45 kHz, 60 W) in the ionic liquid 1-butyl-3-methylimidazolium bis(trifluoromethane)sulfonyl amide, simply sealed in a glass tube under ambient conditions for 10 h (see Fig. 10).

Five crystalline forms of BaTiO\(_3\) are known, of which three are ferroelectric (rhombohedral, orthorhombic, tetragonal) and stable below 120 °C. The cubic form is stable above 120 °C
transformation to the cubic structure type at 1250 °C from the room temperature stable orthorhombic structure to obtain what would normally be regarded as the ‘high temperature’ state reaction of MCO₃ and TiO₂ at temperatures above 900 °C. H₂ evolution from water were tested using 10% methanol polymorphs at ambient temperatures, with minimal energy usage.

The photocatalytic properties of MTiO₃ (M = Ca, Sr, Ba) for H₂ evolution from water were tested using 10% methanol aqueous solution as the sacrificial agent under UV light, against a reference commercial photocatalyst (P25). Without any co-catalysts the H₂ evolution rates of MTiO₃ (M = Ca, Sr, Ba) were already comparable to those found for P25, demonstrating that active surface sites for hydrogen generation are readily available on the catalyst surface. Photodeposition was used to directly deposit 0.025 wt% of Rh onto reactive surface titanate sites on the catalyst surface. Photodeposition was used to directly deposit 0.025 wt% of Rh onto reactive surface titanate sites where electrons are photogenerated. This caused the H₂ evolution rates of MTiO₃ (M = Ca, Sr, Ba) to improve substantially; in the case of SrTiO₃ the photocatalytic activity exceeded the commercial photocatalyst P25, which stimulated investigation of the influence of the IL used in the catalyst synthesis on the photocatalytic activity. To this end bis(trifluoromethane)sulfonimide ([Tf₂N]⁻)-based ILs with various counter cations including 1-butyl-3-methylimidazolium ([C₄mim]⁺), 1-(3-hydroxypropyl)-3-methylimidazolium ([C₃mimOH]⁺), butylpyridinium ([C₄Py]⁺), and tetradecyltriethyl phosphonium ([P₆₆₆₁₄]⁺) were investigated. Astonishing differences in the photocatalytic activity were observed, ranging from negligible (for SrTiO₃ prepared from [P₆₆₆₁₄][Tf₂N]), to higher than commercial with lower co-catalyst loading (material prepared in [C₄mim][Tf₂N]). Scanning electron microscopy (SEM) showed that the obtained SrTiO₃ had an IL-dependence; morphologies varied from relatively isolated nanospheres in [C₄mimOH][Tf₂N] to aggregates which are cubelike in [C₃mim][Tf₂N], raspberry-like in [C₄Py][Tf₂N], and globular in [P₆₆₆₁₄][Tf₂N]. SrTiO₃ prepared in [C₄mim][Tf₂N] shows the highest photocatalytic activity for H₂ evolution (1115.4 μmol h⁻¹) using 0.025 wt% Rh as the co-catalyst. These materials also performed strongly in the degradation of methylene blue, which was investigated as a model substance for the decomposition of organic pollutants in solution. In this case, it was not the material prepared in [C₄mim][Tf₂N], but the nanospheric product of the [C₄mimOH][Tf₂N] synthesis which showed the highest activity for the photocatalytic degradation of methylene blue (88%) under UV irradiation. High resolution transmission electron microscopy (HRTEM) allowed the correlation of photocatalytic activities with the different dominant surface facets of the SrTiO₃ nanomaterial, which show distinct differences for H₂ production and photocatalytic decomposition of organic materials. This therefore demonstrates how the choice of IL used for the synthesis can be used to control the expression of crystal facets, as shown in Fig. 11.

Light phosphors: phase-selective synthesis of light phosphors with record-high quantum efficiencies

CFLs (compact fluorescent lamps) are widely used as energy-efficient light sources. However, this lighting technology generally requires small amounts of mercury in the light bulbs, representing a lifelong safety hazard. An increasing quantity of mercury is now being recorded in landfills, which is ascribed to

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**Fig. 10** Sonochemical ambient condition preparation of perovskite photocatalysts.

![Sonochemical ambient condition preparation of perovskite photocatalysts.](image)

**Fig. 11** HRTEM images of the as-prepared SrTiO₃ in [C₄mimOH][Tf₂N] with {100} expressed (left) and in [C₄mim][Tf₂N] with expression of {110} (right). Reprinted with permission from Wiley-VCH.

