Redox-Switchable Chalcogen Bonding for Anion Recognition and Sensing

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ABSTRACT: Inspired by the success of its related sigma-hole congener halogen bonding (XB), chalcogen bonding (ChB) is emerging as a powerful noncovalent interaction with a plethora of applications in supramolecular chemistry and beyond. Despite its increasing importance, the judicious modulation of ChB donor strength remains a formidable challenge. Herein, we present, for the first time, the reversible and large-scale modulation of ChB potency by electrochemical redox control. This is exemplified by both the switching-ON of anion recognition via ChB oxidative activation of a novel bis(ferrocenyltellurotriazole) anion host and switching-OFF reductive ChB deactivation of anion binding potency with a telluroviologen receptor. The direct linking of the redox-active center and ChB receptor donor sites enables strong coupling, which is reflected by up to a remarkable 3 orders of magnitude modulation of anion binding strength. This is demonstrated through large voltammetric perturbations of the respective receptor ferrocene and viologen redox couples, enabling, for the first time, ChB-mediated electrochemical anion sensing. The sensors not only display significant anion-binding-induced electrochemical responses in competitive aqueous-organic solvent systems but can compete with, or even outperform similar, highly potent XB and HB sensors. These observations serve to highlight a unique (redox) tunability of ChB and pave the way for further exploration of the reversible (redox) modulation of ChB in a wide range of applications, including anion sensors as well as molecular switches and machines.

INTRODUCTION

Sigma-hole interactions, in particular halogen bonding (XB) and more recently chalcogen bonding (ChB), defined as the attractive interaction between Lewis bases and an electron-deficient region of a group 17 and group 16 atom, respectively, have emerged as highly potent and versatile noncovalent interactions.1 Their utility is becoming well-established across diverse fields, including crystal engineering,2,3 materials chemistry,4–9 organocatalysis,10–15 and anion supramolecular chemistry.16–20 The latter includes anion recognition,21–24 transport,25–29 and sensing,28,30–39 wherein a notably enhanced performance of sigma-hole-based anion receptors, in particular XB hosts, in comparison to their traditional hydrogen bonding (HB) analogues is well-documented. This includes enhanced anion binding affinities, selectivities, transport efficacies, and sensory performances, attributable to the more stringent linearity requirements of the electron-deficient XB/ChB donor atom—anion interaction as well as their unique electronic properties and tunability.19,40–42

Indeed, recent reports have demonstrated that the ChB donor potency is highly sensitive to the local electronic environment and can be modulated through either covalent substituent variation24,26,42,43 or heteroditopic ion-pair recognition.44 In the context of anion recognition, this tunable ChB donor potency provides an attractive strategy to generate stimuli-responsive host systems. For example, Gabbai and co-workers have recently reported the notable increased potency of Te(IV) cations as ChB anion receptors and transporters in comparison to their parent, neutral Te(II) analogues.26 This was achieved by the oxidative methylation of diaryltellurides, a powerful, but irreversible, means of tuning the ChB donor properties.

Conceivably, control of local ChB donor potency could also be achieved by coupling of a chalcogen center to a redox-active moiety, wherein oxidation or reduction reversibly switches-ON or -OFF chalcogen centered electrophilicity.

While thus far unprecedented for ChB, the reversible (electrochemical) redox control over XB donor strength is already established, in particular with XB ferrocene and TTF receptor-based systems.32,33,43–49 This is saliently reflected in an associated cathodic perturbation of the redox potential of the electroactive XB receptor in the presence of Lewis bases, thereby presenting a simple, yet powerful, means of electro-
chemical anion detection, as increasingly exploited in sensors.32,36,37,39

Given the compelling evidence for the tunable nature of ChB interactions, it was envisaged that the construction of redox-responsive ChB receptor systems would facilitate a potentially powerful means to electrochemically modulate sigma-hole donor potency with high degrees of ON–OFF state fidelity. To explore this concept, we investigated two redox-active chalcogen-containing anion receptor motifs, namely, chalcogenoviologens and a bis(ferrocenyltellurotriazole), wherein the ChB donor potency can be reversibly switched-OFF or -ON by redox control (Figure 1).

