**Influence of the acid–base stoichiometry and residual water on the transport mechanism in a highly-Brønsted-acidic proton-conducting ionic liquid**

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In this study, Brønsted-acidic proton conducting ionic liquids are considered as potential new electrolytes for polymer membrane fuel cells with operating temperatures above 100 °C. N-Methyltaurine and trifluoromethanesulfonic acid (TfOH) were mixed at various stoichiometric ratios in order to investigate the influence of an acid or base excess. The proton conductivity and self-diffusion of the ‘neat’ and with 6 wt% water samples were investigated by following electrochemical and NMR methods. The composition change in the complete species and the relative proton transport mechanism based on the NMR results are discussed in detail. During fuel cell operation, the presence of significant amounts of residual water is unavoidable. In PEFC electrolytes, the predominating proton transfer process depends on the cooperative mechanism, when PILs are fixed on the polymer matrix within the membrane. Due to the comparable acidity of the cation [2-Sema]⁺ and the hydroxonium cation, with excess N-methyltaurine or H₂O in the compositions, fast proton exchange reactions between the protonated [2-Sema]⁺ cation, N-methyltaurine and H₂O can be envisaged. Thus, an increasing ratio of cooperative proton transport could be observed. Therefore, for polymer membrane fuel cells operating at elevated temperatures, the highly acidic PILs with excess bases are promising candidates for future use as electrolytes.

**Introduction**

Polymer electrolyte fuel cells (PEFCs) operating at elevated temperatures (>100 °C) offer significant improvements over low-temperature PEFCs, such as no humidification of the feed gas, no water recirculation, a more efficient cooling of the cell and a higher tolerance against feed gas impurities.¹⁻³ The proton conductivity of NAFION®-based proton exchange membranes (PEMs), used in PEFCs for low operation temperatures, depends mainly on the polymer’s water uptake. For operation at elevated temperatures (>100 °C), the conductivity of a new membrane material should be maintained in anhydrous conditions. Currently, (high temperature-) HT-PEFCs are based on polycarbazimazolazo (PBI) membranes doped with phosphoric acid (H₃PO₄).⁴⁻⁷ However, the presence of H₃PO₄ leads to slow cathodic oxygen reduction reaction (ORR) kinetics.⁸ There is a specific adsorption of H₃PO₄ species on active sites in the redox catalyst platinum, which causes an inhibition (poisoning) effect.⁹,¹⁰ In addition, the insufficient solubility and diffusivity of oxygen is discussed.¹¹ Thus, there is a need for new non-aqueous proton-conducting electrolytes to be operational for temperatures of 100–120 °C.

Proton-conducting ionic liquids (PILs) are promising candidates as non-aqueous electrolytes at operating temperatures >100 °C. Ionic liquids (ILs) are ionic compounds with bulky cations and anions and thus a low lattice energy.¹² PILs have received much attention as a potential electrolyte in PEFCs due to their good conductivity, wide electrochemical windows and low flammability.¹³⁻¹⁸ In a PIL, the cation or anion may act as a protonic charge carrier, and so either the cations or anions are Brønsted-acids. In the case of cations, i.e., a PIL of the type HB’A⁻, it consists of an (organic) base B, protonated by a very strong acid HA, respectively a super acid:

\[
B + HA ⇌ HB’ + A^–
\]  

The anions of super acids, such as trifluoromethanesulfonic acid or bis-trifluoromethanesulfonimid, have a less inhibiting effect on electrocatalytically-active electrode surfaces than H₃PO₄. The triflimid (CF₃SO₂)₂N⁻ and triflate CF₃SO₃⁻ anions interact only very weakly with metal atoms, resulting in weak adsorption on a Pt surface.¹⁹,²⁰ In a water-free PIL of the type HB’A⁻, protons can only move in an electric field via the
protonated cations HB\(^+\) by means of a vehicle mechanism. A drawback of ILs or PILs is often poor conductivity because of relatively high viscosity. A proton transfer back to the anion of the superacid, e.g., [TfO\(^-\)], has only a very small probability because the protolysis equilibrium in eqn (1) is on the far right side. Thus, a cooperative transport mechanism involving the anions is not possible. In the case of vehicular mechanism, conductivity and viscosity are coupled to each other according to the Stokes–Einstein relation.

However, to avoid the leakage of the liquid electrolyte during operation, a PIL applied in a PEFC must be immobilized in a polymer matrix. A study of a [Dema][TfO]-doped PBI membrane by Liu et al. shows an activation energy of the conductivity in the range of the cooperative mechanism,\(^ {21}\) whereas the pure [Dema][TfO] is vehicular.\(^ {22}\) In this case, the vehicular transport of the cation HB\(^+\) is constrained and a cooperative proton transport mechanism would be advantageous. This was shown in a study by Noda et al. that the excess in the base B or impurities in the PIL of the type HB\(^-\)A\(^+\), which provides the cooperative proton transport and improves conductivity.\(^ {23}\) The excess base B acts as a proton acceptor that is protonated by the proton donor HB\(^-\). The cooperative transports through the excess base B only necessitate the reorientations of the involved particles B an HB\(^-\). This results in increased proton conductivity and reduced activation energy for the conduction process.\(^ {12,23,24}\)

