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Separating convective from diffusive mass transport mechanisms in ionic liquids by redox pro-fluorescence microscopy

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ABSTRACT. The study of electrochemical reactivity requires analytical techniques capable of probing the diffusion of reactants and products to and from electrified interfaces. Information on diffusion coefficients are often obtained indirectly by modelling current transients and cyclic voltammetry data, but such measurements lack spatial resolution and are accurate only if mass transport by convection is negligible. Detecting and accounting for adventitious convection in viscous and wet solvents, such as ionic liquids, is technically challenging. We have developed a direct, spatiotemporally resolved optical tracking of diffusion fronts which can detect and resolve convective disturbances to linear diffusion. By tracking the movement of an electrode-generated fluorophore we demonstrate that parasitic gas evolving reactions lead to ten-fold overestimates of macroscopic diffusion coefficients. An hypothesis is put forward linking large barriers to inner-sphere redox reactions, such as hydrogen gas evolution, to the formation of cation-rich overscreening and crowding double layer structures in imidazolium-based ionic liquids.

INTRODUCTION

Room temperature ionic liquids (RTILs) are salts with melting point below 25 °C.1 They are non-volatile ionic conductors, with emerging applications in technologies and processes ranging from batteries,² to supercapacitors,³ electrocatalysis,⁴ and sensing.^{5,6} For all RTIL-based electrochemical applications it is important to first obtain a correct and complete understanding of mass transport of reactants and products towards and away from the electrode. 7-9 Equally important is to access quantitative information on structures and dynamics of the electrode-RTIL interface. 10-14 For instance, insights on double layer order obtained by atomic force microscopy (AFM), 15, 16 Raman spectroscopy, 17 and more recently even through simple measurements of open circuit potential, 13 have shown the existence of compact cation or anion-rich layers near polarized electrodes.¹³ It can be supposed that such highly ordered and compact RTIL arrangements, which can persist for several hours if formed in response to negative electrode biases, 13 will introduce a kinetic limitation to heterogeneous charge-transfer reactions, especially for reactions highly dependent on the nature of the electrode interface, such as inner-sphere reduction of water, oxygen or protons. 18, 19 Opposite to outer-sphere reactions, which proceed at appreciable rates even when there exist a solvent layer between electrode and reactant, inner-sphere reactions require a

strong interaction between electrode's surface and molecule being oxidized or reduced. $^{\rm 18,\,20,\,21}$

Hence a near-surface compact arrangement of large organic cations is likely introduce a barrier to a common unwanted inner-sphere reaction: evolution of hydrogen as bubbles. 18, 22 Parasitic evolution of gas bubbles at electrodes is a common occurrance, 23-26 and partial masking of the electrode's surface by a gas cavity can decrease its capacitance, 27 or cause an increase in electrical conductivity during electrolysis. ²⁸⁻³⁰ Beside the loss of electroactive area, a convective contribution to mass transport will appear as the surface-pinned bubble grows and eventually departs the electrode.31 Such bubble-induced "disturbance" to diffusion – a localized and transient stirring – is often neglected and/or hard to account for.32 In this work we seek to develop a simple tool to quantify, and to map across a macroscopic electrode, the impact of adventitious gas evolution reactions on the analysis of mass transport in RTILs. The study of reactant and product diffusivity is a central feature of most electrochemical investigations, 33, 34 but in conventional "oneelectrode, one-lead"35 measurement, local information on convection disturbances to quiescent diffusion are inevitably lost.