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(its Curie point) up to 1460 °C; beyond this temperature a hexagonal structure is formed. CaTiO₃ faces a transition from the room temperature stable orthorhombic structure to a tetragonal polymorph between 1100–1150 °C followed by a transformation to the cubic structure type at 1250 °C. SrTiO₃ undergoes a cubic to tetragonal transition on cooling at T = 105 K (−168.15 °C). Thus, only for SrTiO₃ is the cubic aristotype the thermodynamically stable form at room temperature. Quite remarkably, under the chosen synthesis conditions the cubic polymorph was obtained for all three, though it was only expected for SrTiO₃. It is suspected that this is due to ultrasonic pressures favour the formation of the high temperature phase; as the pressure and temperature then drop quickly, no phase transformation to the thermodynamically stable polymorph can occur. Typical methods to prepare MTiO₃ rely on solid-state reaction of MCO₃ and TiO₂ at temperatures above 900 °C; quite often the obtained products contain agglomerated particles of different sizes and morphologies as well as impurities owing to incomplete reaction. These example demonstrate how synthesis conditions in IL-based methodologies can be exploited to obtain what would normally be regarded as the ‘high temperature' polymorphs at ambient temperatures, with minimal energy usage. The photocatalytic properties of MTiO₃ (M = Ca, Sr, Ba) for H₂ evolution from water were tested using 10% methanol aqueous solution as the sacrificial agent under UV light, against a reference commercial photocatalyst (P25). Without any co-catalysts the H₂ evolution rates of MTiO₃ (M = Ca, Sr, Ba) were already comparable to those found for P25, demonstrating that active surface sites for hydrogen generation are readily available on the catalyst surface. Photodeposition was used to directly deposit 0.025 wt% of Rh onto reactive surface titanate sites where electrons are photogenerated. This caused the H₂ evolution rates of MTiO₃ (M = Ca, Sr, Ba) to improve substantially; in the case of SrTiO₃ the photocatalytic activity exceeded the commercial photocatalyst P25, which stimulated investigation of the influence of the IL used in the catalyst synthesis on the photocatalytic activity. To this end bis(trifluoromethane)sulfonimide ([Tf₂N]⁻)-based ILs with various counter cations including 1-butyl-3-methylimidazolium ([C₄mim]⁺), 1-(3-hydroxypropyl)-3-methylimidazolium ([C₃mimOH]⁺), butylpyridinium ([C₄Py]⁺), and tetradecyltriethyl phosphonium ([P₆₆₆₁₄]⁺) were investigated. Astonishing differences in the photocatalytic activity were observed, ranging from negligible (for SrTiO₃ prepared from [P₆₆₆₁₄][Tf₂N]), to higher than commercial with lower co-catalyst loading (material prepared in [C₄mim][Tf₂N]). Scanning electron microscopy (SEM) showed that the obtained SrTiO₃ had an IL-dependence; morphologies varied from relatively isolated nanospheres in [C₄mimOH][Tf₂N] to aggregates which are cubelike in [C₃mim][Tf₂N], raspberry-like in [C₄Py][Tf₂N], and globular in [P₆₆₆₁₄][Tf₂N]. SrTiO₃ prepared in [C₄mim][Tf₂N] shows the highest photocatalytic activity for H₂ evolution (1115.4 μmol h⁻¹) using 0.025 wt% Rh as the co-catalyst. These materials also performed strongly in the degradation of methylene blue, which was investigated as a model substance for the decomposition of organic pollutants in solution. In this case, it was not the material prepared in [C₄mim][Tf₂N], but the nanospheric product of the [C₄mimOH][Tf₂N] synthesis which showed the highest activity for the photocatalytic degradation of methylene blue (88%) under UV irradiation. High resolution transmission electron microscopy (HRTEM) allowed the correlation of photocatalytic activities with the different dominant surface facets of the SrTiO₃ nanomaterial, which show distinct differences for H₂ production and photocatalytic decomposition of organic materials. This therefore demonstrates how the choice of IL used for the synthesis can be used to control the expression of crystal facets, as shown in Fig. 11.

**Light phosphors: phase-selective synthesis of light phosphors with record-high quantum efficiencies**

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Improper disposal of CFLs. It would be possible to replace mercury in CFLs with environmentally benign materials such as noble gases, but this requires special nanophosphors with a quantum efficiency higher than 100% (so-called quantum-cutters) to still allow for energy efficiency. Basically, this approach relies on the conversion of electrical energy into UV light and the subsequent transformation of one energy-rich UV photon into several photons in the visible spectrum, as opposed to just one lower-energy photon, which occurs in the commercially available CFLs, where the energy delta is wasted. In the RGB color ensemble, this problem is therefore most pertinent for the generation of long-wavelength (red) light. When doping Eu$^{3+}$ into GdF$_3$ a material can be made that allows for the conversion of one UV photon into two red ones,$^{157}$ but this is not easy to manufacture at the nanoscale due to its special requirements. Firstly, appropriately small and uniform particles are needed to minimize stray-light effects, and the material has to be absolutely oxygen-free for maximum efficiency: if oxygen is present, an efficient radiationless deactivation through a charge transfer process opens up which reduces the quantum yield beyond an acceptable value. Finally, the material has to be obtained in the correct tetrasilicate polymorph, not as orthorhombic YF$_3$. So, in this example there is a clear need to achieve a polymorph-specific synthesis route. Typically, oxide-free fluorides are prepared by treating (Gd, Eu)$_2$O$_3$ (the commercial source) with an excess of HF at high temperatures, which consumes a lot of energy and is patently unsafe to human health. Furthermore, it is expensive and difficult to control, yielding inconsistent products without sufficient quality to be used as a light phosphor (Fig. 12).

IL-based production methods were used to overcome these shortcomings, through microwave heating of Gd- and Eu-acetate in the IL choline tetrafluoroborate for 10 min at 80 °C.$^{158}$ The yielded nanomaterial consisted of small and uniform particles that were absolutely oxygen-free and contained 100% of the desired phase. The choice of IL dictated the resultant fluoride phase, of tysonite or the orthorhombic YF$_3$ polymorph. It must be noted that the desired material is not obtained under conventional heating. However, by conducting material conversion under microwave irradiation in the right IL, materials were obtained with a quantum efficiency close to the theoretical limit of 200%.$^{159}$ The same method was also applied to produce efficient quantum-cutting green phosphors.$^{157}$

In addition to being a fast and energy-efficient low-temperature process, the use of hazardous HF is totally omitted. This synthetic concept can be easily transferred to (white light) emitting phosphors for light emitting diodes (LEDs), as seen in Fig. 12.$^{160,161}$ Even composite materials can be obtained for this purpose.$^{162}$ Phosphates can be made in a similar way to fluorides.$^{163–165}$ If instead of quantum-cutting materials,$^{166,167}$ simple down-shifting$^{167,168}$ or up-converting materials (for example, for bioimaging) are desired, the synthetic protocol does not even require a protective atmosphere.$^{165}$ Still, care has to be taken that the desired polymorph is obtained, which the IL can uniquely help to achieve.$^{168–172}$ We recently also employed the physical vapour deposition approach,$^{132}$ which can also be used for the manufacturing of broad-band emitters based on divergent rare earth cations.$^{173}$