In particular, PILs contain strong Brønsted-acidic cations that are usually highly hygroscopic. The water absorption is difficult to prevent. Moreover, under fuel cell operation, water will be generated on the cathode side. The presence of residual water acts as a proton acceptor and gives rise to a protolysis equilibrium with the cation HB\(^+\):

\[
\text{HB}^- + \text{H}_2\text{O} \rightleftharpoons \text{B} + \text{H}_3\text{O}^+ \tag{2}
\]

Its extent depends on the acidity of the cation. In a preceding NMR study, it was shown that in Brønsted-acidic PILs of the type HB\(^-\)A\(^+\) cooperative proton transfer will dominate, depending on the cation acidity and the residual water content.\(^ {25}\) There will also be fast exchange between HB\(^-\), B, H\(_2\)O and H\(_3\)O\(^+\). In general, as discussed above, the introduction of a proton acceptor would improve the cooperative transport. The coexistence of excess base B and residual water offers the possibility to improve the technically utilizable conductivity of PIL electrolytes.

In this experimental study, the effect of the PIL acid–base stoichiometry on the proton transport mechanism in a system with residual water is investigated. In general, cooperative proton transport in an IL system requires the presence of a proton acceptor and proton donor with comparable acidity.\(^ {26}\) A highly acidic PIL, 2-sulfoethylmethylammonium trflate [2-Sema][TfO] (pK\(_{A1}\) ≈ -1),\(^ {27}\) is used. The acidity of this is comparable to the hydroxonium cation (pK\(_{A}\) = 0). The 2-sulfoethylmethylammonium cation is prepared by means of the protonation of 2-methylaminoethanesulfonic acid (N-methyltaurine), which exists as a zwitterion due to tautomerism. Because of the presence of the sulfonic acid functionality, it is a very strong acid that can protonate the residual water at a significant percentage.\(^8\)

Appropriate amounts of N-methyltaurine (MTau) and trifluoromethanesulfonic acid (TfOH) are mixed at various molar ratios to vary the PIL compositions from the TfOH-excess to MTau-excess. The interactions between the cations, the excess base and H\(_2\)O are determined by means of electrical conductivity measurements and \(^1\)H NMR spectroscopy. Using a pulsed-field gradient (PFG-) NMR technique, the self-diffusion coefficients of the individual protons in the PILs are obtained. The effect of stoichiometry and residual \(\text{H}_2\text{O}\) on the prevailing proton transport mechanism\(^ {26}\) is discussed by comparing the measured macroscopic and microscopic properties.

### Experimental

**Materials**

Equimolar 2-sulfoethylmethylammonium trflate [2-Sema][TfO] is prepared by slowly adding trifluoromethanesulfonic acid (triflic acid, reagent grade: 98%, Sigma Aldrich) to 2-methylaminoethanesulfonic acid (N-methyltaurine, ≥99%, Sigma Life Science). In the following, N-methyltaurine is abbreviated as MTau in analogy to triflic acid (TfOH). The total amount of [2-Sema][TfO] is divided into 5 bottles, each of about 5 g. Appropriate amounts of TfOH and N-methyltaurine are added to each bottle to maintain the defined molar ratios \(x, i.e., x\text{[MTau]}(1 - x)\text{[TfOH]}\). The mole fraction of N-methyltaurine was from \(x = 0.3–0.7\). Thus, after mixing and heating samples containing 6 wt% \(\text{H}_2\text{O}\), the samples are obtained.

Using Karl-Fischer titration, it can be observed that the samples prepared from the starting materials have a water content of 0.75–0.8 wt%. To analyze the water content, two series of samples were investigated. The first series were the as-prepared [neat] \(x\text{[MTau]}(1 - x)\text{[TfOH]}\) samples with a residual water content of 0.75–0.8 wt% and the second series were samples diluted with water to an \(\text{H}_2\text{O}\) content of 6 wt%, i.e., \(x\text{[MTau]}(1 - x)\text{[TfOH]} + 6\text{ wt% H}_2\text{O}\). The (molar) ratio of \(x\text{[MTau]}(1 - x)\text{[TfOH]}\) : \(\text{H}_2\text{O}\) is in the range of 0.98–1.0, due to the similar molar mass between \(M_{\text{MTau}} = 139\) and \(M_{\text{TfOH}} = 150\). The different stoichiometries of the samples are illustrated in Fig. 1. The samples with the composition \(x = 0.3–0.5\) have a “TfOH-excess”. Conversely, the samples with the composition \(x = 0.5–0.7\) (or 0.65) have an “MTau-excess”.

**Conductivity measurement**

The AC conductivity measurements were performed in a four-probe conductivity cell, using platinum electrodes. The cell constant as a function of the sample volume was determined by using a 0.1 M KCl solution for calibration. The intended water contents of the binary PIL + \(\text{H}_2\text{O}\) mixtures were checked using Karl-Fischer titration at the beginning of each measurement. The total ohmic resistance \(\sigma\) as a function of the temperature \(T\) of the neat PIL and of the PIL + \(\text{H}_2\text{O}\) samples was determined by means of impedance spectroscopy. The temperature \(T\) was increased in increments of 10 \(^\circ\)C from 60 to 110 \(^\circ\)C and vice versa. The excitation amplitude was adjusted to 10 mV. The specific conductivity \(\sigma\) was calculated by using the cell constant.

