Systematic Study of Quaternary Ammonium Cations for Bromine Sequestering Application in High Energy Density Electrolytes for Hydrogen Bromine Redox Flow Batteries

Michael Küttinger 1,2, Paulette A. Loichet Torres 1, Emeline Meyer 1, Peter Fischer 1,* and Jens Tübke 1,2

Abstract: Bromine complexing agents (BCAs) are used to reduce the vapor pressure of bromine in the aqueous electrolytes of bromine flow batteries. BCAs bind hazardous, volatile bromine by forming a second, heavy liquid fused salt. The properties of BCAs in a strongly acidic bromine electrolyte are largely unexplored. A total of 38 different quaternary ammonium halides are investigated ex situ regarding their properties and applicability in bromine electrolytes as BCAs. The focus is on the development of safe and performant HBr/Br2/H2O electrolytes with a theoretical capacity of 180 Ah L−1 for hydrogen bromine redox flow batteries (H2/Br2-RFB). Stable liquid fused salts, moderate bromine complexation, large conductivities and large redox potentials in the aqueous phase of the electrolytes are investigated in order to determine the most applicable BCA for this kind of electrolyte. A detailed study on the properties of BCA cations in these parameters is provided for the first time, as well as for electrolyte mixtures at different states of charge of the electrolyte. 1-ethylpyridin-1-ium bromide [C2Py]Br is selected from 38 BCAs based on its properties as a BCA that should be focused on for application in electrolytes for H2/Br2-RFB in the future.

Keywords: electrochemistry; bromine; quaternary ammonium salts; sequestration; hydrogen bromine redox flow battery; electrolyte; liquid/liquid phase equilibrium

1. Introduction

Within the last ten years, numerous chemical combinations of negative and positive half-cell chemistries for redox flow battery application have been published. The majority of these compositions have been investigated on laboratory scale [1–3]. One commercialized and noticeable redox flow battery system for stationary energy storage application is the hydrogen bromine redox flow battery (H2/Br2-RFB) [4–6].

One of the challenges attributed to the H2/Br2-RFB application is the high vapor pressure of bromine in the electrolyte solutions commonly used in this system [7]. In this study, 38 cost-effective additives based on quaternary ammonium halide salts are investigated for their applicability as bromine complexing additives (BCA) and their effects on electrolyte properties. The most promising additive is selected for further investigation in a cell. This systematic investigation is carried out for the first time for electrolytes in H2/Br2-RFBs.

1.1. State of the Art

In this flow battery system, the negative half cell resembles the negative half cell of a proton exchange membrane fuel cell anode, but is operated in a reversible mode. Liquid posolytes based on hydrobromic acid (HBr), bromine (Br2) and water (H2O) are
pumped through the positive half cell, while bromide (Br\textsuperscript{−}) is oxidized to Br\textsubscript{2} during charge operation (Equation (1)) and vice versa during discharge operation [4,5,8]. In some research articles, gaseous positive half cells operating with bromine vapor are also discussed [4]. The thermodynamic cell voltage is 1.09 V [9]. Br\textsubscript{2} itself is hardly soluble in water [10–14]. Nevertheless, if bromide ions are present, Br\textsubscript{2} forms polybromides such as tribromide (Br\textsubscript{3}−), pentabromide (Br\textsubscript{5}−) or higher polybromides Br\textsubscript{2n+1}− [13,15,16] following Equation (2):

\begin{equation}
2 \text{Br}^{−} ⇌ \text{Br}_{2} + 2 \text{e}^{−} \tag{1}
\end{equation}

\begin{equation}
\text{Br}^{−} (\text{aq}) + n \text{Br}_{2} (\text{aq}) ⇌ \text{Br}_{2n+1}^{−} (\text{aq}) \tag{2}
\end{equation}

Bromine and bromide form addition bondings by complexation, resulting in the previously mentioned polybromide cations Br\textsubscript{2n+1}− [17]. Complexation allows the maintaining of high concentrations of Br\textsubscript{2} in aqueous bromide solutions. These electrolytes are particularly interesting for the application of Br\textsubscript{2}/Br\textsuperscript{−} electrolytes in RFBs due to their high theoretical energy density of up to 254 Wh L\textsuperscript{−1} and a theoretical volumetric capacity of up to 233 Ah L\textsuperscript{−1} (HBr 48 wt%). Ideally, in order to achieve high power densities in H\textsubscript{2}/Br\textsubscript{2}-RFBs, high concentrations of HBr and Br\textsubscript{2} are indispensable [18]. Nevertheless, Br\textsubscript{2} itself and in polybromide form is already quite volatile at ambient temperature [19–21] and can react with different electrode and cell components of the H\textsubscript{2}/Br\textsubscript{2}-RFBs [4,22–26]. One major concern is the crossover of Br\textsubscript{2} from the positive to the negative half cell during cell operation. When Br\textsubscript{2} comes into contact with the platinum-based catalysts in the hydrogen half cell, it can induce corrosive dissolution of platinum resulting in the formation of bromoplatinates [4,22–26]. The decay of the H\textsubscript{2} electrode catalyst can be avoided by reducing the concentration of Br\textsubscript{2} in the positive half cell electrolyte.

Stabilization of polybromides by conversion into a less volatile form would reduce the bromine vapor pressure, thus leading to a safer battery electrolyte [7]. Additives used to lower bromine’s volatility have been applied in zinc bromine RFB (Zn/Br\textsubscript{2}-RFB) [8,27–39] and vanadium bromine RFB (V/Br\textsubscript{2}-RFB) [40,41] electrolytes for a long time. These additives are mostly based on organic quaternary ammonium cation compounds and are called bromine complexing agents (BCA). Their bromide or chloride salts are usually highly soluble in aqueous electrolytes. However, their polybromide salts are only slightly soluble or even insoluble in aqueous media [33,36,42]. A further advantage of the polybromide salts of quaternary ammonium BCAs is their low melting point, leading to an insoluble ionic liquid at room temperature. In the electrolytes, these liquids form emulsions that coalesce over time by forming a heavy second phase in the electrolyte [36], as shown in Equation (3). For illustration, Figure 1 shows a photo of an electrolyte solution with a two-phase liquid Br\textsuperscript{−}/Br\textsubscript{2}/H\textsubscript{2}O/[BCA]\textsuperscript{+} electrolyte in the center.

\begin{equation}
[\text{BCA}]^+ (\text{aq}) + \text{Br}_{2n+1}^{−} (\text{aq}) ⇌ [\text{BCA}]\text{Br}_{2n+1} (\text{aq}) ⇌ [\text{BCA}]\text{Br}_{2n+1} (\text{fs}) \tag{3}
\end{equation}

This heavy phase is often referred as the fused salt phase (fs), which sometimes tends to crystallize [31,42]. In the past, many BCAs have been evaluated for their application on zinc bromine batteries, out of which BCAs such as 1-ethyl-1-methylpyrrolidin-1-ium bromide [MEP]Br and 1-ethyl-1-methylmorpholin-1-ium bromide [MEM]Br [8,27–34,38,39,42,43] have received a lot of attention and have been investigated thoroughly. Other BCA structures such as heteroaromatic BCAs based on alkylated pyridine, picolines or imidazole [27,29,30,34,44], symmetrical and unsymmetrical alkylated aliphatic BCAs [33,40,42] and alkylated cyclic cations [27,29,36,42] have also been reported in the literature. Different side chains in N-position such as n-alkyl [27,29,31,33,34,36] or iso-alkyl groups [31] have been used to investigate their steric influence on melting point, complexity and solubility of the fused salt in Zn/Br\textsubscript{2}-electrolytes. 1-carboxymethyl [30], hydroxalkyl groups [27,29] and 1-chloromethyl groups [33,36,42] have been identified to increase Br\textsubscript{2} concentration in the aqueous phase in order to reduce precipitation of [BCA]Br\textsubscript{2n+1}. Polar side chain groups of BCAs have been found to also increase the solubility of the BCA in aqueous solutions. Mixtures of BCAs have been chosen to overlap
Comprehensive work on BCAs has been carried out for Zn/Br₂-RFB electrolyte mixtures and the properties have been described by Cathro et al. [42], Eustace [36] and Lancry et al. [31]. While the work on BCAs for Zn/Br₂-RFB electrolytes is extensive and only a few studies applying BCAs in V/Br₂-RFB have been published so far, no systematic study is available on the application of BCAs in HBr/Br₂/H₂O electrolytes for H₂/Br₂-RFB. In addition, the influence of the different polybromides on the electrolyte properties has not been investigated and is hardly documented in the literature for Zn/Br₂-RFB [17].

1.2. Work Plan

In this work, we investigate 38 quaternary ammonium halides ex situ as prospective BCAs for their suitability for application in Br⁻/Br₂ electrolytes. The aim is to obtain electrolytes for the positive half cell of a H₂/Br₂-RFB with a theoretical volumetric capacity of 180 Ah L⁻¹. All investigated BCAs are shown in Table 1. They comprise of substances from six building blocks: (i) pyridines, (ii) pyrrolidines, (iii) morpholines, (iv) piperidines, (v) 3-methylimidazoles and (vi) tetraalkylammonium compounds. In this research article, the abbreviations listed in Table 1 are used for simplicity. In this study, the BCAs are examined for solubility in the electrolyte, the stability of a second liquid phase at room temperature and their bromine binding strength for a defined electrolyte composition. BCAs that do not form a stable liquid second phase are either not soluble in the electrolyte or show too low bromine binding strength and are excluded from a detailed study as they are not suitable for a battery application. BCAs, which are still of interest, are intensively investigated over the entire state of charge (SoC) range of the electrolyte with a focus on battery performance-relevant parameters: stability, concentration of Br₂ in the aqueous electrolyte, redox potential as well as the electrolytic conductivity of the aqueous phase. For the first time, detailed investigations regarding parameter development are carried out on the basis of the measured distribution of Br₂ on the polybromides tribromide (Br₃⁻), pentabromide (Br₅⁻) and heptabromide (Br₇⁻) in the aqueous phase. Properties of the fused salt phase are not discussed in detail here in order to retain the clarity of the article. However, fused salt generally has low conductivity [36] and is therefore not the preferred option for cell applications. The criteria are defined to select a BCA for the application; the
observed phenomena are explained in detail, and lastly, one BCA is selected for further use in a H₂/Br₂-RFB. The principle of this selection process is depicted in Figure 1.