**Record figure-of-merit thermoelectrics**

Solid state energy converters, such as thermoelectrics, are as a main technology to enable waste heat harvesting, e.g. in the cooling system of combustion engines.$^{174}$ At a given temperature $T$, the conversion efficiency of a thermoelectric material is closely related to the dimensionless thermoelectric material's figure of merit $zT$ (eqn (1)):

$$zT = \frac{S^2}{\rho \kappa} T$$

where $S$ is the Seebeck coefficient, $\rho$ is the electrical resistivity and $\kappa$ is the thermal conductivity. Good thermoelectric converter materials have a high Seebeck coefficient, expressing the generated thermovoltage per temperature difference. Moreover, a high electrical conductivity is needed in order to reduce how many Joules of energy are dissipated within the thermoelectric generator. A low thermal conductivity, with contribution from the electronic (el) and phononic (ph) system, is further required so that the thermal shortcut due to Fourier's heat conduction is small. This combination of properties is rather rare. Over decades, the best figure of merits were around $zT = 1$, and typically found in binary and ternary group 15/16 materials containing the heavy elements bismuth and/or tellurium, of the general type: (Sb$_2$Te$_3$, Bi$_2$Te$_3$, Sb$_2$Bi$_2$Te$_3$). Within the last decade, materials concepts like alloying with heavy elements, the implementation of nanostructures and precipitates, the transition to materials with complex elemental cells, or the incorporation of rattling atoms have greatly improved the conversion efficiency of today's thermoelectric generators. Further improvement of the thermoelectric figure of merit $zT$ can be achieved by precise control of the chemical composition combined with nanostructuring of the material.$^{174,176–178}$ While the first is indispensable for the optimisation of the electronic transport properties, the latter reduces the phonon thermal conductivity due to the high number of grain boundaries, which scatter phonons more effectively than electrons.

Most recently a completely new, general strategy was developed for the preparation of thermoelectric materials which allows for record-high figure of merits. It involves thermolysis of a single-source precursor in a specific IL under microwave irradiation and a subsequent mild compaction process, shown in Fig. 13.$^{179,180}$

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**Fig. 12** Schematic illustration of the synthesis of light phosphors in ionic liquids in a microwave.
Deep eutectic solvents (DES) and their general benefits as synthetic media in the manufacturing of inorganic nanomaterials

DES and properties

Deep Eutectic Solvents (DES) were first reported in 2001 by Abbott et al., who noted IL-like properties and behaviour for the eutectic compositions of certain salt: H-bond donor (HBD) mixtures, especially choline chloride (ChCl) systems, where the eutectic depression was inexplicably profound. In their broadest definition they are a combination of a hydrogen bond acceptor (HBA) which frequently is a salt, often satisfying itself the definition of an IL, alongside an H-bond donor (HBD); see Fig. 14 for some selected examples.

Strictly speaking, these eutectic mixtures are distinct from ILs. They are not necessarily ionic in nature, nor do they have the same properties as ILs, nor are they even a particularly new phenomenon, with the first reports now dating back 138 years. However, a certain class of eutectics for materials synthesis have developed into a burgeoning field worthy of attention here, where we will specifically address the applications of Deep Eutectic Solvents (DES). The field of DES has now reached 20 years of age at the time of writing, and as such the basic properties and applications of DES are well-reviewed.

The excitement around DES is largely because they offer IL-type performance in applications but with an additional molecular component (the HBD), which makes them cheap and simple to produce, while increasing the accessible chemical space of designer ‘green’ solvent systems; the discussion on IL ‘greenness’ remains equally pertinent for DES (vide supra). In terms of nanosynthesis, relevant properties include generally low surface tensions similar to ILs which promote high nucleation rates, and potential structural effects such as templating, capping, and stabilisation arising from the high ionic strength of potentially interacting compounds. Finally, as well as being solvent and template, the species of the DES can be used as reactive (i.e. non-catalytic) reagents towards materials such as (doped) carbons and polymers.

DES are most frequently mixtures of ammonium salts with H-bond rich donor species, as in the archetypal system ChCl:urea where $x_{\text{urea}} = 0.67$. However, many different systems are known and subdivided into different ‘Types’ I–V, which encompasses quaternary ammoniums, metallic DES, and hydrophobic systems with myriad HBD and salt combinations. DES can therefore be chemically similar to ILs, such that the boundaries can become blurred, with the presence of common species such as cholinium. Because the term ‘deep eutectic’ remains nebulous, works can be found where common eutectic mixtures are presented as ‘DES’ without exact characterisation of the phase diagram and behaviour. Thermodynamic interpretations suggest that the term ‘DES’ should refer only to mixtures where the melting point depression exceeds ideality. There is no arbitrary ‘eutectic depth’ criterion to be met, so given that non-ideality is relatively simple to achieve by introducing H-bonding, the scope remains broad to engineer new solvents with a wide compositional liquid window.

Nanostructure

Nanostructure is one of the key properties of ILs in nanosynthesis, so it is pertinent to revisit this understanding for DES following a few recent reviews. More advanced analyses generally refute the initial understanding of DES as simple ILs containing a complex anion–HBD chelate species at the supposed ‘magic composition’ of $1:2$, though the anion still plays a key role. Introducing another component convolutes the structure relative to ILs due to more possible interactions (i.e. HBD–HBD). The energy landscape of interactions in DES therefore appears quite flat, with many interactions, configurations and spatial arrangement of the constituents. In ILs, the default structure is of amphiphilically separated domains. These domains are evident in small-angle scattering through defined $d$-spacing and computational simulations, and are exploitable in synthesis (vide supra). In DES, the
default case is that there is not a defined d-spacing between hydrophobic and hydrophilic domains, because most common DES are generally not sufficiently amphiphilic in character, given that extensive H-bonding is generally introduced to create a strong non-ideal eutectic depression; however this motif can be induced (Fig. 15). 214,215 Moreover, small quantities of water are often introduced to DES 216 because, below a certain threshold, this affects the bulk structure negligibly217 (see Fig. 16) while improving transport properties and is thus potentially appealing in materials synthesis, where the precipitation of nanoparticles in an already highly viscous medium can cause handling difficulties. However, the addition of water can affect the interfacial structure significantly,218–220 and thus play a role in nanomaterial synthesis (notwithstanding effects such as the increased surface tension reducing nucleation rates). That is not to say that DES cannot be nanostructured nor that ILs must be. It has been observed that introducing additional H-bonding in ILs can eliminate the L3-like bicontinuous phase.218,221 More recent similar work has shown that careful design of DES, for example introducing long alkyl chains, can ‘switch on’ nanostructure, as outlined in Fig. 15.222,223 In DES such as these, hydrophobic solutes experience local compositional effects where they affiliate with specific domains within the solvent, tuning solubility and the solvation environment.224 There is therefore precedent for DES to act as multipurpose amphiphilic agents in nanosynthesis, just like ILs.225 Further structural insights will soon be attainable via computational methodologies such as AIMD (ab initio molecular dynamics)209 and MD (molecular dynamics) with polarisable force fields,226,227 in combination with experimental scattering data,206 with synergy between these bulk studies and those of the interface.220,228,229