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**1H NMR parameters**

The acquisition of the NMR spectra was performed using a Bruker 600 MHz spectrometer, equipped with a 5 mm cryo-probe tuned to 1H. A capillary filled with D2O was enclosed the sample tubes as a field lock. The measurements were performed at 90 °C, because at lower temperatures, the increasing viscosity leads to high relaxation times.

**Measurement of the self-diffusion coefficients**

The self-diffusion coefficients of the observable protons were measured using the diffusion-ordered spectroscopy (DOSY) technique at 90 °C. The measurements were performed by applying 30 field gradient increments with a gradient strength g from 1.3 to 32.5 G cm⁻¹. The values of the gradient pulse length δ and the diffusion time intervals Δ were optimized to aim for least 85% signal attenuation at the strongest field gradient. The value of the (self-) diffusion coefficient Di of a certain proton species i was obtained from the decay of its measured echo intensity vs. the gradient field strength g.

**Results and discussion**

**Total conductivity vs. stoichiometry and water content**

The measurements of the total conductivity σ were performed in the temperature range between 60 and 110 °C. The total conductivity includes cationic and anionic charge transport. In the x[M Tau]·(1−x)[TFOH] and the x[M Tau]·(1−x)[TFOH] + 6 wt% H2O samples, the viscosity was strongly dependent on the composition. In general, the viscosity was rising with an increasing fraction x of the (at room temperature solid) base N-methyltaurine, i.e., with increasing MTau-excess. In the case of an increasing TFOH-excess, the viscosity was generally decreasing, i.e., with decreasing x. A higher content of H2O also leads to a lower viscosity. The dependence of the total conductivity σ, respectively of the specific total conductivity λ on the viscosity η, can be explained according to the Stokes–Einstein and Nernst–Planck relations:

\[
\lambda \eta = \frac{e^2 N_A}{6\pi} \sum_{i} \frac{z_i^2 r_i^4 \alpha_{\text{Diss}}}{\epsilon_{\text{Diss}}} \ \text{const. with } \alpha_{\text{Diss}} = \frac{\epsilon_i}{\epsilon_0} \quad \text{and} \ \lambda = \frac{\sigma}{C_0} 
\]

(3)

Assuming an ionic compound A_i B_j C_k ... X_m, \( r_i \) denotes the stoichiometric factor, \( z_i \) the charge number and \( r_i \) the hydrodynamic radius of the ionic species i. In the case of a dissociation degree \( \alpha_{\text{Diss}} \) not being equal to unity, the concentration of the ionic species i is denoted with \( c_i \) and the initial concentration of the ionic compound \( A_i B_j C_k ... X_m \) with \( \epsilon_0 \). Thus, a decrease in the viscosity will accelerate the vehicular proton transport by PIL cations and H3O⁺.

The dependency of the total conductivity \( \sigma \) on the neat samples and samples with a water content of 6 wt% on the temperature \( T \) and the stoichiometry \( x \) is depicted in Fig. 2(a) and (b), respectively. The course of the conductivities corresponds to the change in the viscosity. In the case of the neat samples, depicted in Fig. 2(a), the sample with the smallest MTau molar fraction \( x = 0.3 \) exhibits the highest total conductivity. Correspondingly, the total conductivities of the samples with a water content of 6 wt% are generally higher compared to the neat sample, as is depicted in Fig. 2(b).

For all of the investigated samples, the total conductivity \( \sigma \) increases as a function of the temperature \( T \). However, the extent of the conductivity increase vs. temperature is different. The highest impact on the conductivity was found for the sample with the highest MTau molar fraction of \( x = 0.7 \). For this MTau molar fraction, in the case of the neat samples, \( \sigma \) increases by a factor of 8.4 when \( T \) rises from 60 °C to 110 °C. In the case of the sample with a molar fraction of \( x = 0.3 \), the increase was only by a factor of 3. The samples with a water content of 6 wt% and a molar fraction of \( x = 0.7 \) and \( x = 0.3 \) show a similar behaviour, with the conductivity increasing by a factor of 8.52 and 2.75, respectively. The different factors may

![Fig. 1](image_url)
indicate that there is not only a change in viscosity but also of the proton transport mechanism responsible for the increase in the total conductivity when increasing the molar fraction of MTau. The total conductivity describes the bulk charge transport macroscopically and includes all mobile charge carriers ([2-Sema]$^+$, [TfO]$^-$ and H$_3$O$^+$). The underlying ionic charge (proton) transport mechanisms can only be discerned by techniques sensitive to the local environment of the mobile charge carriers, i.e., the NMR.

**$^1$H-NMR of neat x[MTau]·(1 − x)[TfOH] samples**

In the following, $^1$H-NMR and $^1$H-PFG-NMR are used to measure the local dynamics of the mobile protonic charge carriers, as well as their self-diffusion coefficients. As discussed above, in a PIL of the type HB$^+$A$^-$, an excess of the base B as well as a certain water content are able to provide cooperative transport and thus improve the proton conductivity. Therefore, depending on the excess of the base and water content, both transport mechanisms, vehicular and cooperative, may be present in a sample.