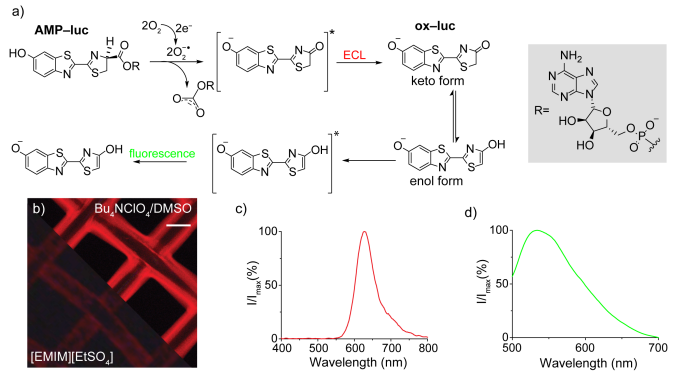


Figure 1. (a) Proposed mechanism for the cathodic electrochemiluminescence (ECL) of AMP–luc. (b) The reaction of AMP–luc (4.0 × 10⁻³ M, oxygen-saturated, -2.0 V vs Ag/AgCl) with electrode-generated superoxide leads to the excited state of ox–luc, which relaxes emitting red light. The red ECL emission is intense for electrolyses in 2.0 × 10⁻¹ M Bu₄NClO₄/DMSO (10× magnification, CMOS sensor camera, unmodified micrograph, Video S1) but very faint in [EMIM][EtSO₄] (Laowa 25 mm F/2.8 2.5–5X, backilluminated CMOS sensor, micrographs edited to maximise contrast). The scale bar is 100 μm. (c) Cathodic AMP–luc ECL spectrum recorded in [EMIM][EtSO₄]. (d) Optical excitation of an electrolyzed AMP–luc/[EMIM][EtSO₄] triggers the green fluorescence of the ox–luc enol product.

³⁶⁻³⁸ In the 1990s Engstrom and co-workers began developing fluorescence microscopy-based approaches to access two-dimensional information on concentration gradients across macroscopic electrodes.³⁹⁻⁴¹ The same group also applied electrochemiluminescent (ECL) reactions to map current heterogeneity due to non-uniform diffusion at the edge of microelectrodes,⁴²⁻⁴⁴ demonstrating the electroanalytical value of optical techniques.⁴⁵⁻⁴⁷ Other groups have since then explored similar research lines, and have for example by developing electrofluorochromic redox couples to monitor the evolution over time of electrode concentration profiles.^{48, 49}

In this paper we develop a strategy for the visualization of diffusion fronts by means of tracking the movement of electrochemically generated fluorophores. Such spatiotemporally resolved data on diffusion coefficients allowed us to detect and quantify bubble-induced convective disturbances of nominally quiescent electrode systems. We have applied this optical approach – redox profluorescence microscopy – to RTILs that have a different propensity of forming compact double layer cation-rich structures, hence a different intrinsic inhibition of adventitious inner-sphere gas-evolving redox reactions. The electrode-generated fluorophore is oxyluciferin (**ox-luc**, Figure 1a), ⁵⁰ which is the final product of the cathodic electrochemiluminescent (ECL, Figure 1b and Video S1, Supporting Information) light-path of non-fluorescent firefly's luciferin adenylate ester (**AMP-luc**). ⁵¹

EXPERIMENAL SECTION

Materials. Unless stated otherwise, all reagents were of analytical grade and used without further purification. Milli-QTM water (> 18.2 M Ω cm) was used for cleaning procedures and, where specified, for adjusting the water content of the RTIL samples. D-luciferin sodium salt (≥95%, Cayman Chemical Company, Michigan) was used as starting material for the synthesis of firefly luciferyl adenylate (hereafter AMP-luc). **AMP-luc** was prepared and titrated as described previously.⁵¹, ⁵² Pyridine (≥99.5%) and acetonitrile (≥99.9%, MeCN) were purchased from Honeywell (North Carolina). 1-Butyl-1bis-(trifluoromethylsulfonyl)imide methylpyrrolidinium (99.5%, [BMPyrr][NTf₂]), 1-ethyl-3-methylimidazolium tetrafluoroborate (>98%, [EMIM][BF₄]) were purchased from Iolitec (Germany). Adenosine 5'-monophosphate monohydrate $(\geq 97\%, AMP), N,N$ -dicyclohexylcarbodiimide (99%), 1-ethyl-3-methylimidazolium ethyl sulfate (≥95%, [EMIM][EtSO₄]), tetrabutylammonium hexafluorophosphate $(\geq 99.0\%,$ Bu₄NPF₆), bis(cyclopentadienylcobalt(III) hexafluorophosphate (98%, Cc⁺), bis(pentamethylcyclopentadienyl)cobalt(III) hexafluorophosphate (98%, Me₁₀Cc⁺) and ferrocene (98%, Fc, sublimed before use) were purchased from Sigma. The water content in RTIL samples was estimated by Karl-Fisher (KF) titrations using a C20S compact KF coulometer (Mettler-Toledo, Ohio). HydranalTM Coulomat AG reagent for the KF titration was purchased from Honeywell (North Carolina), and the 0.1% water standard was from Merck (Germany). At least three KF titrations were performed for each RTIL. The reported values ([EMIM][EtSO₄], 930 ppm; [BMPyrr][NTf₂], 125 ppm; [EMIM][BF₄], 2400 ppm) are the arithmetic average of the three titrations.