DES as inorganic nanomaterial synthesis media

The similarities and differences between DES and ILs in nanosynthesis will now be highlighted through synthesis examples.230 By far the most popular application of DES has been in electrochemistry, especially deposition of metallic films231–239 and particles which can have elaborate high-index-terminated morphologies, possibly due to particle-DES interactions where the surface is capped along certain planes preferentially.240–243 This review will not focus on electrochemistry however, as it is extensively covered elsewhere;22,23,232 we will instead focus on comprehensively reviewing present wet-chemical and solvo-/ionothermal routes towards inorganic nanomaterials, since there is currently a gap in this area.

**Bottom-up, wet-chemical synthesis of nanoparticles employing DES**

In the first instance, DES can be used as drop-in hosts for synthesis of nanoparticles using known chemistry for the

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**Fig. 14** Example Hydrogen Bond Donor (HBD) and Hydrogen Bond Acceptor (HBA) species known to support DES formation.

**Fig. 15** EPSR model snapshots of nanostructured butylammonium bromide-glycerol DES (xglycerol = 0.67). In the right part of the figure, cations are shown in grey, NH3+ in blue, bromide in maroon, glycerol –OH in red, and CH3 in yellow. In the middle and left, respectively, ions only and glycerol only are shown, in conventional colours. Reprinted with permission from ACS.222

**Fig. 16** Cartoon diagram showing structural changes in a DES starting with the pure choline Cl:urea DES (left). As the water content increases the hydrated DES regime is reached (middle, ca. 20 vol%), then the high concentration regime where everything is fully hydrated (ca. 50 vol%). Reprinted with permission from Elsevier.207
synthesis of metallic nanoparticles. This was first shown by Liao et al. who demonstrated the synthesis of very high-index faceted gold nanoparticles in snowflake, star, and thorn morphologies, in the ChCl:urea DES.\textsuperscript{244} This work inspired numerous subsequent studies, who have shown various DES-based methodologies for the synthesis of AuNPs,\textsuperscript{245} with control over morphology\textsuperscript{246} enabling Au nanoflowers,\textsuperscript{247} nanofoams,\textsuperscript{248} highly fractal nanostars,\textsuperscript{249} and nanosheets.\textsuperscript{250} Using a ‘natural DES’ (NADES) allows this to be achieved without adding a reducing agent,\textsuperscript{251} but morphology control, such as the formation of nanosheets, can still be achieved.\textsuperscript{252} Gelated eutectics were applied in the size-controlled preparation of a series of noble metal nanoparticles.\textsuperscript{253} Much like in ILs, laser ablation can also be used in a DES to create narrow-size-distribution Ag NPs,\textsuperscript{254} and noble metal NPs such as Au assemble on the DES surface after sputtering via the same mechanism as in ILs.\textsuperscript{255} Polydisperse Pd NPs were prepared by a short heat treatment of dilute K$_2$PdCl$_4$ in ChCl:EG DES, and loaded onto N-doped porous carbon in an integrated microfluidic approach.\textsuperscript{256} Adhikari et al. introduced a methodology for the synthesis of large quantities of coinage metal colloids (Ag, Au) using a continuous flow millifluidic system. This was achieved using a designed dimethylammonium nitrate–polyl DES, which provides high atom economy and is intrinsically a capping reagent. Adhikari et al. introduced a methodology for the synthesis of large quantities of coinage metal colloids (Ag, Au) using a continuous flow millifluidic system. This was achieved using a designed dimethylammonium nitrate–polyl DES, which provides high atom economy and is intrinsically a capping reagent.\textsuperscript{257} Nanoparticulate CuCl was isolated from ChCl:urea DES, following the reaction of copper chloride with ascorbic acid in the presence of the templating agent PVP (polyvinylpyrrolidone).\textsuperscript{258} While this example was successful in CuCl preparation, it is also an example of a reaction which used an ‘off-the-shelf’ solvent, rather than taking the opportunity to use a reducing agent as the reactive HBD.