Due to the high viscosity of the samples at room temperature, especially in the case of MTau-excess, the NMR measurements were all performed at 90 °C to avoid an FWHM of the signals too broad to evaluate. The $^1$H NMR spectra of x[MTau]·(1 − x)[TfOH] (x = 0.3–0.7) at a temperature of 90 °C are depicted in Fig. 3(a) and the chemical shift δ of the protons situated in the SO$_3$H and NH$_3^+$ group vs. the stoichiometry x in Fig. 3(b). The signals of the protons are labelled with (a), (b), (c, c’), and (d), according to Fig. 1. In the case of the stoichiometric [2-Sema][TfO] sample, i.e., x = 0.5, the protons of the NH$_3^+$ group (b) show up at a chemical shift δ of 6.83 ppm and the proton of the SO$_3$H group (a) at 12.25 ppm. When increasing the mole fraction x of the base MTau from 0.3 to 0.7, the signal (b) of the NH$_3^+$ protons shifts about 0.6 ppm towards the lower magnetic field, i.e., from a chemical shift of 6.49 to 7.10 ppm. In the case of SO$_3$H protons (a) there is a slight shift of +0.06 ppm towards lower magnetic fields when increasing x from 0.3 to 0.4 and a slight shift of −0.17 ppm towards higher magnetic fields from 0.4 to 0.5. Beyond the stoichiometric composition, for a molar fraction x of 0.5 to 0.7, the signal shifts about +0.9 ppm towards lower magnetic fields with increasing the base excess. The signal of the protons of the CH$_3$ group (d) show up at a chemical shift δ of 2.63 ppm and of the CH$_2$CH$_2$ backbone (c, c’) at about 3.37 ppm. There is no appreciable shift in the CH$_3$
(d) and the CH₂CH₂ (c, c’) protons. In the case of samples with a base-excess, i.e., $x > 0.5$, the protons of the CH₂CH₂ backbone (c, c’) are difficult to distinguish due to a general increase of the FWHM in the spectra.

The NMR chemical shift $\delta$ of a proton depends on the local screening of the external magnetic field by the local electron density. In the timescale of the NMR experiment, a chemical shift to a lower field represents a generally lower local electron density or higher “delocalization” of the proton. The local electron density can be affected by intramolecular interactions with adjacent groups or intermolecular interactions by hydrogen bonds. A high acidic proton is accompanied by a low local electron density. In addition, the formation of a hydrogen bond leads to the deshielding of a proton. Thus, the (NMR-) active protons of a Bronsted-acidic PIL in particular are influenced not only by the water content but also by the acid–base stoichiometry, as the active protons can easily form hydrogen bonds and are subject to intramolecular interactions. In the case of the water-free samples, the molar fractions of the species [2-Sema]$^+$, MTau, TFOH and [TFO]$^-$ should vary as a function of the stoichiometry $x$, as depicted in Fig. 4. TFOH is a much stronger acid compared to the [2-Sema]$^+$ cation, and so we can safely assume a complete proton transfer from TFOH to MTau for the entire range of stoichiometry $x$. This should lead to a decreasing fraction of TFOH in the stoichiometric range from $x = 0$ to 1/2 and an increasing fraction of MTau in the range from $x = 1/2$ to 1.

As the acidity of the NH₂⁺ moiety in the [2-Sema]$^+$ cation and MTau is not sufficient to protonate SO₃⁻ moieties (intra/intermolecular) to a noticeable extent, the residence time prior to re-transfer should be very short. Thus, protolysis equilibria should not contribute to the observed stoichiometry $x$-dependent shift of these protons ($pK_A$ of NH₂⁺ in MTau, about 10.2).27–29 As is shown in Fig. 4, the sum of the molar fractions of [MTau] and [TFO]$^-$ is increasing when increasing $x$ from 0.3 to 0.7. Thus, in the case of the NH₂⁺ protons, only the probability of forming H-bonds with neighboring [TFO]$^-$ anions and [2-Sema]$^+$ cations and excess MTau molecules, respectively, can be considered to explain the observation that the $\delta$ value of the NH₂⁺ protons shifts monotonically towards lower fields with a slope of 0.15 ppm/0.1 $\Delta x$ (deshielding).

In the case of SO₃H protons, a stronger effect can be observed compared to the NH₂⁺ protons. Despite the fact that SO₃H protons are present on [2-Sema]$^+$ cations and TFOH, there is only one signal shifting from 12.14 to 13.15 ppm to lower fields. The sites cannot be distinguished on the NMR spectrum. In addition to the possibility of forming H-bonds, the possibility/frequency of intermolecular proton transfers must also be considered. In the case of MTau-excess, i.e., for samples with a stoichiometry of $x = 0.5$ to 0.7, there will be only variable fractions of [2-Sema]$^+$, MTau and [TFO]$^-$.

The acidity of TFOH ($pK_A = -14$) is much higher than that of [2-Sema]$^+$ ($pK_A = -1$).27 The proton transfer from the cation [2-Sema]$^+$ back to the [TFO]$^-$ anion will not take place to a noticeable degree. Thus, TFOH is fully deprotonated. There are only fast intermolecular proton transfers between SO₃⁻ moieties of [2-Sema]$^+$ and MTau, resulting in the strong deshielding of the proton with increasing $x$:

$$\text{CH}_3\text{NH}_2^+(\text{CH}_2)_2\text{SO}_3\text{H} + \text{CH}_3\text{NH}_2^+(\text{CH}_2)_2\text{SO}_3^- \rightarrow \text{CH}_3\text{NH}_2^+(\text{CH}_2)_2\text{SO}_3^- + \text{CH}_3\text{NH}_2^+(\text{CH}_2)_2\text{SO}_3\text{H} \quad (4)$$

In the stoichiometry range between $x = 0.5$ to 0.7, the signal shifts with a slope of 0.45 ppm/0.1 $\Delta x$; see Fig. 4.