Fluorescence and ECL imaging. Unless specified otherwise, time-resolved fluorescence and ECL images were recorded on a Nikon ECLIPSE Ti2-U inverted microscope fitted with a custom-built optoelectrochemical single-compartment, three-electrode cell (Figure S1, Supporting Information). Microscopy experiments were performed in a dark room, at room temperature (23 \pm 2 °C) in air, and within 10 min of exposing the electrolytic AMP-luc solution to the atmosphere. For the fluorescence experiments the microscope was fitted with a Plan Apo λ 10×/0.45 objective (part n. 88-379, Nikon, CFI Plan Fluor), a back-illuminated CMOS monochrome camera (DS-Qi2, Nikon), and a FITC filter/dichroic mirror cube (LED-FITC-A-NTF-ZERO, single band excitation filter 461.0-487.5 nm, single band emission filter 502.5–547.5 nm, and 495 nm dichroic beamsplitter, BrightLine®, Semrock, California). Fluorescence videos were recorded selecting a 20 ms exposure time, a capture rate of 5 frame/s, and a 1× gain. Microscopy experiments to map ECL in organic solvents (AMP-luc in Bu₄NClO₄/DMSO, e.g. video frame in Figure 1b, upper right section) were recorded using a colour camera (DS-Fi3, Nikon), setting the exposure to 1.0 s, and without filtering nor light excitation. ECL intensities for solutions of AMP-luc in RTILs were however to dim to be captured with the DS-Fi3 camera, so that ECL images (e.g. lower left section of Figure 1b, cathodic electrolysis of AMP-luc in [EMIM][EtSO₄]) were recorded with a more sensitive DSLR camera (Nikon D850) equipped with back-illuminated CMOS image sensor and using an ultra macro lens (Laowa 25 mm F/2.8 2.5-5X). The fluorescence of non-fluorescent **AMP-luc** solutions $(4 \times 10^{-3} \, \text{M}, \sim 10 \, \text{mL})$ was triggered electrochemically by applying a short (30 s) cathodic bias (-2.0 V vs reference electrode) to a platinum mesh working electrode (SEC-C Gauze, 80 mesh, 80 µm wire diameter, 7 × 6 mm outer size, purchased from BASi, Indiana). The reference electrode was a plastic body "leakless" Ag/AgCl electrode (eDAQ, part n. ET072-1, with 3.4 M aqueous potassium chloride as filling solution) and a platinum coil served as the counter electrode (0.5 mm diameter wire, 99.99+%, Goodfellow Cambridge Limited). The electrodes were connected to a portable potentiostat (Emstat3 blue, PalmSens BV, Netherlands). The **AMP-luc** solutions were bubbled with oxygen gas (>99.95%. Coregas) for at least 20 min prior to the experiments. The platinum electrodes were cleaned daily through at least 50 consecutive cyclic voltammetry cycles (from -2.0 V to 1.0 V, potential scan rate of 0.01 V s⁻¹) in 5.0×10^{-1} M aqueous sulphuric acid. Local diffusivity values were calculated by analyzing the movement of the **ox-luc** fluorescent front away from the platinum surface. Time-stamped fluorescence intensity profiles were analysed with the open source image processing package Fiji, 53 in order to determine the distance (r) from the electrode surface at which the fluorescence intensity falls to half of its maximum value. This distance (r) was tracked as function of the time (t)after the cathodic pulse, and used to estimate diffusion coefficients (D) by assuming an Einstein's random walk ($r^2 = 2Dt$).⁵⁴, 55 This procedure was repeated for 10 different videos for each RTIL type, with at least 10 different locations analyzed in each video. Fluorescence images were not background subtracted,

and no attempts were made to improve contrast and sharpness. ECL images for RTIL samples were background-subtracted.

Photon counting. A single-photon counting module (SPCM-AQR-14, Excelitas Technologies), interfaced with an avalanche photodiode (APD) controller (Nanonics Imaging Ltd. time constant was set to 1.0 ms), was used to perform quantitative ECL measurements. A data logger (DrDAQ, Pico Technology) was used to record photon count rates. Electrolytic **AMP-luc** solutions and the three electrodes described for the fluorescence imaging experiments were placed inside a quart cuvette (10 mm optical path, Starna Pty Ltd., Australia). The cuvette was fitted with a custom PTFE cap/electrode holder to ensure reproducible electrodes positioning.

Fluorescence and ECL spectroscopy. ECL and fluorescence spectra were recorded on a Cary Eclipse (Varian, California) fluorescence spectrophotometer operated either in Bio/Chemi-luminescence mode (ECL, no excitation) or in Fluorescence mode (474 nm, excitation). The experiments were performed using the cuvette described in the Photon counting section, and the electrodes are described in the Fluorescence and ECL imaging section. The spectrometer's photomultiplier voltage was set to 800 V and the emission slit to 20 nm for both ECL and fluorescence measurements. The fluorescence excitation slit was set to 20 nm.