In the case of the natively dicationic chalcogenoviologens, we demonstrate a complete deactivation of ChB sigma-hole donor strength and associated anion binding affinity upon electrochemical reduction. Conversely, an activation of anion binding potency of up to 3-orders of magnitude is achieved by oxidation of the neutral ChB bis(ferrocenyltellurotriazole) host system. Not only does this provide a powerful and unprecedented means of reversible control over ChB donor strength, but also enables highly sensitive voltammetric anion detection. This work thereby constitutes a very rare example of ChB-mediated sensing28,38 and the first exploitation of ChB in electrochemical sensing.

### RESULTS AND DISCUSSION

#### Synthesis of Redox-Active ChB Anion Receptors

The synthesis of the chalcogenoviologens 1.Te2+ and 1.Se2+ was achieved via reaction of the chalcogen-functionalized bipyridine precursors50 with benzyl bromide, followed by salt metathesis with sodium tetrakis(3,5-bis(trifluoromethyl)-phenyl)borate (NaBArF4), to afford the target receptors 1.Te2+ and 1.Se2+, in yields of 61% and 57%, respectively (Scheme 1). In addition, for the purposes of delineating the role of nonspecific electrostatic interactions in the anion sensing behavior of the viologen systems, an unfunctionalized derivative 1.H2+ was also prepared.

The novel ChB bis(ferrocene-telluro-triazole) receptor 2.Te was constructed by direct appendage of ferrocene (Fc) redox reporters onto an established bis(tellurotriazole) anion receptive scaffold (Scheme 2).25,42,44 This was achieved by reaction of pyridine bis(silveracetylide)42 with in situ generated ferrocenyl tellurobromide, obtained from reaction of diferrocenyl ditelluride51 with bromine, according 3,5-pyridine bis(ferrocenylalkyne) 4. This compound was immediately subjected to copper(I)-catalyzed azide-alkyne cycloaddition (CuAAC)22 with two equivalents of azido pentafluorobenzene52 affording the title compound 2.Te in 54% yield over two steps.53 Further detail and full compound characterization of all receptors by 1H and 13C NMR as well as ESI-MS, can be found in the Supporting Information.

| Chemicals | Reactions | Products |
|-----------|-----------|----------|
| 1. Te2+   | 1. Te2+   | 1. Se2+  |
| 1. H2+    | 1. H2+    | 1. H2+   |

#### Switchable Anion Recognition and Sensing via Voltammetry

Figure 1. Illustrative overview of the redox control of chalcogen bonding (ChB) potency. We demonstrate both reversible electrochemical reductive deactivation of ChB in telluroviologens as well as oxidative activation of ChB in ferrocenyltellurotriazole receptors.

Scheme 1. Synthetic Route for Dicationic Redox-Active Chalcogenoviologen Hosts 1.Te2+ and 1.Se2+ as well as the Control Proto-Congener 1.H2+.
Having established the ChB potency of the dicationic 1.Te\(^{2+}\), attention turned to the assessment of the anion binding capability of the neutrally native bis(ferrocenyltellurotriazole) 2.Te receptor in the switch-OFF state.\(^{53}\) Analogous \(^1\)H NMR anion binding studies revealed, as expected with the neutral receptor, that the ChB potency was strongly diminished.

Indeed, no measurable \(^1\)H NMR perturbations for a range of halide or o xoanion guests were observed even in the much less competitive d\(_6\)-acetone. Only the addition of H\(_2\)PO\(_4\) induced significant chemical shift perturbations of the internal pyridine proton of 2.Te, wherein analysis of the binding isotherm determined a weak 1:1 host–guest stoichiometric association constant of 111 M\(^{-1}\) (Figure S12 and Table S1). Structurally related pyridine bis(tellurotriazole) ChB receptors typically display stronger anion binding in this solvent,\(^{32}\) indicating a somewhat decreased neutral ChB potency of 2.Te, presumably arising due to a combination of an increased electron density at the chalcogen center and steric constraints from the large and comparably electron-rich ferrocene substituents.

These findings support a very weak ChB sigma-hole donor capability of the neutral 2.Te receptor (ChB OFF) in acetone and acetonitrile, while the dicationic 1.Te\(^{2+}/\)Se\(^{2-}\) receptor systems expectedly display much improved native anion binding capabilities even in competitive aqueous media (CD\(_3\)CN/D\(_2\)O 9:1; ChB ON), providing a first indication of the strong dependence of ChB sigma-hole donor strength on the respective receptor’s redox state.