For oxides, Chen et al. used a co-precipitation method to isolate spherical magnetic Fe$_3$O$_4$ nanoparticles.\textsuperscript{259} A DES-based method was developed by Dong et al., where aqueous Tris ((HOCH$_2$)$_2$CNH$_3$) was added as an antisolvent to precipitate ZnO nanocrystals from ChCl:urea.\textsuperscript{260} Xiong et al. synthesised hematite nanospindles, also using a DES-based antisolvent method.\textsuperscript{261} Mn$_3$O$_7$ nanoparticles were prepared by reduction of Mn$^{IV}$ by ascorbic acid in hydrated ChCl:ethylene glycol DES at 40 °C.\textsuperscript{262} It has been shown by Hammond et al. that direct calcination of lanthanide nitrate hydrate:urea DES yields the corresponding lanthanide oxide or mixed hydroxide, depending on combustion temperature.\textsuperscript{197} A similar pyrolytic method where $T = 500–800$ °C was used to prepare nanoparticulate Sn/SnO$_2$@C composites from ChCl:SnCl$_2$ DES.\textsuperscript{263} Various DES such as ChCl:glucose have also been explored as reducing agents for K$_2$MnO$_4$ in the preparation of nanostructured Mn$_3$O$_7$; the high reactant concentration enabled by the DES led to the reaction to completion in less than 1 minute.\textsuperscript{264} Nanocrystalline NiO of ca. 100 nm with high energy surface facets was precipitated from ChCl:urea by addition of NaOH, with subsequent centrifugation and calcination.\textsuperscript{265}

DES have also been used in wet-chemical preparations of multinary compounds. A ChCl:citric acid DES was used to precipitate nanostructured BiOCl/BiVO$_4$ p–n heterojunctions from aqueous solutions;\textsuperscript{266} the same materials were previously prepared by sol–gel synthesis from ChCl:urea.\textsuperscript{267} Zhang et al. used the ChCl:glycerol DES as a crystallisation medium for the octahedral NiCo–NH$_3$ complex, suitable for calcination into NiCo$_2$O$_4$ nanooctahedra.\textsuperscript{268} In an interesting synthesis strategy, Ni foam was introduced into a FeCl$_3$–H$_2$O:urea DES; the abundance of redox-active Fe$^{3+}$ in the DES oxidised the foam into Ni$^{2+}$, thus creating layered NiFe double hydroxide, effective in electrolysis of urea and water.\textsuperscript{269} Nanosheets of Cu$_2$(OH)$_3$NO$_3$ were precipitated from a 44 : 1 M ratio mixture of PEG200 and NaOH, that was described as a DES, but without presented phase characterisation.\textsuperscript{270} This category of polyl–hydroxide systems were expanded to different alkali metal hydroxides and higher PEG weights (PEG400, PEG600) in the preparation of nanoparticulate transition metal oxides, cobaltites, manganites, and cuprites, under mild conditions and in sheet morphologies,\textsuperscript{271} and was also used for CdCo$_2$O$_4$.$\textsuperscript{272}$ Nanosheets of either single-metal or high-entropy metal phosphides were achieved by direct calcination of DES comprising tetrabutylphosphonium chloride:ethylene glycol, and containing either the respective metal chlorides, or an equimolar mixture of CoCl$_2$–6H$_2$O, CrCl$_3$–6H$_2$O, FeCl$_3$–6H$_2$O, MnCl$_2$–4H$_2$O, and NiCl$_2$–6H$_2$O.$\textsuperscript{273}$ Various nanoporous (silver) selenidostannates were prepared by Wang et al. from eutectic mixtures of alkylamine hydrochlorides, with interesting tuneable properties such as reversible thermochromism.\textsuperscript{274} It has been shown that the choice of DES strongly dictates the growth process.\textsuperscript{275} Even sheet and flower-like nanostructured hierarchical intermetallic nitrosulfide compounds have been synthesised: here, mixed-metallic DESs containing FeCl$_3$–6H$_2$O, CoCl$_2$–6H$_2$O and NiCl$_2$–6H$_2$O were directly annealed at 350 °C for 12 hours.\textsuperscript{276} Direct annealing has also been used to prepare Zn and Cu vanadates of the type M$_2$V$_2$O$_7$–$\alpha$ from the respective oxides dissolved in ChCl:urea.$\textsuperscript{277}$ This approach takes advantage of the often high metal oxide solubility in DES,$\textsuperscript{278}$ and was expanded to synthesis of vanadates

![Fig. 17 TEM images and particle size distribution histograms for Au nanoparticles produced in continuous flow in task-specific DES at different Ag (sic) concentrations. Reprinted with permission from ACS.$\textsuperscript{257}$](image-url)
of the type $\text{MV}_n\text{O}_{6-n}$, where the DES was noteworthy for introducing oxygen vacancies, which reduced the band gap relative to the defect-free material. Cobalt vanadate ($\text{CoV}_2\text{O}_4$) was also prepared by Thorat et al., via a sol–gel route followed by annealing, and in this case the reaction temperature was significantly reduced by the presence of the ChCl:malonic acid DES.\textsuperscript{280} Ni/Na nanocomposites have been prepared by Gage et al., who directly annealed a unique ChCl:urea:nickel acetate gel to form core–shell nanoparticles with well-defined facets. This approach shows how DES can be valuable in N-doping of materials, as well as the usual structure-directing and surfactant-free positive aspects.\textsuperscript{281} Composite tungsten-molybdenum oxides were prepared from aqueous ChCl:ethylene glycol DES, with the DES–water mixture yielding composites with prismatic morphologies.\textsuperscript{282} Calcium phosphate nanoparticles were rapidly precipitated from the reaction of $\text{CaCl}_2$ and $\text{K}_2\text{HPO}_4$ in ChCl:urea, with the final product size dictated by the synthesis temperature,\textsuperscript{283} and further by the choice of precursor reactants and solvents.\textsuperscript{284} DES have also been used in the synthesis of nanoparticulate $\text{SiO}_2$, following the common sol–gel route but substituting the DES as the reaction solvent or additive.\textsuperscript{285} A photochromic composite of $\text{WO}_3$–MoO$_3$ was formed by reaction of acetic acid with ammonium tungstate and ammonium heptamolybdate in a ChCl:EG solution, with the solvent acting simply as a low-cost synthesis medium.\textsuperscript{286} Boston et al. demonstrated a 2-step heating process using DES; first, the DES and its (hydrated) metal precursors are dehydrated at $<100 \, ^\circ\text{C}$, then calcined at high temperature to yield nanostructured materials such as $\text{BaTiO}_3$.\textsuperscript{288} Gómez Rojas et al. have been particularly prolific in their deployment of this methodology in the preparation of the superconducting materials $\text{Yb}_2\text{Cu}_3\text{O}_{7-x}$ (aka. YBCO or 123),\textsuperscript{289,290} and $\text{Bi}_2\text{Sr}_2\text{Ca}_{n-1}\text{Cu}_n\text{O}_{2n+4+x}$ (where $n = 2$; aka. BSCCO, Bi-2212),\textsuperscript{291} as well as various strontium niobates.\textsuperscript{292,293} The authors contrasted the performance of DES with ILs in this synthetic methodology, concluding that common chloride-containing DES can lead to lower phase purity through concomitant metal chloride formation, noting that the chloride phase can still recombine to the respective oxide if the calcination temperature is sufficiently high.\textsuperscript{294}  