Table 1. Quaternary ammonium halides for investigation of their properties as bromine complexing agents represented by their structure, name and abbreviation (bold). Letters in brackets identify substances that are synthesized at Fraunhofer ICT (S) and are used from commercial suppliers (C). Details are mentioned in the Supplementary Materials (SM).

| Quaternary Ammonium Compounds Used as BCAs |
|-------------------------------------------|
| 1-Methylpyridin-1-ium hydrobromide, [HMP]Br (S) |
| 1-Ethyl-1-methylpyridin-1-ium bromide, [C2MP]Br (C) |
| 1-α-Butyl-1-methylpyridin-1-ium bromide, [C4MP]Br (C) |
| 1-α-Hexyl-1-methylpyridin-1-ium bromide, [C6MP]Br (S) |
| 1-α-Butyl-1-methylmorpholin-1-ium bromide, [C2MM]Br (S) |
| 1-α-Butyl-1-methylmorpholin-1-ium bromide, [C4MM]Br (C) |
| Pyridin-1-ium hydrobromide, [HPy]Br (S) |
| 1-Ethylpyridin-1-ium bromide, [C2Py]Br (S) |
| 1-α-Butylpyridin-1-ium bromide, [C4Py]Br (S) |
| 1-α-Hexylpyridin-1-ium bromide, [C6Py]Br (S) |
| 1-α-Hexylpyridin-1-ium bromide, [C6Py]Cl (C) |
| 4-Methylpyridine hydrobromide, [H4MP]Br (S) |
| 1-Ethyl-4-methylpyridine hydrobromide, [C4Mpy]Br (S) |
| 1-α-Butyl-4-methylpyridine hydrobromide, [C4Mpy]Br (S) |
| 1-α-Hexyl-4-methylpyridine hydrobromide, [C4Mpy]Br (S) |
| 3-Methylpyridine hydrobromide, [H3Mpy]Br (S) |
| 1-α-Propyl-3-methylimidazol-1-ium bromide, [C3Mim]Br (S) |
| 1-α-Butyl-3-methylimidazol-1-ium bromide, [C4Mim]Br (S) |
| 1-α-Butyl-3-methylimidazol-1-ium chloride, [C4Mim]Cl (C) |
| 1-α-Hexyl-3-methylimidazol-1-ium bromide, [C6Mim]Br (S) |
| 1-α-Hexyl-3-methylimidazol-1-ium chloride, [C6Mim]Cl (C) |
| 1-Methylpiperidin hydrobromide, [HMPip]Br (S) |
| 1-α-Butyl-1-methylpiperidin bromide, [C4MPip]Br (S) |
| 1-α-Hexyl-1-methylpiperidin bromide, [C6MPip]Br (S) |
| 1,1,1-Trime-1-α-tetradecylammonium bromide, [MTA]Br (C) |
| 1,1,1-Trime-1-α-hexadecylammonium bromide, [CTA]Br (C) |
| Tetraethylammonium bromide, [TEA]Br (C) |
| Tetra-α-butylammonium bromide, Tetra-α-octylammonium bromide, [TOA]Br (C) |
| Tetra-α-octylammonium chloride, [TOA]Cl (C) |

2. Results and Discussion

2.1. Stability of Two-Phase Electrolytes and Bromine Binding Strength of BCAs

A total of 38 BCAs are synthesized for this work (marked with “(S)” in Table 1) or ordered from commercial suppliers (marked with “(C)” in Table 1). The component structure of the synthesized BCAs is detected and confirmed by means of 1H NMR and 13C NMR. NMR results of all synthesized components are available in the SM (Section 2) and are verified on the basis of NMR results from the literature [45–51].
2.1.1. Solubility of [BCA]-Salts in Aqueous HBr Solutions

For their application on the H₂/Br₂-RFB, the bromide or chloride salts of the BCAs (Table 1) need to be soluble in aqueous acidic electrolyte at room temperature. This behavior is studied by a simple solubility test. For all investigated BCAs, a solubility test in electrolyte solutions with 1.11 M [BCA]X (X = Br⁻ or Cl⁻) and 5.47 M HBr in H₂O is carried out at θ = 23 ± 1 °C. Most of the investigated substances are soluble in the electrolyte. Exceptions are the three BCAs [CTA]Br, [TOA]Br and [TOA]Cl. These compounds are not completely soluble in the mixture of HBr and H₂O due to their long nonpolar alkyl side chains. Therefore, [CTA]Br, [TOA]Br and [TOA]Cl are not considered as BCAs for an application in a battery and are not investigated further in this work. All the other BCAs of Table 1 are soluble bromide or chloride salts and are investigated on their bromine binding strength and solubility properties related to the addition of Br₂.

2.1.2. Stability of Two-Phase Electrolytes and Bromine Binding Strength of BCAs

The soluble salts should form a liquid fused salt phase in a bromine electrolyte at room temperature, as crystallization of BCA salts in the pump or cell would lead to reduced volume rates and rising parasitic pressures. For the 35 soluble BCA salts, a complexation test in electrolyte solutions by adding 1.11 M Br₂ was performed. The total electrolyte mixture corresponds to a mixture at an SoC of 33% (5.47 M HBr, 1.11 M Br₂, 1.11. [BCA]X in H₂O) at θ = 23 ± 1 °C. The SoC value is chosen, as it represents a 1:1 molar ratio of Br₂ and BCA in the electrolyte. (A definition of the SoC range is provided in the Materials and Methods section.) In the presence of aqueous polybromides, the BCA cations and the polybromides form either an orange-brown liquid fused salt phase or crystallize into reddish-brown or orange crystals. At the same time, a part of the Br₂ remains in the aqueous electrolyte, resulting in a yellow to brown-colored aqueous electrolyte phase, depending on the applied BCA. The concentrations of Br₂ in the aqueous solution and state of aggregation of the second bromine-rich, heavy phase of all BCAs are shown in Figure 2. The results in the figure are sorted according to the decreasing Br₂ concentration in the aqueous electrolyte solution, which reflects the increasing strength of the BCA to bind Br₂ in the second phase. The bromine binding strength of the BCA is defined by the molar fraction of Br₂ stored in the fused salt phase compared to the total amount of Br₂ in the sample. From left to right in Figure 2, the strength of the BCA in bromine binding increases. Additionally, the solubility of [BCA]⁺ cations in the aqueous phase in the presence of polybromides Br₂n⁺1⁻ decreases from left to right following Equation (3). Fused salts that crystallize are shown as red bars, labelled by “S”, while stable liquid phases at room temperature are shown as black bars.

Figure 2 allows an initial classification of the BCAs with regard to the stability of their liquid fused salt phase and their binding strength towards Br₂. The length of the alkyl side chain in N-position of BCAs has a decisive influence on the solubility of the cation in the presence of polybromide anions in the aqueous phase and on the stability of the fused salt. As the length of the alkyl side chain increases, bromine concentrations in the aqueous electrolyte phase (c(Br₂(aq))) decrease, resulting in a rising binding strength of the BCA versus Br₂/polybromides in the fused salt. Already for n-propyl, n-butyl and some BCAs with ethyl side chains in N-position, a concentration of c(Br₂(aq)) < 0.1 M is present in the aqueous phase, whereby more than 91 mol% of Br₂ is transferred into the fused salt. This increased binding of Br₂ may increase the safety of the electrolyte, but it also sustains less Br₂ in the aqueous electrolyte. This can lead to mass transport limitations during cell operation. Liquid fused salts with BCAs which are protonated in N-position [HMP⁺, [H₄MPy]⁺, [H₃MPy]⁺ and [HMPip]⁺ show a lower bromine binding strength with Br₂ concentrations between 0.22 to 0.33 M, which corresponds to a binding strength of only 70 to 80 mol% versus Br₂ in the fused salt. For the right choice of BCA, a compromise between the safety due to strong Br₂ binding property and the availability of polybromides in the aqueous electrolyte at the electrode has to be made. The aqueous phase is essential as it transports bromide, polybromides and protons, all required for the electrochemical
reaction. The concentration values of Br$_2$ in aqueous phases of the 35 samples are shown in detail in Figure S1 in the SM.

![Graph showing BCA binding strength towards Br$_2$](image)

**Figure 2.** Concentrations of Br$_2$ in the aqueous electrolyte phase and aggregate state of the fused salt (black bars = liquid, red bars = solid) of all BCAs shown in Table 1 for a selected state of charge (SoC) of 33% at $\theta = 23 \pm 1$ °C. The sorting is carried out for increasing bromine binding strengths (BCA complexation strength) of the fused salt. Non-soluble BCAs are shown on the right side. The binding strength (text in grey color) of the BCA indicates the fraction of Br$_2$ related to 1.11 M Br$_2$ bound in the fused salt at an SoC of 33%.

On the basis of our studies, we assume that the electron-donating inductive effect (+I-effect) influences the observed trend in Figure 2. The +I-effect increases with rising alkyl side chain length towards the nitrogen atom of the quaternary ammonium and leads to higher bromine binding strength with longer alkyl side chain BCAs and lower [BCA]Br$_{2n+1}$ solubility in the aqueous electrolyte. In detail, due to the +I-effect, the polarity of the N atom is reduced, making the atom increasingly nonpolar and less soluble in the polar HBr/H$_2$O environment as the chain length is increased. In these cases, stronger interactions with the polybromides might be given, while, due to reduced polarity, the interaction with Br$_3^-$, Br$_5^-$ and Br$_7^-$ is preferred compared to polar Br$^-$ ions. To further explain the differences in bromine complexation with increased chain length, in particular on N-heterocycles, former studies on bromine interaction with quaternary ammonium ions can be considered. Easton et al. [17], for example, have calculated the interaction of Br$_3^-$ and Br$_5^-$ with different BCAs by DFT calculation and simultaneously determined the influence of polybromides on the hydrogen atoms in the region around the nitrogen atom in the BCA by $^1$H NMR spectroscopy. The protons in the α-positions of pyridinium, piperidinium and pyrrolidinium rings interact most intensively with the polybromide anion [17], which has also been shown by Lungwitz et al. for the application of polyiodines [52].

### 2.1.3. Influence of Bromide or Chloride Counter Ion of the BCA on Bromine Binding Strength

Aqueous electrolytes containing BCAs with chloride counter ions have an increased bromine binding strength compared to the same [BCA]$^+$ cations with bromide counter ions. They bind bromine 27.9 to 64.3% more strongly in the fused salt in comparison to the equivalent [BCA]$^+$ cation with bromide anions. Next to [C2MIm]Cl, all further [BCA]Cl have a bromine binding strength of 93 mol% or higher, leading to a very low Br$_2$ concentration in the aqueous electrolyte, which significantly limits their application in the
RFB. Based on these results, this study concludes that chloride-based BCAs bind Br$_2$ too strongly, yet a detailed study on the reason for this observation is not within the scope of this research article. It is only known in the literature that bromine forms with chloride mixed polyhalides such as ClBr$_2^-$ or ClBr$_4^-$ [53–55], which may have different properties when in contact with [BCA]$^+$ cations. An evaluation of the differences between the complexation of polybromides or polyhalides for these electrolytes requires a further study.