The significant ChB anion recognition potency of 1.Te\(^{3+}\) was further supported by UV–vis anion binding studies in the same solvent system as employed for \(^1\)H NMR studies. In CH\(_3\)CN/H\(_2\)O 9:1, 1.Te\(^{3+}\) displays significant absorbance changes upon titration with halide anions, whereby, in the presence of iodide, the absorbance band at \(\lambda_{\text{max}} = 481\) nm shifts bathochromically with appearance of an isosbestic point at 505 nm (Figures 3 and S13–S15). These perturbations are large enough to be detectable by the naked eye (change from

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**Table 1.** Anion Association Constants \(K (\text{M}^{-1})\) of 1.Te\(^{2+}/\)Se\(^{2-}/\)H\(^{2+}\) as Determined by \(^1\)H NMR Titrations in CD\(_3\)CN/ D\(_2\)O 9:1

| Anion     | 1.Te\(^{2+}\) | 1.Se\(^{2-}\) | 1.H\(^{2+}\) |
|-----------|---------------|---------------|-------------|
| Cl\(^-\)  | 181 ± 1       | 127 ± 1       |             |
| Br\(^-\)  | 182 ± 2       | 139 ± 1       |             |
| I\(^-\)   | 179 ± 3       | 130 ± 1       |             |

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**Figure 2.** (A) Proposed anion binding mode of 1.Te\(^{2+}\) with iodide as a representative halide anion. (B) Representative \(^1\)H NMR chemical shift perturbations of protons \(a–c\) of 1.Te\(^{2+}\) upon titration with TBA-iodide in CD\(_3\)CN/D\(_2\)O 9:1. (C) Corresponding halide anion binding isotherms.
yellow to red) and present a rare example of ChB-mediated optical anion sensing.7,24,28,56,57 Pleasingly, the 1:1 host−guest stoichiometric binding constants obtained by global fitting of the UV−vis titration data are in good agreement with those obtained by 1H NMR titrations; Br− is bound strongest with K = 873 M−1, while binding of Cl− and I− is slightly weaker with 690 and 627 M−1, respectively (Figures S13−S15).

Furthermore, 1.Se2+ displays only minor UV−vis perturbations toward halide anions (Figure S16), too small to accurately quantify. Similarly, and again in good agreement with 1H NMR studies, the proto analogue 1.H2+ is neither colored nor optically responsive to anions (Figure S17).

**Electrochemical Characterization of Receptors.** The electrochemical characterization of all the ChB redox-active receptors was carried out by cyclic voltammetry (CV) and square-wave voltammetry (SWV) in the presence of 100 mM TBAPF6 as the supporting electrolyte. For solubility reasons, 1.Te2+/Se2+/H2+ were only studied in the same solvent system of CH3CN/H2O 9:1 as employed for 1H NMR and UV−vis studies. As shown in Figure 4A, all the viologen receptors displayed two well-defined, one-electron reductive couples (V2+/V+ and V+/V) at similar, moderately cathodic potentials (Table S2), in good agreement with related previous studies.50,58 Both couples display a high degree of reversibility as evidenced by a close-to-unity ratio of anodic and cathodic peak currents (i/p) and excellent linearity of these currents with the square root of the scan rate (Figures S18−S20), confirming a diffusion-controlled process. These observations highlight that introduction of the chalcogen donor atom into the viologen scaffold does not significantly affect its electrochemical properties. For example, the half-wave potentials E1/2 of all three viologens are very similar and lie between −0.720 and −0.770 V for the first reductive couple (V2+/V+) and between −1.160 and −1.210 V for the second reductive couple (V+/V) (Table S2).

The improved solubility of neutral 2.Te in comparison to dicatonic 1.Te2+ enabled electrochemical studies across a wider range of solvent systems. In all tested solvents, DCM, acetone, CH3CN (Figure 4B), and MeOH containing 100 mM TBAPF6 supporting electrolyte, 2.Te displayed a single, well-defined, reversible, diffusion-controlled redox couple (see also Figures S21−24). This suggests that both Fc motifs undergo one-electron oxidation simultaneously and independently of one another, generating the dicationic 2.Te2+. Only in the presence of the much larger, less-coordinating BARF4− anion as the electrolyte in DCM, was electronic communication between the Fc motifs observed, as evidenced by appearance of two separate redox couples with a peak separation of ≈90 mV (Figures S25 and S26).59

As a result of the strongly electron-withdrawing perfluorobenzene appendages, the receptor’s half-wave potential E1/2 = 0.218 V in CH3CN is significantly more positive than that of parent Fc (0 V) (Table S3).
Significant problems precluded quantitative binding data being determined. As shown in Figure 5, the concentration and ionic strength.