In the case of TFOH-excess, i.e., for samples with a stoichiometry of $x = 0.3$ to 0.5, the shift in the signal is comparably small and not monotonic. Due to the TFOH excess, the SO₃H moiety of the MTau is fully protonated. There are only variable fractions of [2-Sema]$^+$ cations, TFOH and [TFO]$^-$.

Thus, there is primarily an intermolecular proton transfer between the anion [TFO]$^-$ and TFOH, which also results in a de-shielding of the proton:

$$\text{CF}_3\text{SO}_3\text{H} + \text{CF}_3\text{SO}_3^- \rightarrow \text{CF}_3\text{SO}_3^- + \text{CF}_3\text{SO}_3\text{H} \quad (5)$$

Changing the stoichiometry from $x = 0.5$ to 0.4 leads to an increasing delocalization of the SO₃H proton and to an initial downfield shift, as seen in the NMR chemical shift in Fig. 3(b). In the system TFOH/[TFO]$^-$, the presence of [TFO]$^-$ should be equivalent to the excess base MTau in the system [2-Sema]$^+$/MTau. Thus, a further increase in the molar fraction of TFOH may inhibit the delocalization again, resulting in the observed shift of the SO₃H proton back towards the higher field when changing the stoichiometry from $x = 0.4$ to 0.3, due to the restricted mobility.

In the case of the neat, nearly water-free samples, a change in the stoichiometry will affect the probability and duration of forming hydrogen bonds to the NH₂ proton on cations in the timescale of the NMR measurement. In the case of the SO₃H proton, the MTau-excess leads to a significant delocalization between the SO₃⁻/SO₃H sites in the timescale of the NMR measurement and TFOH-excess results to a delocalization between the [TFO]$^-$/TFOH. Both of these may explain the observed signal shift to high magnetic fields in the $\text{^1H}$ NMR spectrum. The acid-base stoichiometry also influences the proton transport processes. The observation of a proton delocalisation on the NMR timescale indicates the possible presence
TFOH is the much stronger acid compared to the [2-Sema]⁺ cation and the acidity of the hydroxonium cation H₂O⁺ is on the same order as the acidity of the [2-Sema]⁺ cation. A preceding work confirms the fast exchange of the H₂O proton and SO₃H proton, resulting in a single signal in the spectrum of SO₃H/H₂O/H₂O at an average NMR shift. Moreover, the signal of the SO₃H proton shows a corresponding increase in its integral area with increasing H₂O content, which is not the case for the NH₂⁺ or alkyl protons.

Considering the protolysis reactions in eqn (6) and (7), below a stoichiometry x of 0.24, all existing MTau and H₂O will be protonated by the excess TFOH in the x[MTau] · (1 − x)[TFOH] + 6 wt% H₂O samples. Part of the excess TFOH should be undissociated. There will only be [2-Sema]⁺, H₂O⁺, TFOH and [TFO]⁻ in the samples. At a stoichiometry x above 0.24, unprotonated MTau and H₂O should be present, as all TFOH is dissociated, i.e., only [TFO]⁻ should exist in this range. With an increasing fraction of H₂O, a portion of the [2-Sema]⁺ cations can also be deprotonated. Taking the preceding investigations of the analog compound 2-sulfoethylammonium triflate into account, a protolysis degree of the [2-Sema]⁺ cation of about 0.3 can be estimated for a sample with a stoichiometric composition (x = 0.5). These assumptions are underpin the construction of the tentative plot in Fig. 6.

In the case of the investigated x[MTau] · (1 − x)[TFOH] + 6 wt% H₂O samples, the signal of the SO₃H proton shifts in the opposite direction compared to the (neat) x[MTau] · (1 − x)[TFOH] samples, i.e., towards higher fields (from 12.1 to 9.3 ppm). The additional H₂O in x[MTau] · (1 − x)[TFOH] + 6 wt% H₂O samples acts as another proton acceptor. Depending on the stoichiometric composition x, it can be protonated by the very high acidic excess TFOH or by the high acidic [2-Sema]⁺ cation according to the protolysis equilibria shown in eqn (6) and (7):

\[
\begin{align*}
\text{CH₃NH₂⁺}(\text{CH₂})₂\text{SO₃H} + \text{H₂O} & \rightleftharpoons \text{CH₃NH}_{2⁺}(\text{CH₂})₂\text{SO₃}⁻ + \text{H}_3\text{O}⁺ \\
\text{TFOH} + \text{H₂O} & \rightarrow \text{TFO}⁻ + \text{H}_3\text{O}⁺
\end{align*}
\]

(6)

(7)

\( \rightleftharpoons \) balance

\( \equiv \) equilibrium constant

\( \text{SO₃H} / \text{H}_3\text{O}⁺ \) is the water-dependent equilibrium constant in water.