Open circuit potentiometry. Time-resolved open circuit potential (OCP) measurements were performed with a Emstat3 Blue potentiostat (PalmSens BV, Netherlands). A small RTIL sample (~5 mL) was loaded in a 10 mL beaker and degassed by means of bubbling argon gas (99.997%, Coregas) through it for at least 20 min. A platinum mesh (SEC-C Gauze, 80 mesh, BASi, Indiana) was used as working electrode, a platinum coil as counter electrode, and an Ag/AgCl "leakless" as the reference electrode (eDAQ, part ET072-1). The reference electrode potential was calibrated against the apparent formal potential of the ferrocene/ferricenium couple (Fc/Fc⁺) as measured by cyclic voltammetry at a platinum disk electrode (eDAQ, ET052, 3 mm diameter) using the above specified reference and counter electrodes and a 1.0×10^{-3} M ferrocene solution in 1.0×10^{-3} M Bu₄NClO₄/MeCN. The working electrode was polished with alumina slurry (0.05 µm, eDAQ, ET033) and then cleaned electrochemically as described in the Fluorescence Imaging section. The OCP measurements were started immediately after a 60 s cathodic pulse (-2.0 V relative to the initial, rest, OCP). To determine the working electrode's rest OCP, electrodes were let to equilibrate in the RTIL sample for at least 5 min after the end of the bubbling procedure. The length of the equilibration period was such to generally obtain a sufficiently stable OCP (dV/dt below |0.0001|). All OCP experiments were performed inside an acrylate glove box (Molecular Imaging, model GB306) kept under nitrogen gas that was dried with a DrieriteTM gas-drying unit.

Cyclic voltammetry. Cyclic voltammetry experiments were carried out inside a custom-built acrylate glovebox kept under positive argon flow (4.0 L×min⁻¹, >99.999%, Coregas, Australia) using an Emstat3 Blue potentiostat (PalmSens BV, Netherlands) and a single-compartment, three-electrode glass cell. The cell was loaded with ~25 mL of either ferrocene (Fc), cobaltocenium hexafluorophosphate (Cc^+), or decamethylcobaltocenium hexafluorophosphate ($Me_{10}Cc^+$) solutions (1.0×10^{-3})

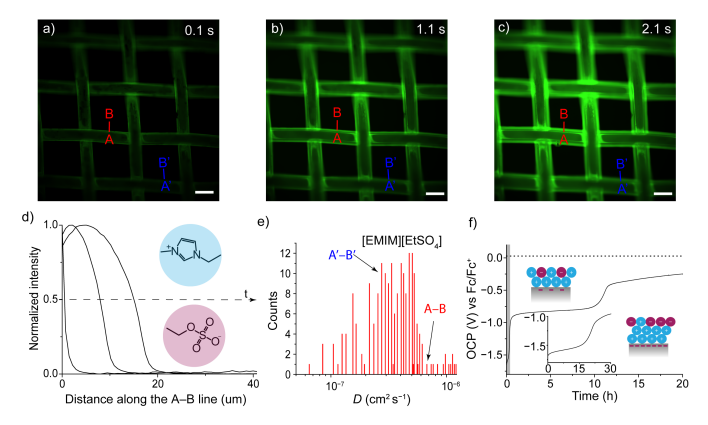


Figure 2. (a–c) Selected time-stamped fluorescence micrographs (10× magnification) recorded during the cathodic electrolysis (-2.0 V vs Ag/AgCl) of an oxygen-saturated AMP–luc solution (0.4 × 10⁻³ M in [EMIM][EtSO₄]) at a platinum mesh electrode (Video S2, Supporting Information). Scale bars in (a–c) are 100 μm. (d) Representative fluorescence intensity plot profiles, sampled along the A–B line marked in (a–c), recorded 0.1, 1.1 and 2.1 s after the working electrode bias is stepped from open circuit to -2.0 V. The analysis of the movement over time of the superoxide diffusion front (triggering the ox–luc formation), as shown in (d), was repeated for at least 20 different working electrode locations per experiment, and repeated in at least 10 different independent experiments to build the histogram plot shown in (e). (f) Representative OCP–time measurement for a platinum mesh working electrode immersed in [EMIM][EtSO₄] (930 ppm of water content). The electrode's OCP was recorded continuously for 20 h, starting after the application of a 60 s negative pulse (-2.0 V vs the initial OCP). The horizontal dotted line represents the initial (rest) OCP value. The cartoons show the schematic depiction of overscreening and crowding near-electrode cation-rich structures RTILs. 11, 13 The overscreening OCP signature is stable for several hours, while the crowding (inset) lasts for ~15 min). The grey shad area in (f) indicates the data expanded in the figure's inset.