2.2. Crystallization Behavior of Fused Salt Phases

2.2.1. Influence of the Organic [BCA]$^+$ Cation Structure on the Crystallization of Fused Salt Phases

In order to understand the crystallization behavior of the various BCAs in the form of their fused salts, the structure of the BCA main component and the length of the alkyl side chain in N-position are considered. Crystallization of the fused salts of BCAs with N-hexyl side chains does not occur for all tested BCAs, and in the case of n-butyl substituents it occurs only sporadically. BCAs with an ethyl substituent ([C2MM]$^+$) and 4-methylpyridine-1-ium compounds tend to crystallize more frequently due to less steric hindrance and a higher localized charge. Apart from [C2MM]$^+$, [C2MPy]$^+$ and [C4MPy]$^+$, the fused salt of all heteroaromatic BCAs alkylated in the N-position remains liquid at SoC = 33% and room temperature. The stability of their liquid phase at room temperature has also been shown in the literature for Zn/Br$_2$-RFB electrolytes, but based on different electrolyte composition [27,29,30,34]. Reports on the precipitation of heteroaromatic BCAs could not be found previously in the literature. The previously mentioned [C2MIm]$^+$, [C2MPy]$^+$ and [C4MPy]$^+$, however, offer a pronounced symmetry and predominantly short alkyl side chains. Protonated heteroaromatic BCAs ([HPy]Br, [HMIm]Br) without additional alkyl groups at the aromatic ring also tend to crystallize, while [H4MPy]Br and [H3MPy]Br form a liquid fused salt phase. A stable phase exists for [HMP]$^+$ and [HM Pip]$^+$. Both of the latter remain liquid, and all stable BCAs protonated in the N-position have comparatively high Br$_2$ concentrations between 0.2 M < c(Br$_2$(aq)) < 0.35 M in the aqueous solution. Their bromine binding strength is between 70 and 80 mol% (Figure 2). All examined tetraalkylammonium bromides ([TEA]Br, [TBA]Br and [MTA]Br) are soluble in HBr/H$_2$O but crystallize with polybromides in the given conditions. The symmetry of the BCA molecules leads to crystallization in H$_2$/Br$_2$-RFB electrolytes, which is also reported for this group in Zn/Br$_2$ electrolytes with a different electrolyte composition [42].

In conclusion, organic BCA cations with a high symmetry in structure and short alkyl side chains tend to crystallize in contact with polybromides, while BCAs with long alkyl side chains and/or an unsymmetrical BCA structure stay liquid when forming a fused salt with polybromides.

2.2.2. Polybromide Species in [BCA]Br$_{2n+1}$ Crystals

Beside the structure of BCAs, polybromides are found in the crystals. It raises the question of whether individual polybromides have a decisive influence on the tendency to crystallize with BCA cations. To investigate the reason behind the crystallization, Raman spectra of dry crystals are studied and are shown in Figure S2 of the SM for most of the [BCA]Br$_{2n+1}$ crystals (except for [TBA]Br and [CTA]Br) and in Figure 3a using the example of the BCA [C2MIm]Br.

The Raman spectra of all show the same distinct features: a very strong peak in the Raman spectrum at a Raman shift (υ) between υ = 158–167 cm$^{-1}$ and a weak peak at υ = 180–200 cm$^{-1}$, which is an adjacent shoulder of the strong peak. An example of the Raman spectrum of [C2MIm]Br$_3$ is shown in Figure 3a. Both peaks are characteristic for tribromide ions Br$_3^-$ due to its symmetrical (υ = 163 cm$^{-1}$) and antisymmetrical (υ = 198 cm$^{-1}$) stretching vibration known from [56–58]. The absence of water’s characteristic Raman bands, such as υ = 1600 cm$^{-1}$ and between υ = 3000 cm$^{-1}$ and 3800 cm$^{-1}$ [59–62], ensures that dry crystals have been studied. Peaks of higher polybromides Raman shifts normally appear at υ = 210 cm$^{-1}$ (Br$_5^-$, antisymmetric stretching vibration), υ = 253 cm$^{-1}$ (Br$_5^-$, symmetric stretching vibration) and υ = 270 cm$^{-1}$ (Br$_7^-$, symmetric stretching vibra-
However, higher polybromides seem to be bound in the liquid fused salts. The spectra of [C3MIm]Br2n+1, [C4MIm]Br2n+1 and [C6MIm]Br2n+1 are shown in Figure 3b, where the absence of pentabromide and probably heptabromide is evidenced by the appearance of a peak at ～ν = 255 cm^{-1}. Figure 3b also shows an increase in peak intensity at ～ν = 160 cm^{-1} (Br_3^{-}) with increasing alkyl side chain length of the BCA, whereas the identified peak at ～ν = 255 cm^{-1} (Br_3^{-}/Br_7^{-}) remains constant. In general, the Raman peak shifts of the crystal samples are similar to those of the fused salts [58]. With an increasing side chain length at an SoC of 33%, bromine is preferentially added as Br_3^{-} ions. This means that BCAs with longer side chains preferentially form a fused salt in which Br_3^{-} acts as the counter ion to the [BCA]^+ cation, but does not crystallize due to steric hindrance from the long alkyl side chain. On the other hand, fused salts formed from BCAs with short alkyl side chains are stable liquid phases when, due to a mixture of Br_3^{-}, Br_5^{-} and Br_7^{-}, the fraction of higher polybromides increases in comparison to Br_3^{-}. In this case, the steric hindrance of the polybromides and charge distribution prevents the formation of crystals. The distribution of Br_2 in the polybromides depends on the BCA and the specific interactions between BCA and the polybromides. Haller et al. [63] assume that higher polybromides are less strongly bound by BCAs than lower polybromides or bromide ions.

In summary, symmetric and small BCAs tend to crystallize at room temperature with Br_3^{-} ions because of their higher charge density and lower steric hindrance. Both long side chains of heteroaromatic BCAs and large polybromides do not crystallize at room temperature due to steric hindrance. An exception is [C2Py]Br, which also remains liquid at room temperature.

For further consideration and application, therefore, N-alkylated pyridine-1-ium compounds and 1-alkylated 3-methylimidazol-1-ium compounds are of particular interest. With the alkyl side chains ethyl, n-propyl, n-buty1 and n-hexyl, they all exhibit a bromine binding strength of more than 80 mol% at an SoC of 33% resulting in a reduced vapor pressure in the aqueous electrolyte solution. In order to compare their properties with well-investigated BCAs such as [C2MP]Br and [C2MM]Br, these two BCA groups are included in further experiments. In this first analysis, the focus has been set on the electrolyte mixture of 1:1 ratio of Br_2 to BCA at an SoC of 33%. However, electrolytes at an SoC of 33% only
provide a selective insight into the electrolyte system of a flow battery. In the following, the properties of the electrolytes are examined over the entire capacity range as a function of the SoC.

2.2.3. Stability of Liquid Fused Salts within the Whole SoC Range

In the first preselection process, mainly heteroaromatic 1-alkylpyridin-1-ium and 1-alkyl-3-methylimidazole-1-ium bromides showed stable liquid fused salt phases at room temperature. This has been investigated for a fixed electrolyte mixture of 1.11 M [BCA]Br, 5.47 M HBr and 1.11 M Br₂. To operate the battery with electrolytes containing BCAs, electrolyte stability over the entire SoC range between an SoC of 0% (7.6 M HBr, 1.11 M [BCA]Br and H₂O) and an SoC of 100% (1 M HBr; 1.11 M [BCA]Br, 3.35 M Br₂ and H₂O) is required. Table 2 lists the investigated BCAs: [C2Py]Br, [C4Py]Br, [C6Py]Br, [C2MIm]Br, [C3MIm]Br, [C4MIm]Br and [C6MIm]Br, as well as [C2MM]Br (known as [MEM]Br) and [C2MP]Br (known as [MEP]Br), and indicates their usable SoC range for θ = 23 ± 1 °C.

For different mixtures, it is shown that all BCAs form a second phase over the entire SoC range, except for an SoC of 0%. If Br₂ is added to the aqueous solution, a separate fused salt phase is formed immediately. For all 1-alkylpyridin-1-ium compounds investigated for the entire SoC range, all fused salts remain liquid at room temperature over more than three years. With the exception of [C2MIm]Br, all investigated 3-methyl-imidazolium compounds form a liquid fused salt phase at room temperature within the entire SoC range. Electrolyte samples with the aqueous and fused salt phase in the whole SoC range for [C2MIm]Br, [C2Py]Br and [C6Py]Br are shown in Figure 4.

[C2MP]Br, [C2MM]Br and [C2MIm]Br form a solid crystalline phase with polybromides in the lower SoC range (Table 2). With these three BCAs, a maximum of 40 to 70% of the capacity range (71.8 to 125.7 Ah L⁻¹) can be used at room temperature, as a crystallization of the fused salt cannot be accepted in the battery during operation. For [C2MM]Br, [C2MP]Br and [C2MIm]Br, dry crystals of the second phase in the lower SoC range are again examined by Raman spectroscopy. In the SoC range in which the fused salts tend to form crystals, only tribromide Br₃⁻ salts of the BCAs are identified in addition to the characteristic peaks for the respective organic cation (Table 2).

In previous studies, [C2MM]Br and [C2MP]Br have been applied in electrolytes for Zn/Br₂-RFB individually or in mixtures [8,28,30,33,38,41–43]. While the liquid fused salts of the Zn/Br₂ electrolytes are stable at room temperature, these BCA form crystals for the selected acidic concentration at different SoC of a H₂/Br₂-RFB electrolyte. [C2MIm]Br, [C2MM]Br and [C2MP]Br are therefore excluded for use at room temperature and lower temperatures in H₂/Br₂-RFB since the formation of crystals in the cell or during operation can lead to strong overpressure in the system. [C2MM]Br and [C2MP]Br as BCAs are not investigated further in this study. On the other hand, [C2MIm]Br is reintroduced in the next experiments in order to compare its properties to the other derivatives such as [C3MIm]Br, [C4MIm]Br and [C6MIm]Br.