\( \text{pK}_a \) of typical alkane sulfonic acids is about –1.

\( \text{pK}_a \) of H₂O⁺ is equal to 0.

(+) This corresponds to an initial molar fraction of 0.50 for TFOH, 0.16 for MTau and 0.34 for H₂O, considering the molar weights and a water content of 6 wt%.

Fig. 5  (a) The ¹H NMR spectra; (b) the ¹H NMR spectra chemical shift (ppm) of SO₃H and NH₂ of 6 wt% water concentration x[MTau] · (1 − x) [TFOH] + 6 wt% H₂O at 90 °C.

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The fraction of the [2-Sema]$^+$ cations as a proton acceptor will also accelerate the intermolecular transport. For a stoichiometry of 0.3 to 0.65 should lead to a decreasing fraction of H$_3$O$^+$ and an increasing fraction of H$_2$O. The fraction of the [2-Sema]$^+$ cations will reach a maximum in close proximity to the stoichiometric point $x = 1/2$. There is a delocalization of the active proton in the NMR timescale between H$_2$O, H$_3$O$^+$, MTau and [2-Sema]$^+$, because the neutral MTau is competing with H$_2$O for the protons of the [2-Sema]$^+$ and H$_2$O$^+$ cations. The chemical shift found for the H$_2$O protons (~9.6–10.9 ppm) is at a much lower magnetic field compared to H$_2$O (~3–5 ppm). Thus, the averaged ppm value of the SO$_3$H/H$_2$O/H$_3$O$^+$ proton should also decrease, respectively shifting towards a higher field, as it inclines to the ppm value of H$_2$O; see Fig. 5.

The proton transport mechanism is further discussed, together with the $^1$H-PFG-NMR/DOSY measurements and the self-diffusion coefficient in the next section. The mobile protonic charge carriers in the PIL/H$_2$O system are [2-Sema]$^+$ and, due to protolysis, also the H$_2$O$^+$ cations. In the case of vehicular transport, the H$_2$O$^+$ cations are probably much more mobile than the [2-Sema]$^+$ ones. For a stoichiometry $x > 0.3$, the fractions of H$_2$O$^+$ and [TfO]$^-$ monotonously decrease with increasing $x$. The fraction of [2-Sema]$^+$ reaches a maximum at $x \approx \frac{1}{3}$. If there is only vehicular transport, this should principally lead to a decrease in the (total) conductivity with increasing $x$. If there is also cooperative transport, the presence of H$_2$O, acting as a proton acceptor, will also accelerate the intermolecular proton transfer between the MTau and the [2-Sema]$^+$ cation, leading to faster cooperative transport. For a stoichiometry of $x > 0.3$, the fraction of (free) H$_2$O increases with increasing $x$. A maximum of an additional cooperative transport and thus of the (total) conductivity should be expected when the equimolar fractions of H$_2$O and H$_3$O$^+$ are present (the maximum probability for proton transfers between H$_2$O$^+$ and H$_2$O). This is approximately the case for a stoichiometry $x \approx 0.4$ and corresponds well to the measured values for the total conductivity which exhibits a maximum at this stoichiometry; see Fig. 2(b) and 6. However, the total conductivity is highly coupled with the viscosity. At a higher stoichiometric composition $x > 0.4$, the vehicular transport is again attenuated and thus the conductivity is decreased.

### 1H-PFG-NMR/DOSY

The self-diffusion coefficient of the protons is measured by $^1$H-PFG-NMR at 90 °C. The diffusion coefficient $D_{H^+}$ of the (active) SO$_3$H/H$_2$O/H$_3$O$^+$ proton and $D_{cation}$ of the other protons of the [2-Sema]$^+$ cation/MTau are depicted in Table 1 for neat x [MTau]·(1−x)[TfOH] and for x[MTau]·(1−x)[TfOH] + H$_2$O, i.e., with a water content of 6 wt%. The value of the diffusion coefficient for the NH$_3$ protons is nearly the same as for the CH$_3$CH$_2$ and CH$_3$ protons. Thus, these can be identified as the diffusion coefficient $D_{cation}$ of the entire cation, respectively, to the diffusion coefficient $D_{H^+, vehicle}$ of protons only by means of a vehicular mechanism. The diffusion coefficient $D_{H^+, vehicle}$ of the SO$_3$H/H$_2$O/H$_3$O$^+$ protons is significantly higher compared to the diffusion coefficient $D_{cation}$ of the [2-Sema]$^+$ cation in all of the samples. As discussed, the observed intermolecular protons transfer between the SO$_3$H moieties, H$_2$O and H$_3$O$^+$, indicating the presence of cooperative transport and resulting in higher values of $D_{H^+, vehicle}$ compared to $D_{cation}$.