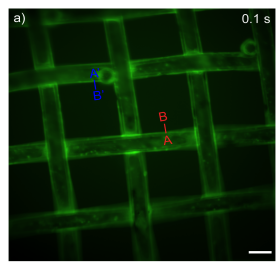
M) in RTILs. Solutions were degassed prior to the measurements by means of a 20-min argon bubbling procedure. The working electrode was a platinum disk (eDAQ, ET052, 3 mm diameter), the counter electrode a platinum coil and the reference electrode a "leakless" Ag/AgCl electrode (eDAQ, part ET072-1). The working electrode was first polished with alumina slurry (0.05 μm , Dace Technology, Arizona) and then cleaned as described in the Fluorescence and ECL imaging section.

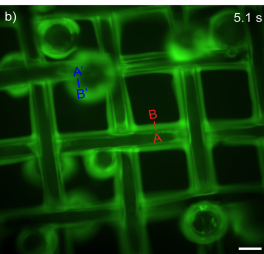
Digital simulations of voltammograms. Digital simulations of cyclic voltammograms were run on DigiElch8 (Gamry Instruments, Pennsylvania). Kinetic, thermodynamic and transport parameters were simulated by fitting data sets that covered a broad range of scan rates (from 1.0×10^{-2} to 5.0 V s^{-1}), and assuming an E mechanism. Charge transfer kinetic parameters were estimated using Butler–Volmer kinetics. Diffusion was modelled as semi-infinite 1D diffusion, and the symmetry factor, α , was set to 0.5. The cell iR drop was left uncompensated during the measurements and the actual value measured by electrochemical impedance spectroscopy prior to each

experiment and then used for the simulations. Cells resistance values were approximately 400 Ω for [EMIM][EtSO₄], 570 Ω for [BMPyrr][NTf₂] and 150 Ω for Bu₄NPF₆/MeCN.

RESULTS AND DISCUSSION

As shown in the top-right section of Figure 1b, the cathodic electrolysis of an oxygenated solution of **AMP-luc** in an organic solvent-based electrolyte (DMSO/Bu₄NClO₄) leads to a red glow visible to the naked eye (Figures S2–S3, Supporting Information). The same electrolysis performed in RTILs, rather than in organic solvents, is however significantly less emissive (Figure 1b, and Figure S3, Supporting Information).⁵⁶ The ECL reaction is triggered by electrode-generated superoxide (Figure 1a).⁵¹ As superoxide diffuses away from its generation site, the ECL's front also moves away from the platinum–RTIL interface.





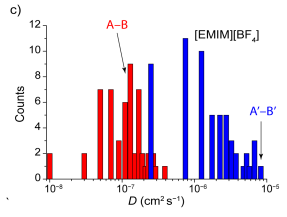
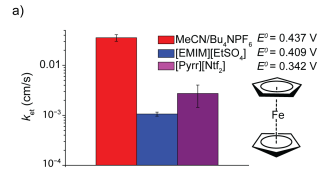
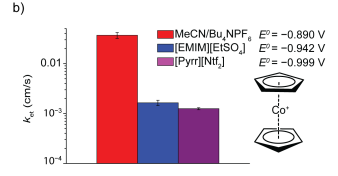


Figure 3. Time-stamped fluorescence micrographs (10× magnification) recorded (a) 0.1 s and (b) 5.1 s after the application of an external bias (-2.0 V vs Ag/AgCl) to an oxygen-saturated AMPluc solution $(0.4 \times 10^{-3} \text{ M in [EMIM][BF}_4], 2400 \text{ ppm of water,}$ Video S3, Supporting Information). Scale bars are 100 um. The movement of the fluorescent front was tracked at ~60 electrode's locations close to sites where gas bubble are visible (e.g. along the A'-B' line), and at ~60 locations at least 100 μm away from bubbles (e.g. A-B line). Histogram plot for the optically determined superoxide D values in [EMIM][BF4]. Blue bars indicate measurements taken at electrode sites with clear convective (bubbles growing) disturbances, and red bars for measurements at quiescent sites.