2.3. Bromine Binding Strength of [BCA]⁺(aq) Cations as a Function of the State of Charge (SoC)

The bromine binding strength of the individual BCAs has been determined based on the concentration c(Br₂) in the aqueous electrolyte solution compared to the total Br₂ concentration at an SoC of 33% for the 35 chosen BCAs in Figure 5. Now, the change of the [BCA]⁺ concentration over the SoC range will be considered to investigate its effect towards the solubility and bromine binding properties for the seven BCAs based on 1-alkyl-1-pyridin-1-ium bromides and the 1-alkyl-3-methylimidazol-1-ium bromides. For the measurement of the BCA concentration in aqueous solution, the intensity of the characteristic peaks with corresponding Raman shifts is investigated. The Raman shifts of the [BCA]⁺ cations in aqueous solution are listed in Table 2. The Raman peaks selected for the evaluation have certain Raman shifts, which are highlighted in Table 2 in bold. The designated Raman shifts for the investigated BCAs are in agreement with the literature [64]. Concentrations of [BCA]⁺ cations in the aqueous phase are shown in Figure 5.
Table 2. Selected BCAs to be investigated over the entire SoC range including the zone in which the second phase is present as a liquid fused salt. Furthermore, characteristic Raman shifts of the [BCA]$^+$ cation are shown, whereby the Raman peak with the Raman shift in bold is used to determine the [BCA]$^+$ concentration. In addition, the Raman shifts of the symmetrical stretching vibration of the individual polybromides are shown, which are used for the evaluation in Section 2.5.

| BCA Structure/Name/Abbreviation | SoC Range of Liquid Fused Salt | Characteristic Cation Raman Shifts/cm$^{-1}$ | Polybromide Raman Shifts in Aqueous Solutions/cm$^{-1}$ |
|---------------------------------|--------------------------------|---------------------------------------------|------------------------------------------------------|
| 1-Ethyl-1-methylpyrrolidin-1-iumbromide, [C2MP]Br (= [MEP]Br) | SoC $\geq$ 30% | 899, 1452, 2954, 2964, 2989 | $\nu_5$ (Br$_3^-$) 165–169, $\nu_5$ (Br$_5^-$) 255 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-Ethyl-1-methylmorpholin-1-iumbromide, [C2MM]Br (= [MEM]Br) | SoC $\geq$ 60% | 685, 701, 2888, 2946, 2985 | $\nu_5$ (Br$_3^-$) 165–167, $\nu_5$ (Br$_5^-$) 254 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-Ethylpyridin-1-iumbromide, [C2Py]Br | SoC 0–100% | 647, 1029, 1177, 2883, 2919, 2945, 2988, 3098 | $\nu_5$ (Br$_3^-$) 165–167, $\nu_5$ (Br$_5^-$) 253–254 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-n-Butlypyridin-1-iumbromide, [C4Py]Br | SoC 0–100% | 651, 1030, 1179, 2880, 2914, 2943, 2976, 3097 | $\nu_5$ (Br$_3^-$) 167–170, $\nu_5$ (Br$_5^-$) 254–257 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-n-Hexlypyridin-1-iumbromide, [C6Py]Br | SoC 0–100% | 649, 1029, 2869, 2910, 2939, 2973, 3097 | $\nu_5$ (Br$_3^-$) 164–169, $\nu_5$ (Br$_5^-$) 255–256 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-Ethyl-3-methylimidazol-1-iumbromide, [C2MIm]Br | SoC $\geq$ 50% | 598, 1024, 1339, 1419, 2961 | $\nu_5$ (Br$_3^-$) 165–168, $\nu_5$ (Br$_5^-$) 253–254 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-n-Propyl-3-methylimidazol-1-iumbromide, [C3MIm]Br | SoC 0–100% | 1023, 1337, 1417, 2961 | $\nu_5$ (Br$_3^-$) 167–170, $\nu_5$ (Br$_5^-$) 254 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-n-Butyl-3-methylimidazol-1-iumbromide, [C4MIm]Br | SoC 0–100% | 1024, 1339, 1419, 2963 | $\nu_5$ (Br$_3^-$) 165–170, $\nu_5$ (Br$_5^-$) 254–255 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
| 1-n-Hexyl-3-methylimidazol-1-iumbromide, [C6MIm]Br | SoC 0–100% | 1025, 1122, 1341, 1418, 2961 | $\nu_5$ (Br$_3^-$) 164–167, $\nu_5$ (Br$_5^-$) 253 |
|                                |                                |                                             | $\nu_5$ (Br$_7^-$) 269 |
Figure 4. Photos of electrolyte samples with different BCAs [C2MIm]Br, [C2Py]Br and [C6Py]Br as a function of SoC for an SoC of 0, 10, 20, 30, 33, 40, 50, 60, 66, 70, 80, 90 and 100%. For electrolytes with [C2MIm]Br, between 0 < SoC < 60%, the second electrolyte phase is shown to precipitate and to form crystals. For [C2MIm]Br at SoC ≥ 60%, [C2Py]Br, as well as [C6Py]Br in the entire SoC range, the formation of a liquid fused salt phase occurs at room temperature. With increasing SoC, the volumes of the fused salt phase increase. In particular for the application of [C6Py]Br, the coloration of the aqueous phase from yellow to brown shows that the bromine concentrations increase with increasing SoC. In addition, the SoC range is specified by total concentrations for an SoC of 0% and 100%.

Figure 5. Concentration of [BCA]^+ cations in the aqueous electrolyte phase containing (a) 1-alkylpyridine-1-ium bromides and (b) 1-alkyl-3-methylimidazole-1-ium bromides used as BCAs with different alkyl radicals in N-position as a function of the state of charge (SoC). All values of Figure 5 are printed in the SM in Table S1.

When the BCA is added to the polybromide solution the solubility equilibrium leads to a separation of [BCA]Br_{2n+1} salts by forming a fused salt phase according to Equation (3), or, as in the case of [C2MIm]Br, to the formation of crystals. The phase separation leads to a decrease in the concentration of [BCA]^+ (aq) regardless of the [BCA]^+ cation type as shown in Figure 5. The concentration dependence of the [BCA]^+ cations on the SoC shows a characteristic curve. At an SoC of 0% and thus c(Br_2) = 0 M, a concentration of c([BCA]^+) = 1.11 M is present, which decreases for all BCAs at SoC > 0%. Especially for 3-methylimidazol-1-ium derivatives, a linear and BCA-independent decrease in the [BCA]^+ concentration between 0 ≤ SoC ≤ 20% can be observed (Figure 5b). For pyridin-
1-ium derivatives, the same trend is observed between 0 ≤ SoC ≤ 10% in Figure 5a. Between 20 < SoC < 66%, the concentration of BCA continues to decrease for both groups of substances, although these changes become smaller as the SoC increases. In this range, the different bromine binding strength and solubility of the [BCA]⁺ cation with respect to the polybromide species become visible. [C6Py]Br and [C6MIm]Br show a considerably lower concentration in aqueous solution and therefore solubility as well as their property to bind bromine more strongly, respectively, than that of BCAs with shorter alkyl side chains. Organic cations of [C6Py]Br and [C6MIm]Br are no longer detectable in the solution even at SoCs of 50% and 40%, respectively. [C4Py]Br, [C4MIm]Br and [C3MIm]Br follow this trend at higher SoCs. [C2Py]Br and [C2MIm]Br show the lowest bromine binding strength versus polybromides. Thus, higher concentrations of [C2Py]⁺ and [C2MIm]⁺ remain in aqueous electrolytes at a higher SoC. From SoC ≥ 70% on, [C2Py]⁺ and [C2MIm]⁺ are not detectable in the Raman spectrum. Experimentally determined detection limits for [BCA]⁺ cations in aqueous solution are listed in Table 3/line 1. Due to an increasing total Br₂ concentration with an increasing SoC, the [BCA]⁺ cations are gradually transferred into the fused salt phase. In principle, the SoC range can be divided into two sections: Depending on the [BCA]⁺ cation, there is a section for lower SoC values where [BCA]⁺ cations are present in the aqueous solution, and a section for higher SoC values, where [BCA]⁺ cations are not present in the aqueous solution but are completely present in the fused salt phase.

Table 3. Detection limits and detection range for [BCA]⁺ in the aqueous electrolyte phase, maximum fractions of Br₂ in the aqueous phase compared to the global concentration and c(Br₂)aq for an SoC of 33%.

| [BCA]⁺ | [C2Py]⁺ | [C4Py]⁺ | [C6Py]⁺ | [C3MIm]⁺ | [C4MIm]⁺ | [C6MIm]⁺ |
|--------|---------|---------|---------|----------|----------|----------|
| SoC limit for detectable [BCA]⁺ [%] [a] | ≤70    | ≤66    | ≤40    | ≤60    | ≤50    | ≤40    | ≤33    |
| c(Br₂)(aq)/c(Br₂)total [%] [b] | ≤10.1  | ≤10.3  | ≤8.5  | ≤10.4  | ≤10.0  | ≤10.6  | ≤9.5  |
| C(Br₂)(aq) at an SoC of 30%/mM | 121   | 91     | 9      | 105    | 63     | 34     | 8      |

[a] The SoC value corresponds to the highest SoC at which [BCA]⁺ cations can still be detected in the aqueous phase. [b] Highest ratio of Br₂ concentration in the aqueous phase versus the total Br₂ concentration in the sample detected at an SoC of 90% or 100%.

The individual, BCA-dependent bromine binding strength is determined from the [BCA]⁺ concentrations in the range of 20 ≤ SoC ≤ 60% and is applicable to investigate the solubility behavior of the BCA. The BCAs can be sorted according to their concentration in the aqueous phase. Low concentrations of [BCA]⁺(aq) in the aqueous phase describe a larger bromine binding strength and vice versa. The bromine binding strength is obtained in the following order: [C6MIm]⁺(aq) ≥ [C6Py]⁺(aq) > [C4MIm]⁺(aq) > [C4Py]⁺(aq) > [C3MIm]⁺(aq) > [C2MIm]⁺(aq) > [C2Py]⁺(aq). Correspondingly, the solubility of [BCA]Br₂n+1 in the aqueous electrolyte for the 1-alkylpyridine-1-ium polybromides and for the 1-alkyl-3-methylimidazole-1-ium polybromides decreases with an increasing length of the alkyl side chain. The detection limits of [BCA]⁺ cations in Table 3 show that 1-alkyl-3-methylimidazol-1-ium is a slightly stronger bromine binding BCA than 1-alkylpyridin-1-ium if both have the same alkyl group in the N-position. As mentioned earlier, the bromine binding strength cannot be the only criteria to choose a BCA. Moderate Br₂ concentrations in the aqueous phase along the whole SoC range are a prerequisite for the operability of the cell and will be investigated in the next section.

2.4. Effect of the Solubility Equilibrium on the Br₂(aq) Concentration for the Whole SoC Range

Br₂ is transported from the tank into the RFB half cell as an ingredient of the aqueous solution. Moderate Br₂ concentrations dissolved as polybromide ions in the aqueous electrolyte phase are a prerequisite for stable cell operation. The bromine concentrations in the aqueous phase of the electrolytes for the investigated BCAs are plotted as a function of the SoC in Figure 6.
2.4.1. General Influence of BCA Application on Br₂ Concentration in Aqueous Solution

Due to the relatively low solubility of the BCA-polybromide salt, the Br₂ concentration in the aqueous solution in equilibrium is low compared to the global concentration of Br₂ indicated by the absolute concentration in Figure 6. While the total concentration of Br₂ varies linearly along the SoC range, between \( c(\text{Br}_2(\text{total})) = 0.35 \, \text{M} \) at an SoC of 10% and \( c(\text{Br}_2(\text{total})) = 3.35 \, \text{M} \) at an SoC of 100% (Figure 6), the solubility equilibrium of the BCAs with the polybromides results in a significantly lower concentration of bromine \( (\text{c(Br}_2(\text{aq})) < 0.35 \, \text{M}) \) in the entire SoC range for all investigated BCAs. In addition, no linear relationship between \( c(\text{Br}_2(\text{aq})) \) and the SoC is observed. The maximum ratio of Br₂ in the aqueous phase compared to the total Br₂ concentration is present for SoC > 80% and is shown in Table 3/line 2. A maximum fraction of 11 mol% Br₂ is present in the aqueous solution, while a fraction of more than 89 mol% Br₂ is present in the fused salt phase. Although vapor pressures are not measured in this work, the safety of the electrolytes increases tremendously due to the low Br₂ concentrations and the resulting reduced outgassing of Br₂ from the aqueous electrolyte. Additionally, low concentrations of c(Br₂(\text{aq})) would decrease the available bromine that can crossover across the membrane to the H₂ electrode inside the battery.