### I onothermal methods

When used in the synthesis of materials such as zeolites and organic–inorganic frameworks, DES can function in the same way as ILs, providing solvent-directed templating of the resultant materials. This ‘ionothermal’ methodology is a rather attractive and versatile strategy,\textsuperscript{295} which has gained significant attention following its recent demonstration and review by Walton et al. and thus will also not be covered exhaustively here.\textsuperscript{87,296–299} Moreover, it is less pertinent to this review since much of the literature is focused towards open-framework structure synthesis, rather than nanoparticulate or nanostructured media.\textsuperscript{300} A DES can provide a versatile IL-type synthesis environment, at a fraction of the synthetic complexity and cost, for example in the preparation of tubular $\text{BiVO}_4$,\textsuperscript{301} Mg-Al layered double hydroxide,\textsuperscript{302} or Mn-doped $\text{SnHPO}_4$.\textsuperscript{303} Such high-temperature synthesis strategies in DES exist somewhere between the extremes of true ionothermal and solvothermal methods. Generally, compared to molecular liquids, DES have high thermal stability and lower vapour pressure. This is not always the case, of course, with low stabilities for certain DES; for example, ChCl decomposes, urea begins to thermally hydrolyse at 80 °C, and many acidic choline chloride DES readily esterify even at room temperature.\textsuperscript{304,305} Considered alongside ILs, then, the thermal stability of DES is lower, DES components tend to be more reactive and interacting, and the molecular components (as well as impurities or additives such as water) introduce a nonzero vapour pressure, which depends on the nature of the HBD. Indeed, polymorph-specific crystallisation strategies have even been developed by evaporation of volatile HBDs from DES at room temperature and pressure.\textsuperscript{306} Thus, we make the distinction here between ‘ionothermal’ processes, where the vapour pressure of the DES is low and the DES provides a templating effect, but is not itself a reactant, and ‘solvothermal’ ones.  

### Deep eutectic-solvothermal synthesis

The ‘Deep Eutectic-Solvothermal’ methodology was first introduced by Hammond et al. in the preparation of nanostructured $\text{CeO}_2$ for CO oxidation catalysis. Here, the DES is not considered a pseudo-IL as in ionothermal synthesis. In a Deep Eutectic-Solvothermal reaction, the solvent contains a significant molecular component which imposes non-negligible vapour pressures at the chosen synthesis temperature, and the solvent plays more than a spectator role in the reaction.\textsuperscript{307} In this example, the ChCl:urea ($x_{\text{urea}} = 0.67$) DES was used alongside $\text{Ce(NO}_3)_3\cdot 6\text{H}_2\text{O}$ as the cerium source. Solvothermal conditions caused thermal hydrolysis of the urea HBD into $\text{NH}_4^+$ and $\text{CO}_3^{2-}$ ions, precipitating orthorhombic monohydrated cerium oxy-carbonate $\text{Ce}_6\text{O}((\text{CO}_3)_2)\cdot \text{H}_2\text{O}$, which could then be calcined into $\text{CeO}_2$. At low water content, the surface tension of the DES is lowest, favouring rapid nucleation of small nanoparticles. Increasing the water content decreases nucleation rate, but increases the hydrolysis reaction rate by increasing the relevant reagent ($\text{H}_2\text{O}$) concentration and overall system diffusivity, which was the limiting condition. This was combined with neutron diffraction studies showing pre-structuring of reactive compounds (i.e. urea, water, nitrate) around $\text{Ce}^{3+}$ centres, potentially further facilitating rapid reaction and precipitation. Thus, through controlling water content and temperature at constant reaction time, it was possible to select the product morphology, from small nanoparticles to highly catalytically active 1D structures such as nanowires, nanorods, and cubic phases (see Fig. 18). However, it should be noted that high water contents are expected to ‘destroy’ the DES structure, such that at some point above around 50 vol% $\text{H}_2\text{O}$, all the dissolved compounds are in a fully hydrated state and bulk water begins to percolate through the mixture.\textsuperscript{217} More recently, this approach has been adapted by Exposito et al. to take advantage of the high heat- and mass-transfer conditions offered by continuous flow reactors.\textsuperscript{308} This approach allowed quantitative yields of nanostructured $\text{CeO}_2$, even with short residence times and conventional heating, overall forming a relatively mild set of synthesis conditions.
Morphology control was also achieved under deep eutectic-solvothermal conditions in a green chemical microwave-assisted preparation of small (low-temperature, low water fraction), 1D (low-temperature, high water fraction) and large crystalline (high-temperature, high water fraction) iron oxide nanoparticles. These were useful as photoanode materials for solar water splitting, imaging via superparamagnetism, or as recyclable catalysts. Here, the ChCl:urea DES was again used as a degradable reaction agent, creating basic conditions upon thermal hydrolysis of urea to precipitate iron oxides, with WAXS data showing minimal solvent degradation, except under the most extreme conditions. A number of advanced techniques were used to study this reaction in depth under conventional heating, as the first in situ study of solvothermal conditions in DES. Neutron diffraction and EXAFS measurements of the initial state showed [Fe(L)(Cl)] solvation complexes for both pure and hydrated DES where ‘L’ is an O-containing ligand. In situ SANS and EXAFS showed high initial kinetics for the conversion of the iron salt into [O–Fe–O–] networks. For the hydrated DES, these species immediately nucleated and grew into 1D structures, followed by a rapid fusion stage. For the reaction in the pure DES, nanoparticles only precipitated after 5000 s due to higher solubility of [O–Fe–O–] species, and following precipitation, the particles grew very slowly to small oblate spheroids (Fig. 19), but were screened from fusion. Isotope substitution suggested a choline-rich surface layer, with the cation acting as a capping surfactant in the same way as ILs. When the same methodology was applied using ChCl:glycerol DES instead by Zhang et al., nanoparticulate iron alkoxides were formed.