But while on one hand in DMSO/Bu₄NClO₄ it is possible to track the movement of this front (Figure S4 and Video S4, Supporting Information), and therefore to estimate optically a diffusion coefficient for superoxide $(2.62 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1} \text{ in agree-}$ ment with previous reports),⁵¹ on the other hand in RTILs the AMP-luc ECL reaction is too dim. Consequently, tracking optically the movement of the ECL front does not constitute a viable means of estimating diffusivities in RTILs. However, and very fortunately, the ECL product, **ox-luc** (Figure 1d), is fluorescent. Excitation with blue light of AMP-luc solutions undergoing electrolysis results in a bright green fluorescence which can be imaged with a CMOS camera and selecting exposure times as low as 20 ms (Figure 2a). Access to such fast capture rates, by focusing on redox pro-fluorescence rather than on ECL, means that the superoxide diffusion front can be visually mapped even in highly viscous RTILs (Figure 2a-c). Spatiotemporal D information become therefore available, data which are not accessible through conventional hydrodynamic measurements of diffusivity, such as rotating disk electrodes.⁵⁷ For example, from microscopy data as in Figure 2a-c the evolution of the fluorescence front can be rapidly sampled at several hundred different locations across the platinum electrode. The plot in Figure 2d shows the movement of the front at one of the \sim 200 locations analyzed (marked as A–B). The distance (r) travelled over time (t) by the front (taken arbitrarily as the point with an intensity half of the maximum value) along the A-B line away from the electrode, was used to compute diffusivity assuming an Einstein's random walk ($r^2 = 2Dt$). Data at electrolysis times beyond ~3 s were discarded since, as shown by Amatore and co-workers, natural convection becomes then dominant leading to underestimate the diffuse layer thickness.⁵⁸⁻⁶⁰ The histogram in Figure 2e shows the distribution across the macroscopic sample of the optically determined D. The mode of D is 4.80×10^{-7} cm² s⁻¹, but while for example the fluorescent front moves along the A–B line with a D of 5.25×10^{-7} cm² s⁻¹, the same measurement along A'-B' indicates a slightly smaller $D (3.35 \times 10^{-7})$ cm 2 s $^{-1}$). Such local differences in D, most likely due to natural convection, 58-60 would however be lost in a conventional electrochemical measurement and even advanced forms of redox microscopy would not be able to access these information.⁴⁵⁻⁴⁷ The average D obtained in [EMIM][EtSO₄] is one order of magnitude smaller than in DMSO,51 as it expected based on the higher viscosity of the former (~100 cP vs. 2.0 cP).⁶¹

cathodic the electrolysis of AMP-luc [EMIM][EtSO₄] the interface remains relatively quiescent and the distribution of D across the electrode surface is therefore relatively narrow (Figure 2e). This is surprising, as despite both the large negative potential used to trigger the pro-fluorescent reaction (-2.0 V vs Ag/AgC for 15 s) and the moderate level of water content in [EMIM][EtSO₄] (930 ppm), yet there was no evidence of hydogen bubbles forming on the electrode (Video S2). The formation and departure from the electrode of gas bubbles would have introduced convection hence caused highly dispersed D values. This leads to the question of what prevents in [EMIM][EtSO₄] gas evolution at such negative voltage biases. As introduced above, some RTILs form compact and longlived cation-rich double layer structures on negatively biased electrodes. The compactness of such RTILs structures has been detected by Atkin and co-workers as a mechanical resistance to AFM tips approaching biased electrodes, 16,62





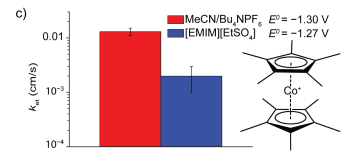


Figure 4. Comparison of the refined (digital simulations of experimental cyclic voltammograms, CVs) charge transfer rate constant ($k_{\rm et}$) of three outer-sphere redox molecules (Fc (a), Cc⁺ (b), and Me₁₀Cc⁺ (c)). Voltammograms were obtained in 1.0×10^{-1} M MeCN/Bu₄NPF₆ (red bars), [EMIM][EtSO₄] (blue bars), and [BMPyrr][NTf₂] (violet bars). Experiments were repeated at least five times for each system, and the CV voltage sweep rate was varied between 0.01 and 5.0 Vs⁻¹. (Supporting Information, Figure S8–S15). The metallocene concentration was 1.0×10^{-3} M. No $k_{\rm et}$ value is reported for Me₁₀Cc⁺ in [BMPyrr][NTf₂] as CVs of this system were poorly reproducible. Refined values of the apparent formal potential (E^0 vs Ag/AgCl) are indicated by labels in figure. The chemical structure of each metallocene is shown next to the relative graph.