2.4.2. Individual Concentration Trends of Different [BCA]+ Cations

For BCAs based on 1-alkylpyridin-1-i um and 1-alkyl-3-methylimidazol-i um, characteristic concentration curves for c(Br₂(\text{aq})) are obtained, whereby both diagrams in Figure 6 can be divided into two sections: In the first section (section I), between \( 0 < \text{SoC} \leq 40\% \) the Br₂ concentrations in the aqueous solution are different for each BCA but are approximately independent of SoC and are also listed for an SoC of 30% in Table 3, on line 3. One exception is the trend of [C2MIm]Br, which will be explained later. There is a clear trend in Br₂ concentrations in the aqueous phase at an SoC of 30% for the applied BCAs, which corresponds to the trend of BCA-concentrations in Section 2.3: \( c(\text{Br}_2; [\text{C6MIm}]) < c(\text{Br}_2; [\text{C5Py}]) < c(\text{Br}_2; [\text{C4MIm}]) < c(\text{Br}_2; [\text{C3MIm}]) < c(\text{Br}_2; [\text{C2Py}]) \). In this range, the bromine binding strength of the [BCA]⁺ cation is clearly visible.

**Figure 6.** Concentrations of Br₂ solved in the form of polybromide ions in the aqueous electrolyte phase depending on the state of charge (SoC) for the investigated BCAs: (a) 1-alkylpyridine-1-ium bromides and (b) 1-alkyl-3-methylimidazole-1-ium bromides as BCAs with different alkyl side chains in the N-position at \( \theta = 23 \pm 1 \, ^\circ \text{C} \). All values of Figure 6 are printed in the SM in Table S2. Section I shows a strong dependence of the Br₂ concentration in the aqueous phase on the alkyl side chain length of the BCA, while in section II the Br₂ concentration is independent of the alkyl side chain length. The total Br₂ concentration in the samples allows a direct comparison of the Br₂ concentration in the sample and the aqueous phase for 10 ≤ SoC ≤ 100%.

Table 3. Detection limits and detection range for [BCA]+ in the aqueous phase of the electrolytes for the investigated BCAs are plotted as a function of the SoC in Figure 6.
again and corresponds exactly to the mentioned trends in Section 2.3: \([\text{C}_6\text{MIm}]^+\text{(aq)} > [\text{C}_6\text{Py}]^+\text{(aq)} > [\text{C}_4\text{MIm}]^+\text{(aq)} > [\text{C}_4\text{Py}]^+\text{(aq)} > [\text{C}_3\text{MIm}]^+\text{(aq)} > [\text{C}_2\text{MIm}]^+\text{(aq)} > [\text{C}_2\text{Py}]^+\text{(aq)}\). The alkyl side chain length in the N-position in all BCAs is responsible for the differences in complexing strength, which is just a handy indicator for the solubility of the \([\text{BCA}]\text{Br}_{2n+1}\). The longer the side chain, the lower the solubility of the \([\text{BCA}]\text{Br}_{2n+1}\) in the aqueous solution. Another finding is that in general, the 1-alkyl-3-methylimidazol-1-ium compounds are slightly stronger BCAs compared to the 1-alkylpyridin-1-ium, when the same alkyl side chains are introduced. The concentrations of \(\text{Br}_2\) and \([\text{BCA}]^+\) cations in the aqueous solution both show the same trend.

For the \(\text{Br}_2\) concentrations of \([\text{C}_2\text{MIm}]\text{Br}\) between \(10 \leq \text{SoC} \leq 50\%\), there is a concentration plateau in Figure 6b which differs from the \(c(\text{Br}_2(\text{aq}))\) curves for \([\text{C}_3\text{MIm}]\text{Br}, [\text{C}_4\text{MIm}]\text{Br}\) and \([\text{C}_6\text{MIm}]\text{Br}\). As mentioned above, the second phase of \([\text{C}_2\text{MIm}]\text{Br}\) is crystalline and consists solely of the tribromide compound ((\text{C}_2\text{MIm})\text{Br}_3). Consequently, the ratio of bromine to BCA cation is fixed to 1:1, and no higher polybromides can be stored in the second heavy phase. This leads to rising \(\text{Br}_2\) concentrations in the aqueous phase until the second phase liquefies again at \(\text{SoC} \geq 50\%\), as presented in Figure 4.

The second section in Figure 6a,b starts from \(\text{SoC} > 40\%\) (section II). Here the \(\text{Br}_2\) concentration increases noticeably for all BCAs, whereby the dependence of the \(\text{Br}_2\) concentration on the choice of the \([\text{BCA}]^+\) cation decreases with increasing \(\text{SoC}\). For both 1-alkylpyridin-1-ium and 1-alkyl-3-methylimidazolium cations as BCAs, the \(\text{Br}_2\) concentration for \(\text{SoC} \geq 60\%\) increases, whereby for each \(\text{SoC}\) value examined, approximately equal concentrations \(c(\text{Br}_2(\text{aq}))\) are present in the aqueous solution. In this section, the length of the alkyl side chain at the N-position of the BCA seems to have no influence on the binding strength of \(\text{Br}_2\) of the second phase in solution. However, 1-alkylpyridin-1-ium bromides bind bromine in section II slightly more strongly and tend to have a lower solubility in the presence of polybromides at \(\text{SoC} > 60\%\) as slightly lower \(\text{Br}_2\) concentrations in the aqueous phase are present (Figure 6). In general, for \(\text{SoC} \geq 80\%\) the \(\text{Br}_2\) concentrations rise only slowly or form a plateau. In that range, the solubility of \(\text{Br}_2\) in the aqueous solution is limited by a lack of bromide ions to form polybromides, known from BCA-free electrolytes [65].

It is noticeable over the whole range that for small amounts of \(\text{Br}_2\) at \(\text{SoC} \leq 40\%\), the complexation strength of the individual BCA types has a major influence on bromine concentration in the aqueous phase, but at \(\text{SoC} \geq 60\%\) neither the type of BCA base nor the length of the alkyl side chain seem to play a noticeable role, where all BCAs are considered to complex polybromides to a similar extent. At the same time, there is a transfer of \(\text{Br}_2\) from the aqueous phase into the fused salt over the entire \(\text{SoC}\) range, but since it has been shown above that no BCA cations are present in the aqueous solution at higher \(\text{SoCs}\), the transfer no longer occurs according to Equation (3).

2.4.3. Theoretical Comparison of \(\text{Br}_2\) Concentrations and Current Densities in the \(\text{H}_2/\text{Br}_2\)-RFB Cell

Overall, the concentrations of \(c(\text{Br}_2(\text{aq}))\) are concentrations of \(\text{Br}_2\) in chemical equilibrium for both phases and are only present during operation if the mass transfer of \(\text{Br}_2\) from the fused salt to the aqueous electrolyte phase takes place without any mass transfer limitation, i.e., the equilibrium state is maintained during the discharge process. Applying this theoretical assumption, the current densities of \(i = 723 \text{ mA cm}^{-2}\) for \(c(\text{Br}_2) = 0.3 \text{ M}\), \(i = 482 \text{ mA cm}^{-2}\) for \(c(\text{Br}_2) = 0.2 \text{ M}\) and \(i = 241 \text{ mA cm}^{-2}\) for \(c(\text{Br}_2) = 0.1 \text{ M}\) would be possible in a cell with a geometrically active electrode/membrane area of 40 \(\text{cm}^2\) and a volume flow rate of 30 \(\text{mL min}^{-1}\). Data for this cell are described in [65]. For electrolytes with lower concentrations, the possible current densities would decrease further. Only \([\text{C}_2\text{Py}]\text{Br}\) forms a liquid fused salt over the entire \(\text{SoC}\) range and has concentrations \(c(\text{Br}_2) \geq 0.121 \text{ M}\) in equilibrium between the two phases. In reality, lower concentrations can be expected in the aqueous solution. Therefore, a current density \(i < 250 \text{ mA cm}^{-2}\) for galvanostatic charging and discharging can also be expected for electrolytes with \([\text{C}_2\text{Py}]\text{Br}\). As \([\text{H}_4\text{MPy}]\text{Br}, [\text{H}_3\text{MPy}]\text{Br}\) and \([\text{HMPpip}]\text{Br}\) upon first glance have aqueous \(\text{Br}_2\) concentrations between
0.20 M and 0.35 M at an SoC of 33% in Figure 2, they seem to be more applicable for effective cell operation. From Figure 6 it is noticed that in the range of an SoC of 33%, Br₂ concentrations in the aqueous solutions are lowest. For the protonated forms it is therefore expected that for higher SoCs, much larger Br₂ concentrations are present and electrolytes cannot still be classified as “safe” electrolytes. From the results in this section, it is expected that concentrations might rise for a higher SoC and lead to greater volatility of Br₂. [C2Py]Br tends to be the [BCA] of choice in order to reach performance and safety of the Br₂/Br⁻ electrolytes.

2.5. Distribution of Br₂ on Polybromides in the Aqueous Phase

As presented in the previous section, the aqueous Br₂ concentration defines the power capability of the cell. For a deeper understanding of the complexation behavior of the different BCAs, the Raman spectra of the aqueous phase have been analyzed regarding the distribution of bromine in the different polybromides. The method has been described in our earlier work [65]. The distribution of Br₂ on Br₃⁻, Br₅⁻ and Br₇⁻ is shown in Figure 7. The Raman spectra of all investigated samples are shown in the SM in Figures S3–S9.