Table 2. Maximum Cathodic Voltammetric Shifts \( \Delta E_{\text{max}} \) of 1,10-Te2+/Se2+/H2+/V2+, in the Presence of 50 mM Anion, and \( K_{\text{ox}} = K_{1,\text{Te2+/Se2+}} \) as Determined by Fitting of the Voltammetric Binding Isotherms According to Eq 1, for the First Reductive Couple of Viologen Hosts toward Various Anions in Degassed CH3CN/H2O 9:1 and 100 mM TBAPF6

| Anion | 1,10-Te2+ | 1,10-Se2+ | 1,10-H2+ |
|-------|-----------|-----------|----------|
| Cl−   | −57       | 156       | 9.3      |
| Br−   | −61       | 205       | 10.8     |
| I−    | −49       | 107       | 6.8      |
| HSO4− | −36       | 77        | 4.1      |
| NO3−  | −22       | n/a       | n/a      |

| Anion | 1,10-Te2+ | 1,10-Se2+ | 1,10-H2+ |
|-------|-----------|-----------|----------|
| Cl−   | 9.3      | 156       | 10.8     |
| Br−   | 6.8      | 205       | 10.8     |
| I−    | n/a      | 107       | 6.8      |
| HSO4− | n/a      | 77        | 4.1      |
| NO3−  | n/a      | n/a       | n/a      |

Reductive Switch-OFF Deactivation of ChB in Telluroviologen. In order to assess the ChB donor strength of the redox-active receptors, we conducted voltammetric anion sensing studies. This provides a particularly convenient and straight-forward means of assessing the ChB potency, as the magnitude of voltammetric perturbation in the presence of the analyte is directly dependent on its binding strength to the different receptor oxidation states, as discussed in more detail below. As shown in Figure S2, the first reductive couple (V2+/V+) of 1,10-Te2+ displayed significant, continuous cathodic voltammetric shifts toward all tested anions in CH3CN/H2O 9:1, with a notable preference for the halides according to eq S2, see the Supporting Information for further details. / not conducted. n/a = not meaningful due to small response.

\[ \Delta E_{\text{max}} = \frac{RT}{nF} \ln \left( \frac{1 + K_{\text{ox}}[A]}{1 + K_{\text{red}}[A]} \right) \]

As shown in Table 2, with the halides and bisulfate is moderate but significant (77−205 M−1), and the largest for Br−. Pleasingly, this is in good agreement with 1H NMR binding studies where the strongest Br− binding was observed (Table 1). The somewhat smaller magnitude of \( K_{\text{ox}} \) determined voltammetrically in comparison to the analogous 1H NMR binding constants is in line with similar prior reports and most likely arises from electrolyte effects as detailed elsewhere.
In all cases, \( K_{1.\text{Te}^{2+}/\text{Se}^{+}} \), that is binding to the monocationic \( 1.\text{Te}^{+}/\text{Se}^{+} \), is negligible (\( K_{1.\text{Te}^{2+}/\text{Se}^{+}} \approx 0 \) M\(^{-1} \)), indicative of complete ChB switch-OFF and anion decomplexation from the \( 1.\text{Te}^{2+} \) host upon monoreduction to \( 1.\text{Te}^{+} \). This is in excellent agreement with negligible voltammetric perturbations observed for the second reductive couple; i.e., in this solvent system, monoreduction is sufficient to completely deactivate ChB-mediated anion binding, such that a further reduction to \( 1.\text{Te} \) has no additional impact on anion binding strength.\(^61\)

We further quantified the redox switching magnitude of \( 1.\text{Te}^{2+} \) by calculating the \( \text{BEF} = K_{1.\text{Te}^{2+}/1.\text{Te}^{+}} \), according to a simplified model (\( \text{BEF} = 10^{\Delta E_{\text{max}}/0.059} \) (eq S2), see Table 2, refs.\(^{33,39} \) and SI Section 6 for further details). This confirmed significant ChB binding modulation (i.e., deactivation) by a factor of up to \( \approx 11 \)-fold in the case of bromide.