and manifest stable and discrete open circuit potential (OCP) plateaus.¹³ For example, OCP-time data in Figure 2f show that with [EMIM][EtSO₄], the imidazolium-rich layer formed at a platinum cathode persists for nearly 10 h once the cathodic bias has been removed. Only mechanical shaking of the electrochemical cell disrupts prematurely such ordered double-layer arrangement.¹³ More specifically, once the exogenous field (the negative electrode bias) is removed and the OCP logging

started, both crowding and overscreening structures persist for ~15 min and ~10 h, respectively (Figure 2f). 11, 13 Notably, in the fluorescence experiments described above the cathodic bias is not interrupted, hence cation-rich interfacial layers are likely to be present for the entire duration of the AMP-luc electrolysis. We believe this compact layer is introducing a kinetic barrier to inner-sphere redox reactions, preventing, despite the presence of substantial levels of water, gas evolution even under very negative voltages. The formation of crowding and overscreening structures in [EMIM][EtSO₄] hinders the reduction of water traces to the point that this relatively hydrophilic RTIL (~1000 ppm of water) behaves alike an hydrophobic RTIL such as [BMPyrr][NTf₂] (~100 ppm of water, Figure S5 and Video S5, Supporting Information). For both RTILs, the lack of bubbleinduced convective disturbances accounts for a relatively narrow distribution of D values across the macroscopic electrode.

By mapping the **ox-luc** fluorescent front, a large dispersion in the diffusivity characteristics were however revealed for RTILs that do not form compact double-layers. For instance, in [EMIM][BF₄] (34 cP, ~2400 ppm of water) mapping diffusion fronts at random locations across the electrode unveiled superoxide D values spread over nearly four orders of magnitude (Figure 3). [EMIM][BF₄] does not have a significant barrier to water reduction (Figure S6, Supporting Information) and hydrogen bubbles form on the electrode during the cathodic electrolysis of **AMP-luc**. Notably, when *D* is measured far away from a growing bubble, such as at the A-B line marked in Figure 3ab, its value is, as expected, lower than for the less viscous [EMIM][EtSO₄] and experimental D values felt within a relatively narrow range (Figure 3c, red bars). Growing bubbles are clearly visible in the fluorescence micrographs (Figure 3b), and tracking the movement of **ox-luc** near the gas cavities – sites where quiescent diffusion is disturbed by convection – led to a significant local overestimate of D (Figure 3c, blue bars).

This spread in the superoxide diffusion coefficient in [EMIM][BF₄] is caused by the lack of a barrier towards innersphere reactions and not by the large hydrophilicity of this RTIL. For instance [EMIM][EtSO₄] does not show any evolution of bubbles even when deliberately spiked with 2% of water (Figure S7a and Video S6, Supporting Information). Even with such large level of water, the [EMIM][EtSO₄]-electrode interface retains its ability to form compact electrostatic structures detectable as long-lived negative OCP plateaus (Figure S7b). [EMIM][BF₄] on the other hand cannot favourably align with the electrode electric field even at water levels as low as 500 ppm.¹³ The formation of stable double layers is strongly influenced by the nature of the RTIL, and for example cations with smaller alkyl chains can more easily pack at the a negative charged surface compared to larger cations.⁶³ However, [EMIM][EtSO₄] and [EMIM][BF₄] share the same cation, thus the strikingly different ability to form stable double layers is due to difference in the anion. Specifically, the reason lies in the [BF₄] anion lacking a dipole moment. The stronger dipole moment of [EtSO₄] facilitates its alignment with the external electric field, and consequently the generation of an internal electric filed, which manifests as a long-lived OCP signature.¹³

It is only when the water level becomes as high as 5.0% that the duration of overscrening plateaus in [EMIM][EtSO₄] is significantly reduced, while the evolution of gas bubbles becomes