![Figure 7. Distribution of Br₂ in the aqueous electrolyte phase on the polybromides Br₃⁻, Br₅⁻ and Br₇⁻, which are present, depending on the SoC with different BCAs with different alkyl side chains in the N-position for (a) 1-alkylpyridine-1-ium bromides and (b) 1-alkyl-3-methylimidazole-1-ium bromides as BCAs at θ = 23 ± 1 °C. All values of Br₂ distribution on polybromides are printed in the SM in Tables S3-S5.](image)

An analysis of the distribution of Br₂ among the polybromides Br₃⁻, Br₅⁻ and Br₇⁻ in the aqueous phase shows basically similar results for both BCA groups as shown in Figure 7a,b. For all investigated BCAs, the storage of Br₂ in the aqueous electrolyte in the form of Br₃⁻ predominates, followed by storage in Br₅⁻ and finally storage in Br₇⁻. This is in accordance with earlier studies on pure BCA-free HBr/Br₂/H₂O-electrolytes [65], leading to the result that [BCA]⁺ has only a minor influence on the fractions of Br₂ in the different polybromide species in the aqueous solution. For SoC < 40%, the n-hexyl side chain BCAs [C6Py]Br and [C6MIm]Br have a slightly increased Br₂ content in Br₃⁻, which means that less Br₂ is distributed to the Br₅⁻ and Br₇⁻ anions. For SoC < 40%, n-butyl, n-propyl and ethyl BCAs have a constant Br₂ distribution, which is independent of the BCA. For BCAs alkylated with an n-hexyl group, more Br₃⁻ is available, while less Br₂
is stored in Br$_5^-$ and Br$_7^-$. In this range, cations of [C6Py]$^+$ and [C6MIm]$^+$ are available in the aqueous solution and already prefer to interact with Br$_5^-$ in comparison to Br$_3^-$ and Br$_7^-$. This is in accordance with the composition of their fused salt phases, shown in Figure 3b.

For an SoC of $\geq 40\%$, there is no dependence of the Br$_2$ distribution in the aqueous phase on the length of the alkyl radical present in the N-position of the BCA. The distribution is independent of the choice of the BCA basic compound pyridine or 3-methylimidazol. Within this SoC range, the fraction of Br$_2$ stored as Br$_3^-$ increases from 30.7% to 41.6–47.3%, while proportionally less Br$_2$ is stored in Br$_5^-$ and the curve decreases from 60% to 46–50%. Br$_2$ remains in Br$_7^-$ within the whole SoC range relatively constant up to a maximum of 13.5%. Presumably, a better solvation of the more stable Br$_3^-$ compared to Br$_5^-$ is present. This effect is obtained also for BCA-free HBr/Br$_2$/Br$_2$O solutions [65]. Since the Br$_2$ concentration of c(Br$_2$(aq)) < 0.35 M and the distribution of Br$_2$ to the polybromides is relatively constant in the whole range, no specific influence of this distribution on the performance or individual parameters is given and therefore cannot be discussed.

### 2.6. Influence of Different BCAs on Electrolyte Conductivity of the Aqueous Phase

For both flow-by and flow-through electrodes in RFBs, a low ohmic cell resistance is required to achieve high cell or stack performance. In RFB, this strongly depends on the electrolyte composition at different SoCs, usually targeting the maximum electrolyte conductivity of the aqueous electrolyte phase. Electrolyte conductivities of the aqueous phases from the seven pre-selected [BCA]/Br$_2$ electrolytes summarized in Table 2 are shown in Figure 8. In addition, the reference conductivity curves of a BCA-free HBr/Br$_2$/H$_2$O electrolyte (from [65]) and a pure HBr/H$_2$O acid solution within the SoC range are shown in Figure 8. All measured values are listed in Tables S6 and S7 in the SM. The values for pure HBr/H$_2$O acid at $\theta = 20$ °C are calculated from molar conductivities from [66] and are adapted to the SoC scale.

![Figure 8](image)

**Figure 8.** Specific electrolyte conductivity $\kappa$ of the aqueous electrolyte phase containing (a) 1-alkylpyridin-1-ium bromides and (b) 1-alkyl-3-methylimidazol-1-ium bromides as BCAs with different alkyl side chains in the N-position as a function of state of charge SoC at $\theta = 23 \pm 1$ °C. In addition, electrolyte conductivities of BCA-free HBr/Br$_2$/H$_2$O electrolytes are shown (green dots) from [65] and of pure HBr/H$_2$O solutions (orange line) from [66] at $\theta = 20$ °C with the same total HBr and/or Br$_2$ concentrations as reference curves.

Electrolytic conductivities in the aqueous phase depend mainly on the high proton concentration and are larger than 300 mS cm$^{-1}$ throughout the SoC range, whereby the Grotthuss mechanism [67,68] is responsible for ionic transport through the solution. In
with longer alkyl side chains lead to lower conductivity in the aqueous phase due to their presence in the entire SoC range compared to BCA-free HBr/Br2/H2O electrolytes, as shown in Figure 5. For 0 ≤ SoC ≤ 60% for both 1-alkylpyridin-1-ium electrolytes and for 1-alkyl-3-methylimidazol-1-ium electrolytes, κ rises approximately from 368.90 to 478.00 mS cm⁻¹ at an SoC of 0% to reach a peak at κ = 643.1–681.84 mS cm⁻¹ between an SoC of 50 and 60%. Within this range, there is a difference from the two reference curves in accordance with the concentrations of the [BCA]+ cation in solution, shown in Figure 5. Initially, the organic character of [BCA]+ cation decreases the conductivity. Organic [BCA]+ cations with longer alkyl side chains lead to lower conductivity in the aqueous phase due to their increasing size. On the one hand, their migration in the electric field is slowed down and the passive transport by the Grotthus mechanism is restricted. Grotthus charge transfer on a direct path between the electrodes is hindered and is forced to bypass the large organic cations. However, as the SoC is increased, the conductivity of the aqueous electrolyte in Figure 8 is shown to increase. This observation can be directly related to the results presented in Figure 5, whereby the [BCA]+ concentration of the investigated BCAs in the aqueous electrolyte is shown to decrease with the SoC as an indication that the cations are now bonded to the fused salt phase and leaving the aqueous electrolyte phase. Thus, based on these results, the decrease in the aqueous electrolyte [BCA]+ concentration translates into the increased electrolyte conductivity trend seen in Figure 8 between 0 ≤ SoC ≤ 60%.

For electrolytes with 1-alkylpyridin-1-ium bromide as a BCA, the complexation strength with respect to the side chain can be read in Figure 8a. Since [C6Py]+ with polybromides Br2n+1− is less soluble in the aqueous phase than [C2Py]+, their concentration decreases more rapidly compared to [C2Py]+. The conductivity of the solution increases correspondingly more quickly in Figure 8a compared to electrolytes containing [C4Py]+ or [C2Py]+.

In the second range for 60 < SoC ≤ 100%, there are hardly any [BCA]+ cations available in the aqueous phase, as presented in Section 2.3. In this range, no direct dependence of conductivity on the alkyl side chain in the N-position of the [BCA]+ cation is obtained. Electrolytes with BCAs show approximately same conductivities. With an increasing SoC, the proton concentration decreases, and the conductivity of the solution decreases subsequently.

However, electrolytes containing BCAs reach higher conductivities compared to BCA-free HBr/Br2/H2O electrolytes for SoC ≥ 50% (green dots, in Figure 8). Due to the transfer of Br2 by use of the BCA from the aqueous phase into the fused salt, only concentrations lower than 0.35 M Br2 to the maximum total concentration of 2.99 M Br2 in BCA-free HBr/Br2/H2O electrolyte (calculated from values in [65]) are present in aqueous electrolytes. Since the absolute Br2 concentration in the aqueous electrolytes with BCA is much lower, while the concentration of protons in both solutions is assumed to be the same, the conductivity is higher when using BCAs and is comparable to pure HBr/H2O solutions as shown in Figure 8. The Groththus mechanism takes place directly between the electrodes and does not need to bypass larger polybromides, as is the case in BCA-free HBr/Br2/H2O electrolytes [65]. As for SoCs > 60%, the [BCA]+ cations are not available in the aqueous phase; the conductivity of the resulting electrolytes in that range is shown to be higher than in BCA-free HBr/Br2/H2O electrolytes.

Overall, in the complete SoC range there are conductivities between a minimum of 324.8 mS cm⁻¹ (SoC 100%) and a maximum of 712.0–745.0 mS cm⁻¹ (an SoC of 50%, depending on the [BCA]+), which are large in comparison to aqueous electrolytes of Zn/Br2-RFB using the example of Eustace [36] with conductivities between 44.4 ≤ κ ≤ 128.2 mS cm⁻¹ (θ = 23 °C). Pure aqueous solutions of comparable BCAs [C2MIm]Cl, [C4MIm]Cl and [C6MIm]Cl offer...
conductivities of \( \kappa \leq 82 \text{ mS cm}^{-1} \) at \( \theta = 25 ^\circ \text{C} \) \[69\], whereby a clear dependence on the alkyl radical in the N-position of the 3-methylimidazole-1-ium can be confirmed from \[69\] with \( \kappa([\text{C2MIm}]\text{Cl(aq)}) > \kappa([\text{C4MIm}]\text{Cl(aq)}) > \kappa([\text{C6MIm}]\text{Cl(aq)}) \).

In this work, the achieved conductivities correspond to those of a pure HBr/H\(_2\)O solution, which for an SoC of \( \geq 60\% \) means an increase in conductivity compared to HBr/Br\(_2\)/H\(_2\)O electrolytes and has a positive effect on cell performance. These high conductivities are due to the presence of the high proton concentration of c(HBr) \( \geq 1 \text{ M} \). The influence of BCAs is directly attributable to the presence of BCAs for an SoC of \( < 60\% \) and indirectly for an SoC of \( \geq 60\% \). These high conductivities allow one to obtain high current densities at a low expected electrolyte overvoltage, depending on the half-cell structure.

2.7. Comparison of the Redox Potential of BCA-Free and BCA-Containing Electrolytes

To obtain a high-power density on an H\(_2\)/Br\(_2\)-RFB cell, a high thermodynamic cell voltage is beneficial. The thermodynamic cell voltage of the H\(_2\)/Br\(_2\)-RFB is composed of the difference of the half-cell redox potentials \( \varphi \) vs. NHE in open circuit state of the RFB. The redox potential of the positive half-cell \( \varphi(\text{Br}_2^-/\text{Br}^-) \) depends, in the simplest scenario, on the concentrations of the redox pair Br\(^-\)/Br\(_2\) or Br\(^-\)/Br\(_{2n+1}^-\) according to Nernst \[70\], while assuming activity coefficients (\( \gamma \)) of \( \gamma = 1 \) for all components of the redox reaction according to Equation (4):

\[
\varphi(\text{Br}_2^-/\text{Br}^-) = \varphi^0(\text{Br}_2^-/\text{Br}^-) - 0.5 \frac{R}{F} T^{-1} \ln(c(\text{Br}^-)^2/c(\text{Br}_2^-)^{-1}) \tag{4}
\]

In Equation (4), \( F \) is the Faraday constant (96485 As mol\(^{-1}\)), \( R \) the molar gas constant (8.3145 J mol\(^{-1}\) K\(^{-1}\)) and \( T \) the temperature in K. \( \varphi^0(\text{Br}_2^-/\text{Br}^-) \) is the standard redox potential of the Br\(_2\)/Br\(^-\) electrolyte with 1.09 V \[9\]. The theoretical potential curve following the Nernst equation (Equation (4)) and global concentrations of HBr and Br\(_2\) at \( T = 298.15 \text{ K} \) are applied. The curve is shown in Figure 9 as a function of the SoC (orange line). It has much higher values at corresponding SoCs compared to the measured redox potential of BCA-free electrolytes in Figure 9 (green dots). In the electrolyte system under investigation, there are no ideally diluted solutions, but strongly concentrated electrolytes. As a result, redox potentials according to the NERNST equation must be considered including activity coefficients as shown in Equation (5) \[70\]:

\[
\varphi(\text{Br}_2^-/\text{Br}^-) = \varphi^0(\text{Br}_2^-/\text{Br}^-) - 0.5 R.T.F^{-1} \ln(c(\text{Br}^-)^2 \gamma(\text{Br}^-)^2/c(\text{Br}_2^-)^{-1} \gamma(\text{Br}_2^-)^{-1}) \tag{5}
\]

For low SoCs with high concentrations c(HBr) \( > 1.2 \text{ M} \) of HBr, the activity coefficients for \( \gamma(\text{HBr}) > 1 \) are available (calculated with data from \[71\]). Increasingly higher activities \( a(\text{HBr}) = c(\text{HBr}) \gamma(\text{HBr}) \) must be taken into account in the calculation of the redox potential of the positive half cell due to constantly increasing activity coefficients of HBr in the aqueous solution \[71\]. Wlodarczyk et al. \[70\] clearly show that the increasing activity coefficients of HBr for low SoCs leads to a strong decrease in the redox potential of the bromine electrolyte at an electrode according to Nernst compared to a pure concentration-based calculation.