As observed for the total conductivity $\sigma$, the diffusion coefficient $D_{H^+}$ of the active proton and $D_{cation}$ of the [2-Sema]$^+$ cation are decreasing with increasing stoichiometry $x$ (from 0.3 to 0.65, respectively, to 0.7). The dynamic viscosity $\eta$ of the samples is directly coupled to the diffusion coefficient of the [2-Sema]$^+$ cation $D_{cation}$, due to the Stokes–Einstein relation. In the case of cooperative transport, there is a distinct decoupling from viscous processes. Thus, if vehicular and cooperative mechanisms are both present, an increase in viscosity will increase the impact of cooperative transport to the total proton transport. The share of cooperative processes in all of the samples is evaluated by calculating the ratio between $D_{H^+, vehicle}$ to $D_{H^+}$ as follows:

$$\frac{D_{H^+, vehicle}}{D_{H^+}} = \frac{D_{H^+, vehicle} - D_{H^+, vehicle}^{cation}}{D_{H^+}}$$ (8)

The diffusion coefficient $D_{H^+, vehicle}$ for vehicular transport is identical to that of the [2-Sema]$^+$ cation. The ratio represents the share of cooperative transport in the total proton diffusion process. The share of cooperative transport is depicted in Fig. 7. With increasing stoichiometry $x$, i.e., with an increasing fraction of MTau, the share of cooperative transport is also increasing.

### Table 1: Self-diffusion coefficients of the SO$_3$H/H$_2$O/H$_3$O$^+$ ($\equiv D_{H^+}$) and the cation protons in neat x[MTau]·(1−x)[TfOH] and 6 wt% water concentration x[MTau]·(1−x)[TfOH] + H$_2$O at 90 °C

| MTau molar fraction x | $D_{H^+}$ $\times 10^{-6}$ cm$^2$ s$^{-1}$ in neat samples | $D_{H^+}$ $\times 10^{-6}$ cm$^2$ s$^{-1}$ in samples with 6 wt% H$_2$O |
|----------------------|----------------------------------------------------------|----------------------------------------------------------|
| 0.3                  | 15.4 ± 0.7                                               | 12.2 ± 1.3                                               |
| 0.4                  | 4.63 ± 0.4                                               | 9.35 ± 1.3                                               |
| 0.4                  | 3.00 ± 0.3                                               | 3.15 ± 0.6                                               |
| 0.5                  | 1.31 ± 0.0                                               | 6.00 ± 0.1                                               |
| 0.6                  | 1.13 ± 0.2                                               | 4.95 ± 0.4                                               |
| 0.65                 | —                                                        | 1.05 ± 0.3                                               |
| 0.7                  | 0.75 ± 0.1                                               | 8.14 ± 0.4                                               |
| 0.7                  | 0.34 ± 0.2                                               | 1.51 ± 0.4                                               |
for the neat samples and for samples with a water content of 6 wt%. When comparing the course of both curves, as expected, the presence of the amphother water generally increases the share of the cooperative mechanism. The ratio of $D_{\text{H}^+}^{\text{coop}}$ to $D_{\text{H}^+}$ reaches a value of 82% in a sample with an stoichiometry of $x = 0.65$ and 6 wt% water content. Corresponding to the chemical shift in the $^1$H-NMR spectrum, the ratio relates to the ability of the protons to be delocalized on the NMR timescale and thus to participate in intermolecular transfer. A higher ability of the protons to be delocalized on the NMR timescale exchange reactions between the $[2\text{-Sema}]^+$ cation, $\text{H}_3\text{O}^+$, MTau $\text{H}_3\text{O}^+$, an excess of MTau or $\text{H}_2\text{O}$ will enhance the fast proton transport mechanism. The excess proton acceptors provide more sites for proton hopping, which accelerate the cooperative proton transport mechanism.

**Conclusion**

$N$-Methyltaurine and TFOH were mixed at various molar ratios in order to obtain samples of the proton-conducting liquid $[2\text{-Sema][TfO]}$ with various amounts of excess free acid TFOH or excess free base $N$-methyltaurine. The “nearly neat” samples and the samples with 6 wt% residual water were investigated regarding proton conductivity and self-diffusion using electrochemical and NMR methods. It could be observed that an excess of the free base MTau retards the vehicular proton transport due to an increase in the dynamic viscosity, which leads macroscopically to a lower (total) conductivity. Due to a comparable acidity of the cation $[2\text{-Sema}]^+$ and the hydroxonium cation $\text{H}_3\text{O}^+$, an excess of MTau or $\text{H}_2\text{O}$ will enhance the fast proton exchange reactions between the $[2\text{-Sema}]^+$ cation, $\text{H}_3\text{O}^+$, MTau and $\text{H}_2\text{O}$ and thus will increase the share of cooperative transport.

In an PEFC electrolyte, based on a PIL immobilised in a polymer matrix, the proton conductivity depends on the presence and the share of cooperative transport as vehicular transport is significantly hampered. The use of a base-excess high acidic PIL would allow a higher fraction of cooperative transport and thus a higher proton conductivity. For the future use as conductive electrolytes in PEFCs at elevated operation temperatures (100–120 °C) and atmospheric (non-humidified) operation, a PIL with a high hygroscopicity to retain $\text{H}_2\text{O}$ formed during operation at the cathode and an excess of the base may be favorable. Enabling fast cooperative transport may help in reaching sufficient proton conductivities.

**Conflicts of interest**

There are no conflicts to declare.