significant (Figure S7c,d and Video S7, Supporting Information). With [BMPyrr][NTf₂], another RTIL where cathodic biasing does not lead to discrete negative OCP signatures (Figure S5f), a water content of 2.0% is already sufficient to cause significant gas evolution under the negative bias of the pro-fluorescent experiments (Figure S7e,f and Video S8, Supporting Information). While, as shown above, an inner-sphere reaction such as hydrogen evolution is influenced by how accessible the electrode surface is, the presence or absence of ordered and compact RTIL structures is unlikely to have a significant effect on outer-sphere reactions. 18 To test this we turned to well-studied outer-sphere metallocenes.64 Crowding and overscreening are unlikely to impair the redox kinetics of molecules such as ferrocenes and cobaltocenes. The choice of Cc⁺ and Me₁₀Cc⁺ is because they both have, especially Me₁₀Cc⁺, 65, 66 redox potentials sufficiently negative so to approach the cathodic bias used for the AMP-luc pro-fluorescent reaction and to closely match the position of the overscreening and crowding OCP signatures. By digital simulations of cyclic voltammetry experiments at different scan rates (from 1.0×10^{-2} to 5.0 V s^{-1}) we looked for evidence of a drop in the electron transfer rate constants ($k_{\rm et}$) when the solvent was changed from MeCN to RTILs of similar viscosity (~100 cP) but that either form or do not form cationrich structures that leads to discrete OCP signatures after a cathodic pulse ([EMIM][EtSO₄] and [BMPyrr][NTf₂] respectively).⁶⁷ The fitting results are summarized in Figure 4. Firstly, in an organic solvent (MeCN/Bu₄NPF₆) the refined $k_{\rm et}$ is about one order of magnitude higher than in both RTILs. Secondly, a comparison between k_{et} obtained in [EMIM][EtSO₄] and in [BMPyrr][NTf₂] suggests for both Fc and Cc comparable, with the uncertainty, redox kinetics in both RTILs. Unfortunately data for Me₁₀Cc in [BMPyrr][NTf₂] were not reproducible, possibly due to the poor stabilization of Me₁₀Cc⁺ by the [NTf₂] anion.⁶⁸ The drop in k_{et} moving from MeCN to [EMIM][EtSO₄] was comparable for Fc, Cc⁺ and Me₁₀Cc⁺. This is important, since in the case of a blocking effect by cathodic overscreening and crowding structures on the electron transfer of outer-sphere couples, a drop in $k_{\rm et}$ would have been negligible for the relatively anodic Fc/Fc⁺ couple, but progressively larger for Cc⁺ and Me₁₀Cc⁺. The redox couple Cc/Cc⁺, has a redox potential 1.33-1.35 V more negative than that of the Fc/Fc⁺ couple (Figure 4a,b), and the Me₁₀Cc/Me₁₀Cc⁺ couple is about 1.67 V more negative than the Fc/Fc⁺ couple, and therefore close to the potential where ordered double layers form in [EMIM][EtSO₄] (Figure 2f) and yet the relative drop in $k_{\rm et}$ is comparable or even smaller for the latter.

CONCLUSIONS

We have developed a spatiotemporally resolved optical measurement of diffusion coefficients that can account for convective disturbances. This approach reduces the likelihood of overestimating diffusion coefficients in viscous systems where adventitious gas-evolving reactions cannot be ruled out. The cathodic generation of fluorescent oxyluciferin (ox-luc), by means of reacting firefly's luciferin adenylate ester (AMP-luc) with electrode-generated superoxide radical anion, was used to map superoxide diffusivity in RTILs. Literature diffusivity data for superoxide in RTILs are highly scattered, presumably because the presence of water impurities^{1, 69, 70} leads to unaccounted and poorly reproducible convective contribution to

mass transport. Visual mapping of electro-generated **ox-luc** diffusion fronts addresses this this issue and allows to measure accurately diffusion coefficients even in systems where convection cannot be removed. For example we demonstrate that in [EMIM][BF₄], D for superoxide increases from 6.40×10^{-8} cm² s^{-1} in quiescent regions, to 2.30×10^{-6} cm² s⁻¹ in regions affected by convective disturbances (in proximity of evolving gas bubbles). A conventional macroscopic one-electrode, one-lead measurement would output an average and therefore overestimate diffusion coefficient. While evolution of gaseous products, and its interference with quiescent diffusion, cannot be removed, it can however be detected, accounted for, and even predicted. With regards to this last point – the likelihood of significant convective disturbances – we have been able to uncover a link between inner-sphere reduction of water and the stability of cation-rich double layer RTIL structures. Our results suggest that RTILs forming stable overscreening and crowding double layers can be biased to very negative voltages without detectable gas evolution from water splitting or proton reduction. Readily available time-resolved open-circuit potentiometry can be used to screen for RTILs likely to yield stable overscreening and crowding double layers under negative electrode biases. We also note that while compact double layer RTIL structures pose a significant kinetic barrier to inner sphere reactions, such structures does not impair the electrode kinetics of outer-sphere redox reactions, such as for widespread metallocenes.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website. Detailed description of electrochemical set up, experimental and simulated cyclic voltammograms, open circuit potentiometry data, fluorescence and ECL images, time resolved fluorescence and ECL intensity profiles, photon counting plots (PDF). Supplementary videos (.mp4).

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