The measured redox potential \( \varphi(\text{Br}_2^-/\text{Br}^-) \) of the pure HBr/Br\(_2\)/H\(_2\)O electrolytes (green curve in Figure 9) increases with an increasing SoC from 0.76 V vs. NHE (at an SoC of 0%) and a maximum value at an SoC of 100% at 1.12 V vs. NHE. For SoC \( \geq 89.4\% \), Br\(_2\) is not completely soluble in the aqueous solution as explained in \[65\]. The aqueous solutions contain less Br\(_2\), and the redox potential flattens following the Nernst equation (Equation (5)).
In principle, the redox potential of the aqueous electrolyte solution with the use of [BCA]Br in the presence of Br₂ is lower within the entire SoC range compared to BCA-free HBr/Br₂/H₂O electrolytes (Figure 9). The transfer of Br₂ into the fused salt reduces the bromine concentration in the aqueous solution, while at the same time the HBr concentrations stay nearly constant in comparison to BCA-free HBr/Br₂/H₂O electrolytes. Hence, according to Equation (4), the redox potential of [BCA]Br electrolytes is slightly lower when compared to the BCA-free HBr/Br₂/H₂O electrolytes. For SoCs < 50%, the solubility of the respective [BCA]* cation is responsible for the trend observed in Figure 9. In accordance with lower bromine concentrations in the electrolyte for increasing the n-alkyl side chain length (Figure 6), lower redox potentials are observed with the BCA electrolytes with larger n-alkyl chain. For SoCs > 50%, the difference between the potentials of the pure HBr/Br₂/H₂O electrolyte and the electrolytes containing BCA becomes increasingly larger. In global terms, the Br₂ concentration increases strongly, which can be seen in the potential of the BCA-free HBr/Br₂/H₂O electrolyte, but at the same time the majority of the Br₂ in the solutions with BCAs is transferred to the fused salt. The redox potentials of the electrolytes with BCAs increase at a slower rate with an increasing SoC.

The use of [BCA]Br leads to an average reduction in the redox potential of the positive half cell and the cell voltage ΔE by 32 ≤ ΔE ≤ 114 mV for all BCAs and over the entire SoC range compared to pure HBr/Br₂/H₂O electrolytes. The performance of the cell is reduced by this effect, while it is already lowered by high activity coefficients of HBr [70,71]. Fabjan et al. [8] described for Zn/Br₂-RFB electrolytes that solutions of [C2MM]Br and [C2MP]Br reduce the concentration of Br₂ in the aqueous phase to approximately 0.01 M, and the redox potential is characteristically reduced by 50 to 70 mV. This applies to our electrolytes and both BCA series in Figure 9 for the range 30 ≤ SoC ≤ 70%. A lower cell voltage due to the use of BCAs must be accepted as a trade-off for limiting the Br₂ concentration in the aqueous solution.

Figure 9. Redox potentials of the redox couple Br⁻/Br₂ in aqueous posolyte vs. NHE as a function of its state of charge SoC measured on a glassy carbon electrode without [BCA]Br and with (a) 1-alkylpyridine-1-ium bromides and (b) 1-alkyl-3-methylimidazole-1-ium bromides as BCAs with different alkyl side chain groups in the N-position of the BCA at \( \theta = 23 \pm 1 \) °C. The orange line shows the simulated redox potential following the Nernst equation and is based on global concentrations of HBr and Br₂. The measured redox potential of BCA-free HBr/Br₂ electrolytes is shown with green dots for comparison and was first published in [70]. Values of redox potentials of the electrolytes containing BCAs are printed in the SM of this study in Table S8.
2.8. Interpretation of the Study and Selection of a Promising BCA for Application in Cell Tests

In the scope of this research, 38 cost-effective BCAs have been investigated and evaluated on the basis of their solubility, Br₂ binding strength, ionic conductivity and equilibrium potential for their possible application as electrolytes on H₂/Br₂-RFBs. In the following, an elimination procedure is applied to select the most suitable BCA for battery operation. The BCAs are excluded on the basis of the results obtained in the previous sections. To start, based on their solubility, [CTA]Br, [TOA]Cl and [TOA]Br do not dissolve in a polar HBr/H₂O solution due to their long nonpolar alkyl groups. In addition, other BCAs such as [HPy]Br, [TEA]Br, [C2MPip]Br, [C2MP]Br, [C4MM]Br, [HMIm]Br, [C2MM]Br, [MTA]Br, [TBA]Br, [C24MPy]Br and [C44MPy]Br form solid crystals in the aqueous solution instead of a liquid fused salt when they come into contact with polybromides. Hence, none of these BCAs can be used in the cell as the spontaneous formation of crystals in the cell, tubing and pumps can lead to operational problems.

Since the H₂/Br₂ RFB is intended to be operated at high current densities due to the good reaction kinetics of both electrode reactions, a high mass flow of Br₂ into the cell during the discharge reaction is necessary, making high concentrations of Br₂ desirable. At the same time, it is the task of the BCAs to reduce the bromine concentration in the aqueous phase in order to increase safety. All BCAs that form a liquid fused salt phase and have long alkyl side chains such as n-butyl or n-hexyl have a bromine binding strength of >95 mol%. Additionally, all BCAs with a liquid fused salt phase and ethyl or n-propyl as a side chain, with the exception of [C2Py]Br, have a bromine binding strength of >90 mol%. These BCAs do not leave sufficient Br₂ in the aqueous phase of the electrolyte to achieve acceptable current densities. [HMP]Br, [H4MPy]Br, [H3MPy]Br and [HMPip]Br show only a bromine binding strength of 70 to 80 mol% at an SoC of 33% with concentrations between 0.22 to 0.33 M, while it is expected that for higher SoC values, larger Br₂ concentrations than 0.5 M are available. The essential safety might not be achieved at these concentrations.

It is therefore proposed to investigate 1-ethylpyridin-1-ium bromide [C2Py]Br as BCA in further studies in the RFB cell. From the seven BCAs investigated over the entire SoC range, [C2Py]Br exhibits the most advantages. Even though there is a reduction in the electrolyte conductivity and the redox potential when compared to the BCA-free one, the concentrations of Br₂ in the aqueous solution are higher compared to solutions with [C4Py]Br and [C6Py]Br. In the [C2Py]Br, electrolytes at least 89 mol% Br₂ remain in the fused salt phase, yet the electrolytes still maintain large Br₂ concentrations, between 0.12 M to 0.26 M, in the aqueous solution. The availability of Br₂ in the liquid electrolyte phase turned out to be a decisive factor in our selection, making this compound clearly stand out over all the other investigated BCAs. The initial cycling tests and investigation of the [C2Py]⁺ cations on the cell performance during the charge and discharge operation are shown in parallel in [72].

3. Materials and Methods

A detailed description of the testing methods is given in the SM (Section 1) for this article to enable reproducibility of the measurements. An abbreviated form is reproduced here.

3.1. Reagents

Reagents used for preparation of bromine electrolyte samples in this work are hydrobromic acid, bromine and distilled water. Chemicals for the synthesis of BCAs are named in the Supplementary Materials (SM) including suppliers and purities.

3.2. Synthesis of Bromine Complexing Agents

The investigated BCAs are listed in Table 1, including chemical structure, name and abbreviation. Non-commercially available N-alkylated quaternary ammonium bromide salts based on pyridine ([C2Py]Br, [C4Py]Br, [C6Py]Br, [C6Py]Cl), on 3-picoline ([C23MPy]Br, [C43MPy]Br, [C63MPy]Br), on 4-picoline ([C24MPy]Br, [C44MPy]Br, [C64MPy]Br), on 1-methylimidazole ([C2MIm]Br, [C3MIm]Br, [C4MIm]Br, [C6MIm]Br, [C6MIm]Cl), on 1-
methylpiperidine ([C2MPip]Br, [C4MPip]Br, [C6MPip]Br,) and on 1-methylpyrrolidine ([C6MP]Br) are synthesized by alkylation reaction [45,73] according to Dzyuba et al. [74]. Some BCAs are available as golden liquids at reaction temperatures between 40 °C and 100 °C, while some precipitate as golden or white solids. \(^1\)H NMR and \(^{13}\)C NMR spectra of the synthesized substances are presented in SM and are consistent with NMR shifts found in the literature [45–49]. Protonated forms of 4-picoline, 3-picoline, 1-methylpiperidine, pyridine, 1-methylimidazole and 1-methylpyrrolidine ([H4MPy]Br, [H3MPy]Br, [HMP]Br, [HMPip]Br, [HPy]Br, [HMIm]Br) are prepared by dissolving the heterocyclic ammonium compound with a calculated excess of HBr in the electrolyte solution.

3.3. Electrolyte Formulation

For each BCA, a sample of 10 mL is mixed in aqueous solution containing 5.47 M HBr, 1.11 M [BCA]Br or [BCA]Cl. HBr 48 wt% and H\(_2\)O are added volumetrically. Solubility of [BCA]Br or [BCA]Cl is evaluated. To solutions, in which the BCA salt is dissolved, Br\(_2\) with a mass of 1.77 g (1.11 M Br\(_2\)) is added gravimetrically. All mixtures correspond to an electrolyte mixture for an SoC of 33%.