An elegant preparation of colloidal nanoparticulate Ag was shown by Adhikari et al., demonstrating that, through careful design of a silver triflate:acetamide DES, the solvent can be used without requiring additives such as metal salts and capping surfactants. Though this was deemed a wet chemical process for metallic nanoparticles as discussed vide supra, it meets our definition of ‘deep eutectic-solvothermal’, since acetamide has non-negligible vapour pressure at the chosen synthesis temperature of 200 °C, and the solvent plays an active role as template and reactive metal ion source. Under microwave irradiation, silver ions are rapidly reduced by the sole reactive additive, oleylamine, to form stable, high colloidal concentrations (up to 54 mg mL\(^{-1}\)) of relatively monodisperse [Ag\(\text{\textsubscript{3}}\text{\textsuperscript{+}}\)] nanocrystals. This derived from the group’s earlier work, where chloride-free cholinium DES (i.e. X = nitrate or acetate) were used to prepare colloidal silver nanoparticles following microwave-solvothermal treatment of solubilised AgNO\(_3\). As with the previous example, this method was shown to be far faster and less energy-intensive than using ChCl DES or neat oleylamine as the solvent in the formation of monodisperse AgNPs.
In addition to noble metals, further deep eutectic-solvothermal nanoparticulate metal oxide synthesis routes exist. Datta et al. were able to achieve morphology control of V$_2$O$_5$, and thus dictate the performance of the resultant nanostructured cathode materials, by altering the water content in a solvothermal reaction using ChCl:urea DES.$^{316}$ Using ESI-MS, the authors were able to identify the vast quantity of species formed during solvothermal decomposition of the solvent, highlighting the mixture complexity. Spherical nanoparticles of Mn$_3$O$_4$ with 25 nm diameter were prepared by reaction of KMN$_3$O$_4$ in ChCl:EG at 140 °C.$^{317}$ Li et al. prepared cubic MgO by calcination of MgCO$_3$ cubes, formed by solvothermal treatment of a MgCl$_2$-6H$_2$O:urea DES.$^{318}$

Several works have focused on the synthesis of nanostructured or mesoporous TiO$_2$ in DES, such as through sol-gel routes.$^{319}$ For example, Sandhu et al. used a novel ChCl:hydroquinone-thiourea DES in a fluoride-free synthesis of titanias with a high population of the (111) anatase surface facet, creating more active dye degradation catalysts.$^{320}$ Applying a DES based on ChCl:oxalic acid, Wang et al. were able to synthesise (001) and (101) facet-rich TiO$_2$ with morphology evocative of crassula perforata. Here, the combination of morphology, defects, and presence of surface heterojunctions gave an order of magnitude improvement in photocatalytic water splitting relative to commercial P25.$^{321}$ Similarly, a 20-fold improvement in H$_2$ production rate was observed for TiO$_2$ derived from ChCl: betaine-oxalic acid DES. The high performance was ascribed to the presence of brookite-rutile heterophase junctions, where the phase ratio could be dictated by the DES mixing ratio.$^{322}$ This highlights both one of the strengths and one of the issues of using DES in this area; as long as synthesis is conducted within the liquid window, the entire phase diagram is relevant, and it is not always productive to overfocus on the eutectic composition with respect to applications when off-eutectic mixtures can make useful or even better materials.

A number of solvothermal routes towards sulfur-containing materials have also been shown using DES. These DES usually involve thiourea, which functions as a primary or secondary HBD in the solvent, and provides a sulfur source upon thermal degradation. Jian et al. synthesised porous NiCo$_2$S$_4$ using the novel PEG200:thiourea DES, where PEG200 offered templating of hierarchical porosity under solvothermal conditions.$^{323}$ The same DES was used by Xu et al. to prepare CdS particles, which were then subjected to a second solvothermal treatment with the same DES to prepare core-shell CdS@CeO$_2$ nanocomposites.$^{324}$ Another example of this PEGylated DES is shown by Zhao et al., where it again provided sulfur and templating in the preparation of Fe$_3$S$_4$ nanosheets, whose porosity yielded strong performance in the electrochemical fixation of N$_2$.$^{325}$ Ni$_3$S$_2$ nanosheets were obtained by solvothermal treatment of electrodeposited Ni-Co on nickel foam, with a ChCl:EG DES containing low concentrations of thiourea.$^{326}$ A similar solvothermal sulphurisation method was used to obtain nanospheric Ni$_3$S$_2$, with strong supercapacitance.$^{327,328}$ Low concentrations of thiourea and metal chlorides in the ChCl:urea DES yielded chalcopryte CuInS$_2$ in a nanorod morphology under microwave irradiation.$^{329}$ Nanoparticles of the popular kesterite semiconductor Cu$_2$ZnSnS$_4$ (CZTS) with a high defect concentration were prepared by Karimi et al. in their solvothermal treatment of the respective metal chlorides, using thiourea as a sulfur source, from the ChCl:urea eutectic.$^{330}$ In the ChCl:EG DES, NiS$_2$ nanospheres were synthesised by solvothermal treatment of nickel sulfate, sodium thiosulfate and EDTA, making it noteworthy as the only synthesis not to use thiourea as a sulfur source.$^{331}$