Importantly, the redox reversibility of both redox couples of the telluroviologen host is generally maintained in the presence of anions, highlighting that ChB potency and anion binding can be judiciously reversibly switched OFF by redox control. As discussed in more detail in a later section, the sensory performance of this simple ChB voltammetric anion sensing system and redox-switch compares very favorably with those of a range of more elaborate XB or HB redox active systems (see Figure S25).

**Oxidative Switch-ON Activation of ChB in Ferrocenyltellurotriazole 2.\text{Te}.** Having established the general utility of (reductive) redox-controlled ChB in simple telluroviologens, our attention focused on a further elaboration of this concept in the more advanced anion receptor scaffold 2.\text{Te}. The ChB potency of this natively neutral receptor can be reversibly switched-ON by redox control of the ferrocene/ferrocenium (Fc/F\(^{+} \)) couple, a more common, and arguably more useful approach for anion recognition and sensing, circumventing the isolation of charged compounds.\(^4\)

To assess this capability, voltammetric sensing studies with halides and oxoanions were undertaken. Due to the improved solubility of neutral 2.\text{Te}, these were conducted in a range of solvent systems. In CH\(_3\)CN, 2.\text{Te} voltammetrically responded to all tested anions (H\(_2\)PO\(_4\), HSO\(_4\), NO\(_3\), Cl\(^-\), and Br\(^-\)) via significant, continuous cathodic shifts, as shown in Figure 6A.\(^66\)

In all cases, a well-defined, monotonic, and reproducible response (Figures S28–S30) of the sensor was observed with the following response magnitude selectivity, defined by the maximum cathodic voltammetric shift \( \Delta E_{\text{max}} \) of H\(_2\)PO\(_4\) > Cl\(^-\) > Br\(^-\) > HSO\(_4\) > NO\(_3\), with an impressively large perturbation magnitude toward H\(_2\)PO\(_4\) of \(-217 \) mV (Table 3). Notably, as discussed above, \(^1\)H NMR anion titration investigations demonstrated that the neutral receptor 2.\text{Te} displayed no detectable binding of most of these anions in acetone. This confirms that, as expected, receptor oxidation to 2.\text{Te}\(^{2+}\) significantly enhances ChB potency and enables voltammetric anion sensing. Quantified analysis of the voltammetric response isotherms determined impressively large binding constants in the oxidized state of \( K_{2.\text{Te}^{2+}} \) of up to 174,000 M\(^{-1} \) for H\(_2\)PO\(_4\) and for the halides with binding constants of 4050 and 10,200 M\(^{-1} \) for Br\(^-\) and Cl\(^-\), respectively (Table 3). The H\(_2\)PO\(_4\) preference in this nonaqueous solvent is, again, in good agreement with the aforementioned \(^1\)H NMR studies in which only this anion induced measurable, but weak binding to 2.\text{Te} in d\(_6\)-acetone.

The large-magnitude anion recognition-induced voltammetric perturbations of 2.\text{Te} toward Cl\(^-\), Br\(^-\), and H\(_2\)PO\(_4\) correspond to an impressively substantial binding switch-ON approach.
upon oxidation with BEFs of 458, 180, and 4760, respectively (Table 3). This reversible ChB donor modulation, by up to over 3-orders of magnitude, is notably greater than that attainable by cooperative ion-pair recognition in related heteroditopic ChB receptors.44 We believe that this anion binding enhancement of 2. Te2+ is predominantly driven by specific ChB interactions (and not by electrostatics), as inferred from the comparisons of 1. Te2+/Se2+/H2+ (vide supra) as well as comparisons with similar bis(ferrocene-(iodo)triazole) XB and HB systems (vide infra).53 However, electrostatic interactions between the anion guest species and the receptor’s Fc’ groups may also play a role. The delineation of these contributions will be the subject of future studies.

Unsurprisingly, the introduction of 5 or 10% water into the CH3CN solvent system is associated with moderately diminished response magnitudes, Kox binding constants, and BEFs (Figures 6B, C, 7, and Table 3).59 Nevertheless, significant shift responses and switching magnitudes are retained in these competitive, aqueous solvent systems, a particularly impressive accomplishment when considering that the native neutral 2. Te receptor exhibits no significant anion affinity in less-competitive pure organic solvent media. For example, even in CH3CN/H2O 9:1, significant cathodic shifts of up to ~41 mV toward Br− are observed, corresponding to a BEF = 5.2.