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**References**

1. Y. Liu, W. Lehnert, H. Janšen, R. C. Samsun and D. Stolten, *J. Power Sources*, 2016, **311**, 91–102.
2. D. J. Jones and J. Rozière, *J. Membr. Sci.*, 2001, **185**, 41–58.
3. E. Quartarone and P. Mustarelli, *Energy Environ. Sci.*, 2012, **5**, 6436–6444.
4. Q. Li, J. O. Jensen, R. F. Savinell and N. J. Bjerrum, *Prog. Polym. Sci.*, 2009, **34**, 449–477.
5. J. A. Asensio, E. M. Sánchez and P. Gómez-Romero, *Chem. Soc. Rev.*, 2010, **39**, 3210–3239.
6. L. Vilčiauskas, M. E. Tuckerman, G. Bester, S. J. Paddison and K.-D. Kreuer, *Nat. Chem.*, 2012, **4**, 461.
7. C. Korte, F. Conti, J. Wackerl, P. Dams, A. Majerus and W. Lehnert, *J. Appl. Electrochem.*, 2015, **45**, 857–871.
8. K. Wippermann, J. Wackerl, W. Lehnert, B. Huber and C. Korte, *J. Electrochem. Soc.*, 2015, **163**, F25–F37.
9. K. Hsueh, E. Gonzalez and S. Srinivasan, *Electrochim. Acta*, 1983, **28**, 691–697.
10. K. L. Hsueh, E. Gonzalez, S. Srinivasan and D. T. Chin, *J. Electrochem. Soc.*, 1984, **131**, 823–828.
11. B. R. Scharifker, P. Zelenay and J. M. Bockris, *J. Electrochem. Soc.*, 1987, **134**, 2714.
12. A. Noda, M. A. B. H. Susan, K. Kudo, S. Mitsushima, K. Hayamizu and M. Watanabe, *J. Phys. Chem. B*, 2003, **107**, 4024–4033.
13. M. A. B. H. Susan, T. Kaneko, A. Noda and M. Watanabe, *J. Am. Chem. Soc.*, 2005, **127**, 4976–4983.
14. M. M. Mannarino, D. S. Liu, P. T. Hammond and G. C. Rutledge, *ACS Appl. Mater. Interfaces*, 2013, **5**, 8155–8164.
15. D. E. Smith and D. A. Walsh, *Adv. Energy Mater.*, 2019, **9**, 1900744.
16. L. Koók, B. Kaufer, P. Bakonyi, T. Rózsensberszki, I. Rivera, G. Bütrén, K. Béla-Bákó and N. J. Nemestóthy, *J. Membr. Sci.*, 2019, **570**, 215–225.
17. J. Escorihuela, A. García-Bernabe, A. Montero, O. Sahuquillo, E. Gimenez and V. Compan, *Polymers*, 2019, **11**, 732.
18. J. Luo, A. H. Jensen, N. R. Brooks, J. Sniekers, M. Knipper, D. Aili, Q. Li, B. Vanroy, M. Wübbenhorst, F. Yan, L. Van
Meervelt, Z. Shao, J. Fang, Z.-H. Luo, D. E. De Vos, K. Binnemans and J. Fransaer, *Energy Environ. Sci.*, 2015, 8, 1276–1291.

19 P. Zelenay, B. Scharfker, J. M. Bockris and D. Gervasio, *J. Electrochem. Soc.*, 1986, 133, 2262–2267.

20 V. S. Murthi, R. C. Urian and S. Mukerjee, *J. Phys. Chem. B*, 2004, 108, 11011–11023.

21 S. Liu, L. Zhou, P. Wang, F. Zhang, S. Yu, Z. Shao and B. Yi, *ACS Appl. Mater. Interfaces*, 2014, 6, 3195–3200.

22 S.-Y. Lee, A. Ogawa, M. Kanno, H. Nakamoto, T. Yasuda and M. Watanabe, *J. Am. Chem. Soc.*, 2010, 132, 9764–9773.

23 H. Nakamoto, A. Noda, K. Hayamizu, S. Hayashi, H.-o. Hamaguchi and M. Watanabe, *J. Phys. Chem. C*, 2007, 111, 1541–1548.

24 M. L. Hoarfrost, M. Tyagi, R. A. Segalman and J. A. Reimer, *J. Phys. Chem. B*, 2012, 116, 8201–8209.

25 J. Lin, L. Wang, T. Zinkevich, S. Indris, Y. Suo and C. Korte, *Phys. Chem. Chem. Phys.*, 2020, 22, 1145–1153.

26 K. D. Kreuer, A. Rabenau and W. Weppner, *Angew. Chem., Int. Ed. Engl.*, 1982, 21, 208–209.

27 K. Wippermann, J. Giffin and C. Korte, *J. Electrochem. Soc.*, 2018, 165, H263–H270.

28 P. Rimmelin, S. Schwartz and J. Sommer, *Org. Magn. Reson.*, 1981, 16, 160–163.

29 P. Batamack and F. Jacques, *Catal. Lett.*, 1995, 35, 135–142.

30 E. J. King, *J. Am. Chem. Soc.*, 1953, 75, 2204–2209.

31 H. Huang, J. Y. Jin, J. H. Hong, J. S. Kang and W. Lee, *Bull. Korean Chem. Soc.*, 2009, 30, 2827–2829.

32 M. Endres, J. Giffin and C. Korte, *RSC Adv.*, submitted.

33 A. Kitada, S. Takeoka, K. Kintsu, K. Fukami, M. Saimura, T. Nagata, M. Katahira and K. Murase, *J. Electrochem. Soc.*, 2018, 165, H121–H127.