For selected BCAs [C2MM]Br, [C2MP]Br, [C2Py]Br, [C4Py]Br, [C6Py]Br, [C2MIm]Br, [C3MIm]Br, [C4MIm]Br and [C6MIm]Br, electrolyte properties are investigated ex situ within the entire SoC range. The SoC range is defined for SoC 0% with 7.7 M HBr, 1.11 M [BCA]Br and 0 M Br\(_2\) and for an SoC of 100% with 1 M HBr, 1.11 M [BCA]Br and 3.35 M Br\(_2\). The SoC definition is derived from another study [65]. Samples with a volume of 30 mL in a test series with SoC steps of 10%, and additionally for an SoC of 33% and 66%, are mixed.

3.4. Stability of Fused Salt Electrolyte

The stability of the fused salt is checked by visual inspection. The criteria for qualification of the liquid range of a fused salt are the absence of visible orange or red crystals in the fused salt.

3.5. Concentration of [BCA]+ Cations in Aqueous Electrolytes

Concentrations of 1-alkylpyridin-1-ium cations and 1-alkyl-3-methylimidazol-1-ium cations in the aqueous electrolyte phase are determined by Raman spectroscopy over the entire SoC range. Both cations show characteristic strong peaks at distinctive Raman shifts especially due to their aromaticity [64,75,76]. Characteristic Raman shifts of peaks of the BCAs are shown in Table 2. Peak areas at this Raman shifts are determined and compared with peak areas at known [BCA]+ concentrations (an SoC of 0% and 1.11 M [BCA]Br). [BCA]+(aq) concentrations as a function of the SoC are therefore calculated (ESI).

3.6. Br\(_2\) Concentration in Aqueous Electrolyte Phase

Br\(_2\) binding strength of the BCA or solubility of the [BCA]Br\(_{2n+1}\) complex is investigated by measurement of Br\(_2\) concentration in aqueous electrolyte phase. Concentrations of Br\(_2\) are investigated by linear chronoamperometry at a rotating disk electrode. At a rotation speed of \(\omega = 1000\) rpm on vitreous carbon electrode, linear sweep scans between 0.8 V and −0.5 V vs. an Ag/AgCl/KCl(sat.) reference electrode and a scan rate of −40 mV s\(^{-1}\) are performed for all samples. Constant reduction currents result for half-cell potentials < −0.1 V vs. Ag/AgCl/KCl(sat.), which represent mass transport limitation by Br\(_2\) from the bulk phase [77,78]. According to Levich [77–80], reduction currents are directly proportional to the c(Br\(_2\)) of the bulk solution. Concentrations c(Br\(_2\)(aq)) in M are calculated from the limiting currents I\(_{lim}\) (in mA) shown in Equation (6):

\[
c(Br_2(aq)) = \frac{0.00342466 \text{ mol L}^{-1} \text{ mA}^{-1} I_{lim}}
\]

A linear calibration curve with known bromine concentrations is investigated before at same conditions. Br\(_2\) concentrations are measured for the aqueous electrolyte phase of 35 samples with different BCAs chosen at an SoC of 33% and the series of electrolyte
samples with [C2Py]Br, [C4Py]Br, [C6Py]Br, [C2MIm]Br, [C3MIm]Br, [C4MIm]Br and [C6MIm]Br at 13 different SoCs.

3.7. Polybromide Determination, Polybromide Distribution and [BCA]$^+$ Concentration Determination by Raman Analysis

By means of Raman spectroscopy on the aqueous electrolyte samples, second phase crystals and fused salt samples, the occurrence of the different polybromides tribromide Br$_3^-$, pentabromide Br$_5^-$ and heptabromide Br$_7^-$ in electrolyte series of 1-alkylpyridin-1-ium cations und 1-alkyl-3-methylimidazol-1-ium cations electrolytes at different SoCs were investigated. Raman shifts for polybromides and bromine were determined between $\tilde{\nu} = 150$ and 320 cm$^{-1}$ as described in the literature [17,56,81,82]. Peak areas of the following stretching oscillations of the polybromides were determined: $\tilde{\nu}$(Br$_3^-$, sym.) = 162–163 cm$^{-1}$ [17,53,56], $\tilde{\nu}$(Br$_3^-$, antisym.) = 190–198 cm$^{-1}$ [17,56,83], $\tilde{\nu}$(Br$_5^-$, antisym.) = 210 cm$^{-1}$ [17,56], $\tilde{\nu}$(Br$_5^-$, sym.) = 250–253 cm$^{-1}$ [17,56,81,83,84], $\tilde{\nu}$(Br$_7^-$, sym.) = 270 cm$^{-1}$ [56,81]. In the literature, pure Br$_2$ shows a strong, single peak at $\tilde{\nu}$(Br$_2^-$, sym.) = 300–325 cm$^{-1}$ [81,83,84]. Since there are differences in Raman shifts between the aqueous solution and the corresponding fused salt phase, Raman shifts actually used for fitting the individual peaks are shown in Table 2 for the investigation of the aqueous electrolyte phase. Investigation of Br$_2$ distribution in polybromides is explained in detail in the SM and literature [65].

3.8. Electrolytic Conductivities of Aqueous Phase

Electrolytic conductivities of the aqueous electrolytes of 1-alkylpyridin-1-ium BCAs and 1-alkyl-3-methylimidazol-1-ium BCAs electrolyte at various SoC are determined at $\theta = 23 \pm 1$ °C in a conductivity cell by potentiostatic impedance spectroscopy. The cell constant is determined in a 1 M KCl aqueous solution at $\theta = 23 \pm 1$ °C with $\kappa$ (1 M KCl at $\theta = 23$ °C) = 103.9 mS cm$^{-1}$ (calculated from [85]). A detailed description can be found in the ESI.

3.9. Redox Potential of the Electrolytes

The redox potentials of the aqueous electrolyte solution are investigated on a glassy carbon stick electrode vs. an Ag/AgCl/KCl(sat.) reference electrode at $\theta = 23 \pm 1$ °C and corrected to present them vs. a normal hydrogen electrode. A detailed description can be found in the ESI.

4. Conclusions

In this study, tertiary ammonium compounds of pyridine, pyrrolidine, morpholine, 3-methylimidazole, tetraalkylammonium and picolines have been alkylated with $n$-alkyl side chains in the N-position to form organic bromide or chloride salts. In combination with some commercially available organic salts, this resulted in 38 organic quaternary ammonium substances from these six N-containing building blocks. The substances have been investigated ex situ on their properties as BCAs in HBr/Br$_2$-electrolytes: BCA solubility, Br$_2$ binding strength, the stability of the second heavy fused salt phase, Br$_2$ concentration for application, electrolyte conductivity and redox potential. This study has been performed to define their applicability of BCAs to limit the vapor pressure of Br$_2$ in aqueous highly concentrated electrolytes for a high-energy density H$_2$/Br$_2$-RFB with a theoretical capacity of 180 Ah L$^{-1}$/196 Wh L$^{-1}$ (7.7 M HBr).

From the 38 substances evaluated, 15 are not soluble in the electrolyte or form a crystal phase in contact with polybromides. This behavior makes them unsuitable for their application in RFBs. We found that BCAs with short side chains and/or symmetric structure tend to form crystals instead of forming liquid fused salts. An investigation of the [BCA]-polybromide crystal composition showed that all BCA crystals are tribromide salts [BCA]Br$_3$.

All the other BCAs form fused salt phases, while most of them exhibit a strong bromine binding strength (14 substances > 95 mol% and four substances between 90 to 95 mol%); four of the N-position protonated BCAs show insufficient binding strengths (<80 mol%).
The binding strength increases with an increase in alkyl side chain length, yet the structure of the six basic components (pyridine, picoline, etc.) only has minor influence on the bromine binding strength in solutions.

1-alkyl-3-methylimidazol-1-ium and 1-alkylpyridin-1-ium bromides have been investigated within the whole SoC range to compare the influence of side chain alkyl groups on the different parameters evaluated throughout this study:

For a low SoC (<40%), the selected BCAs show different bromine binding strengths. A clear trend has been found as short alkyl side chains lead to lower Br$_2$ binding strength compared to those with long alkyl side chains. For an SoC of ≥40%, the concentrations of Br$_2$ in the electrolyte’s aqueous phase increase, while the binding strength becomes independent of the alkylated side chain. However, to obtain a clear insight on the reason behind this behavior, further investigation on the fused salt phase would be needed which is beyond the scope of this study. On a final note regarding this comparison, it is worth mentioning that the ability to form a fused salt phase differed from that of the [C2Py]Br and the [C2MIm]Br BCAs as we observed that while [C2Py]Br forms a liquid fused salt, the cation of [C2MIm]Br tends to crystallize over a wide SoC range.

The ionic conductivity of the aqueous phase is high for the whole SoC range between 325 and 745 mS cm$^{-1}$ and depends primarily on the available proton concentration and its ionic transport by the Grothuss mechanism. The concentration of [BCA]$^+$ in the aqueous phase can, to a lesser extent, influence the measured ionic conductivities of the electrolytes as their presence can slightly reduce the conductivity or increase it indirectly by sequestration of Br$_2$ from the aqueous phase.

The redox potential of the electrolyte is reduced by 32 to 114 mV due to the use of BCAs compared to BCA-free electrolytes in the whole SoC range. Following the Nernst equation, the decrease in Br$_2$ concentration in the aqueous phase leads to reduced redox potentials of the half-cell reaction. This is certainly a trade-off that must be accepted as it is a direct consequence of the ambition to lower bromine vapor pressure by limiting Br$_2$ concentration in aqueous solutions.

Based on the results of fused salt stability over the SOC range, electrolyte conductivity, the trend of the redox potential and especially the concentrations of Br$_2$ remaining in the aqueous electrolyte phase, one BCA was selected from 38 BCAs and needs to be investigated in detail and applied in the positive half cell of the H$_2$/Br$_2$-RFB cell: 1-ethylpyridin-1-ium bromide ([C2Py]Br).

**Supplementary Materials:** The following are available online: Detailed description of the measurement methods, synthesis of BCAs, manufacturers’ data and chemical purities (Chapter S1), as well as $^1$H NMR and $^{13}$C NMR interpretation for synthesized BCAs (Chapter S2), tabulation of measured values for Figures within the main manuscript (Table S1–S8), Raman spectra of solutions (Figures S3–S9), Raman spectra of [BCA]Br$_3$ crystals (Figure S2) and concentrations of Br$_2$ in aqueous solution for all 35 BCAs investigated (Figure S1).

**Author Contributions:** Conceptualization, M.K.; methodology, M.K.; synthesis, E.M. and P.A.L.T.; validation, M.K.; formal analysis, M.K., P.A.L.T. and P.F.; investigation, M.K., P.A.L.T. and E.M.; resources, P.F., J.T. and M.K.; data curation, M.K. and P.A.L.T.; writing—original draft preparation, M.K.; writing—review and editing, M.K., P.A.L.T., P.F. and J.T.; visualization, M.K.; supervision, M.K. and P.F.; project administration, M.K.; funding acquisition, M.K., P.F. and J.T. All authors have read and agreed to the published version of the manuscript.

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