Ionic Liquid Microcapsules: Formation and Application of Polystyrene Microcapsules with Ionic Liquid Cores.

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ABSTRACT

The synthesis of 2-3 µm spherical microcapsules with a polymer shell and a liquid core comprised of ionic liquid (IL) ([Bmim][NTf2]) is described. These discrete IL microcapsules are prepared quickly and in large quantity in a low temperature, one pot synthesis, by a modified coacervation technique. These IL microcapsules show ability to release dye from the IL core into solution through a polymer membrane and also concentrate metal ions from solution into the microcapsules.

Keywords
Core-shell, ionic liquid materials, dye release, extraction, polymer membrane.

INTRODUCTION

We present spherical microcapsules of polystyrene containing ionic liquid in their core. Materials such as these have many potential uses in sustainable applications. Ionic liquids, (ILs) are liquids composed entirely of ions.1,2,3 They have a remarkable ability to be tuned for specific application, providing new approaches in science, technology and industry. Since their early use as replacements for organic solvents,4 uses of ionic liquids continue to expand. In particular, there has been a lot of recent interest in ionic liquids and ionic liquid materials in energy and electronics.5,6,7 Catalysis continues to feature heavily as an area which has seen benefit from ionic liquids8 as does environmental remediation such as CO2 uptake,9 energy materials10 and metal recovery.11 Steady interest and new applications are coming from the pharmaceutical, drug delivery and medicinal areas10, 12, 13, 14 as well as the food industry.15

In the pharmaceutical fine chemicals sector, the use of ionic liquids in drug16 and fragrance delivery17 has been noted. Ionic liquids are being used as drugs themselves. Davis and co-workers18
synthesised an alkylated derivative of miconazole (an antifungal drug), which was combined with [PF$_6$]$^-$ to form the ionic liquid, alkylmiconazolium hexafluorophosphate. The same group reported formation of ILs from artificial sweeteners combining the anions of saccharin and acesulfame with organic cations.$^{13}$ Ionic liquids as drugs are seen to have many potential benefits.$^{19,20}$ Most recently an oral delivery method for insulin featuring an ionic liquid made from choline and geranic acid (CAGE),$^{21}$ which has shown to be effective in protein stabilisation over long periods. Previous work using CAGE showed stabilisation of protein drugs for transdermal delivery.$^{22}$ However, there are some drawbacks to the large scale use of ionic liquids. These include separation issues, fear of possible release of material to the environment, loss of liquid electrolyte from electronic devices and contamination of products in reactions. Finding ways to circumvent these issues using materials synthesis is an important goal. In particular Ionic liquid gels$^{23}$: sol-gel, polymer, and low molecular weight ionic liquid gels$^{24,25}$ continue to play a role. Supported ionic liquid acids,$^{26}$ bases$^{27}$ and entrapped catalysts$^{28}$ are facilitating the use of catalysts with easy separation. Confining ionic liquids can broaden their application further and also help to build theoretical models of IL behavior.$^{29,30}$ Whether for use in catalysis or environmental clean-up, ionic liquids containing actives of high value need to be easily separated for recycling. Extraction of active products or recovery of metals into ionic liquids currently requires large scale separations. Using ionic liquids for extractions$^{31,32}$ could be easily enhanced by using a suitable membrane technology. Confinement of ionic liquids within a membrane would facilitate their use in a wide range of industrial and academic disciplines. The use of ionic liquid membranes shows promise in many applications$^{33}$ but the encapsulation or confinement has yet to be achieved easily in 3 dimensions. Examples of spherical encapsulated ionic liquids are rare. Previous attempts to produce ionic liquids confined within polymer shells include individual coating of spherical IL drops with a polymer$^{34}$ and formation of microgels,$^{35}$ both of these methods have limitations for large scale production. Encapsulation is the process by which a
material is contained within a shell of a second material. Liquid core microcapsules are used for a variety of purposes from flavor protection in food to delayed release of pharmaceuticals. Release from microcapsules can be achieved by a number of mechanisms. For hard rigid shells, such as metals or inorganics a brittle fracture, due to shear or ultrasound can be achieved. For polymeric shells a slow diffusional release profile is more common. There are many possible routes to making microcapsules. These include direct polymerization from a surface layer by layer deposition or self-assembly of particles at a templating interface.

EXPERIMENTAL

We have prepared spherical microcapsules of polystyrene containing ionic liquid in their core. This is achieved at low temperature in one pot with a resultant large quantity of spheres. The basis of the method used in this work is adapted from the work of Loxley and Vincent and shown schematically (Figure. 1) In the original method the polymer shell material is insoluble in the core, which contains a low volatility organic solvent. The two components are mixed with a volatile solvent, typically dichloromethane, forming a single organic phase. This is emulsified in water, containing a small amount of surfactant. The resulting emulsion droplets are templates for the microcapsules. As the volatile solvent evaporates the polymer precipitates and, depending on the wetting conditions, forms a shell surrounding the organic core. We postulated that it may be possible to replace the low volatility organic phase with an ionic liquid. Thus forming microcapsules of ionic liquid contained in a polystyrene shell. This proved to be successful (SI S1.2 Methods).

RESULTS AND DISCUSSION

Initial studies on dye release from the ionic liquid core through the polystyrene membrane, and metal uptake through the polystyrene membrane into the ionic liquid core were performed. The microscopy images (Figure. 2) and (SI S5.0) show that we have successfully produced ionic liquid
core microcapsules. These are easily made in the liter scale. The capsules have a size of a few microns (~2 µm). This is set by the emulsification technique. Smaller capsules can be achieved, for example with vigorous emulsification or the use of ultrasound. The shell thickness is simply determined by the concentration of polymer solution. The release of dye from the capsules (Figure. 3) follows the expected trend, with more polymer in the shell resulting in slower release. The timescale for release is given by the inverse of the rate constant, which is calculated from the graph (Figure. 4).