Similar methodologies have been used to prepare multinary and composite compounds in DES under solvothermal conditions. Spinel ferrites of the form MFe$_2$O$_4$ (M = Mg, Zn, Co, Ni) were synthesised at much lower reaction temperatures than comparable techniques by using a ChCl:maeleic/malonatic/oxalic acid, or ChCl:urea or ChCl:EG DES.$^{332,333}$ Jiang et al. used a ChCl:triethanolamine DES to template a 3D porous nanostructure for NaTi$_2$(PO$_4$)$_3$, interesting as an anode for sodium-ion batteries once calcined with sucrose to cast it onto a carbon substrate.$^{334}$ In a control experiment, it was shown that aqueous mixtures of the HBD and salt respectively formed prismatic and globular nanoparticles. Various metal molybdates (M = Mg, Fe, Mn) have been prepared by Baby et al. using ChCl:urea in solvothermal reactions.$^{335}$ The controlled degradation of the DES was found to template more desirable active morphologies useful in toxin sensing, [Co(NH$_3$)$_4$CO$_3$]Cl with a porous 2D nanosheet morphology was synthesised by Liu et al. solvothermally; following ‘activation’ by casting onto Ni foam, the amorphous CoOOH performed well in the oxygen evolution reaction.$^{316}$ Narayanam et al. used solvothermal heating of a volatile ChCl:phenol DES to produce crystaline polyoxo-titanium clusters, relevant as molecular analogues of TiO$_2$, with nuclearities of up to 28.$^{337}$ RuCu$_2$O$_4$ is a mixed oxide showing high electrical conductivity, stability, and electrochemical performance, and thus interesting in hybrid supercapacitor cells; Karade et al. were able to synthesise this material in nanofibrous form under solvothermal conditions with a rather high water content of 50 vol% in their DES.$^{338}$ NiCo$_2$O$_4$ nanorods were solvothermally grown on pre-prepared MoS$_2$ nanosheets in ChCl:urea by Zhao et al. (110 °C, 16 h),$^{339}$ and on reduced graphene oxide by Ni et al.$^{340}$ Alloyed nanoclusters of PtCu were solvothermally grown on multi-walled carbon nanotubes in a DES, with the solvent apparently facilitating a controlled dispersion of nanoparticles on the nanotube surface.

**Scope of DES in nanomanufacturing**

Having reviewed the applications of DES in nanomanufacturing, it is pertinent to comment on their scope in the field. Clearly, DES show many of the same promising features of ILs, such as structure direction, morphological templating and assembly control, as well as potentially lower toxicity and improved environmental credentials. DES have potentially great potential, since the addition of another species (the HBD) widens the accessible phase space from the pure ionic melt to a wide range of ionic–molecular mixture compositions. They also show significant capability as drop-in replacements for conventional solvents due to these advantages.

DES also share many of the drawbacks of ILs, especially the potentially enormous viscosity,$^{215}$ degradation,$^{341}$ solution
complexity,\textsuperscript{312} and not being intrinsically green nor responsibly sourced.\textsuperscript{187} Moreover, all of the described examples derive from known chemistry. This applies whether the example methodology is: 1. simple, such as using a DES as a purely spectating solvent host for a known reaction; 2. more advanced, such as using a DES as an ionothermal synthesis template facilitating higher reaction temperature without autogenous pressure; 3. the most intricate, using a DES in \textit{i.e.} a microwave-solvolothermal reaction where it functions as solvent, template, and reactant simultaneously. However, the most exciting aspect of DES and ILs is in their designer nature, where they can potentially be altered to make new chemistries viable.\textsuperscript{342} Thus, it will be exciting to watch for new, advanced synthesis protocols with high-level design. We foresee many novel applications of these solvents in ‘tuneable’ sense, societal welfare will be improved through safer, more energy efficient and benign production of nanomaterials.

**Insights**

The lessons learned from these examples are therefore:
- By employing ILs and DES in the manufacturing of nanomaterials it is possible to obtain materials that cannot be obtained using conventional routes.
- The nature of the liquid critically determines the particle morphology and size.
- The IL or DES controls the assembly process of nanostructures.
- ILs and DES enable phase-selective synthesis.
- In contrast to classical surfactants and NP stabilisers, the nanostructures can be cleaned easily of ILs and DES, a requirement for many applications.
- The use of a task-specific IL or DES can prevent the use of toxic and hazardous substances.
- Uncommon, energy-efficient conversion methods like microwave and ultrasound heating synergise with ILs and DES to enhance the manufacturing of nanomaterials.

**Conflicts of interest**

There are no conflicts to declare.

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**Conclusions**

The presented examples illustrate the transformative character and unique opportunities that ILs and DES have in the deliberate, designed synthesis of nanomaterials. Because of their high modularity, they can be tailored for a specific synthesis through chemical structure variations unlike any conventional solvents. Unique to ILs and DES is their truly multifunctional character, which allows them to serve as more than just the solvent and reaction medium. With ILs and DESs, properties and property combinations can be achieved that are unattainable with conventional solvents. This also allows the use of unconventional synthetic methods, and the development of new techniques.

It is expected that the presented, universal methods for the production of nanomaterials from ILs, and soon also from DES, will allow for the manufacturing of improved products and devices and open up new horizons for nanosynthesis. The methods are generally, faster, safer, more energy and atom (material)-efficient, use smaller quantities of less toxic chemicals, negate the use of auxiliary substances (such as surfactants and stabilisers), reduce the risk of pollution, reduce and prevent waste, and thus overall, they reduce the risk to human health and minimise the impact on the environment, while being economically viable. In addition, these unconventional production methods will allow us to develop new and improved nanomaterials, pertinent for a wide number of applications such as catalysis and wastewater treatment, solar cells and solid-state lighting as well as biomedicine. Solutions beyond the current leading nanomanufacturing technologies will be provided that advance and innovate nanomanufacturing to achieve sustainability.

This will, on one hand, lead to new technological and scholarly knowledge with respect to understanding the designed manufacturing of nanomaterials and the discovery of new materials. In a broader sense, societal welfare will be improved through safer, more energy efficient and benign production of nanomaterials.

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