![Diagram of dye release experiments](image)

Figure 3 Schematic of dye release experiments with ionic liquid core microcapsules: a) thymol blue Dye + IL, b) diffusion of dye through polymer shell, c) remaining IL in core after dye diffusion.

The numerical values are shown in the SI, (SI, S6 Dye release, table S1). We assume that all dye is partitioned into the core and that release follows a diffusional timescale, \( \tau \), such that \( \tau = T^2/D \), with \( T \) the shell thickness and \( D \) the diffusion coefficient of dye through the solvated polymer. We derived a simple model to estimate this timescale. The total number of capsules is \( N \) and the volume of the core is \( 4/3 \pi R^3 \). We therefore have
\[ NV_{\text{core}} = \frac{m_{IL}}{\rho_{IL}} = N \frac{4\pi}{3} R^3 \quad \text{(EQ 1)} \]

With \( m_{IL} \) the mass of ionic liquid added to the mixture and \( \rho_{IL} \) the ionic liquid density. The volume of the shell, \( V_{\text{shell}} \), is given by

\[ NV_{\text{shell}} = \frac{m_p}{\rho_p} \quad \text{(EQ 2)} \]

with \( m_p \) the total mass of polymer and \( \rho_p \) the polymer density. The overall volume balance provides

\[ V_{\text{core}} + V_{\text{shell}} = \frac{4\pi}{3} (R + T)^3 \quad \text{(EQ 3)} \]

then

\[ \left( \frac{m_p}{\rho_p} + \frac{m_{IL}}{\rho_{IL}} \right) = \left( \frac{4\pi N}{3} \right)^{1/3} \left( D \tau \right)^{1/2} + \left( \frac{m_{IL}}{\rho_{IL}} \right)^{1/3} \quad \text{(EQ 4)} \]

In this work \( m_{IL} \) is 3.869 g and \( \rho_{IL} \) is 1437 kg/m³. The polymer density \( \rho_p \) is taken as 1050 kg/m³. This provides \( m_{IL}/\rho_{IL} \) as \( 2.69 \times 10^{-6} \) m³. Plotting \( (m_p/1050 + 2.69 \times 10^{-6})^{1/2} \) versus \( \tau^{1/2} \) then produces a straight line as shown in (SI Figure S5) The slope of the best fit line is \( 1.63 \times 10^{-9} \) m s \(^{-1/2} \) and the intercept is given by 0.014 m. Estimating the radius of the capsules, \( R \), to be \( \sim 2 \) µm, EQ 1 provides the number of capsules \( N \) to be \( 8 \times 10^{10} \). This allows an estimate of the diffusion coefficient, \( D \), of thymol blue through polystyrene of \( 5.5 \times 10^{-18} \) m²s⁻¹. This is a sensible order of magnitude for the diffusion of a molecule through a polymer.

Table 1

<table>
<thead>
<tr>
<th>Time in solution (hours)</th>
<th>Integrated (^7)Li intensity (a.u.)</th>
</tr>
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<tbody>
<tr>
<td>0.5</td>
<td>471 ±6</td>
</tr>
<tr>
<td>2.5</td>
<td>482 ±6</td>
</tr>
<tr>
<td>73</td>
<td>488 ±6</td>
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</tbody>
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For metal uptake experiments a 1M solution of Li⁺ was prepared from its chlorinated salt. Samples of microcapsules were then exposed to the Lithium solution (SI S4). Lithium NMR was used to measure the Ion uptake into capsules and this shows that Lithium ions are extracted from the salt solution very rapidly, within the 30 minutes of the first measurement. This implies that the ionic liquid capsules are easily saturated with lithium. (Figure 5) The average error is calculated from the standard deviation of the repeats. (SI S7 metal ion uptake).

CONCLUSION

We have demonstrated the preparation and use of ionic liquid core polymer shell particles. The particles are easy to reproducibly prepare. Confining the ionic liquid in this way provides not just a practical way of using materials but also a neat discrete particle for fundamental study of ionic liquid behaviour in a confined space. We have shown the ability of the ionic liquid core to contain a model active compound, and to release it through the polymer membrane. The rate of release of a dye from intact capsules is shown to be determined by the diffusion of dye through the polymer shell. Despite diffusion coefficients as low as $10^{-18} \text{m}^2\text{s}^{-1}$ the release from ~2 µm capsules occurs within a few hours. We also show the ability of metal ions to diffuse from a solution outside of the microcapsule and be concentrated within microcapsule (Figure 5).

In this work we show that ionic liquid core polymer shell materials can be made in significant quantities. Ionic liquid core polymer shell materials are likely to have several specific practical advantages: they are easy to filter/recover; they can act as distinct confined containers; their uniformity of size gives potential for use in liquid flow systems for printing of ionic liquid based e-inks or painting on to electrodes and surfaces. The ionic liquid core polymer shell particles can be tailored to allow diffusion out of or into the material as well as release at a specific temperature (melting point of polymer shell) or through mechanical destruction. Furthermore, ionic liquids can be chosen/tailored for their functionality, to extract or to release a target of interest, such as drug delivery or metal recovery.
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Supporting information
Details of materials, methods and characterisation.

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SYNOPSIS

Ionic liquid contained within discrete self-organized polymer spheres (microcapsules). A low temp, high yielding, one-pot route to polymer membrane ionic liquid core microcapsules for sustainable applications.