Of further note, the response selectivity of the sensor is altered in the presence of water (Figure 7). The response toward the halides and dihydrogen phosphate, in particular, is attenuated upon introduction of 5% water, most likely a reflection of their respective large hydration enthalpies. This also results in a somewhat altered selectivity pattern in CH3CN/H2O 9:1 of ΔEmax: H2PO4− > Br− > Cl− ≈ HSO4− > NO3−. Upon increasing the water content to 10%, the response and binding magnitudes further decrease, however not as sharply. Again, the most strongly affected anion is H2PO4−, which now induces a similar response to Cl−, both of which are lower than that of Br− (Br− > H2PO4− ≈ Cl− > HSO4− ≈ NO3−). The observation that the halide response is more significantly retained than that of the oxoanions hints at a unique halide selectivity associated with ChB recognition, akin to that observed in XB systems.39

Chalcogen Bonding 1. Te2+ and 2. Te Receptors as Potent Redox Controlled ON/OFF Switches and Sensors. In spite of their weak ChB anion binding strength in their reduced states and comparable structural simplicity, both 1. Te2+ and 2. Te are among the most potent voltammetric anion sensors reported to date, as measured by their response magnitudes.39

In fact, to the best of our knowledge, 2. Te significantly outcompetes every reported solution-phase XB voltammetric anion sensor, as well as the vast majority of redox-active HB anion sensors, for the sensing of most anions in CH3CN, in particular chloride and bromide.33,39,46,48,49,60,69–72 This is particularly noteworthy considering, in general, the significantly weaker sigma-hole donor ability of ChB receptors in comparison to XB systems.16,19,44 As shown in a comparison of the sensing performance of 2. Te with related XB and HB bis(ferrocene-(iodo)triazole) systems (see SI Section 7 and Figures S31–S34),39 Br− especially elicits an up to 2.4-fold larger response than its structurally closest XB congener, in both CH3CN and CH3CN/H2O 19:1. Due to its enhanced response magnitudes, 1. Te2+ even outcompetes these XB and HB sensors (recently reported as some of the most potent voltammetric sensors), for the sensing of Cl− and Br− at high concentrations in CH3CN/H2O 9:1.39 These observations support a developing picture that supports a ChB preference for the “softer” halides in comparison to oxoanions (akin to XB), in spite of their lower basicity.

As discussed above, the large response magnitudes of 1. Te2+ and 2. Te are a direct reflection of their large anion binding enhancement factors (BEF = Kox/Kred), that is the relative change in anion binding constants upon reduction/oxidation.35,39 The impressive sensing performance of both 1. Te2+ and 2. Te can thus be attributed to a uniquely potent switching-ON or -OFF of anion binding, respectively, as previously observed in a range of XB voltammetric anion sensors. We ascribe this to the high sensitivity of the ChB donor strength to electronic inductive substituent effects42 as well as the most intimate spatial and electronic coupling of the sigma-hole donor binding site to the redox transducer, afforded by the direct appendage of the Te sigma-hole donor to Fc or viologen. As a result, the redox modulation of the electroactive transducer more strongly influences the ChB sigma-hole depth and thereby anion binding strength, which in turn translates to a large voltammetric response.

The direct appendage of Te ChB donor atoms with optical or redox transducer groups remains a largely unexplored, but promisingly highly potent, design feature of ChB receptors that not only enables sensing with improved sensitivity but can, in the case of the 2. Te scaffold, also “free up” a substituent, such that other functional groups can easily be incorporated into the receptor (here via the triazole component). This is notably not easily achievable in XB or HB systems in which the XB/HB bond donor atom typically carries only one substituent.

As a result of this structural adaptability, combined with the high-fidelity redox-control of ChB potency, as demonstrated with 1. Te2+ and 2. Te, we envision these electroactive ChB centers as ideal motifs for further exploration in ion sensors as well as molecular redox switches and machines.
CONCLUSIONS

We report, for the first time, ChB-mediated electrochemical anion sensing and reversible redox-controlled switch-ON and switch-OFF of ChB donor potency exemplified by 3,5-bisferrocenettellurotriazole pyridine and telluroviologen anion receptors. Despite their structural simplicity and comparably weak anion binding strength in the reduced receptor states, both systems displayed large scale anion binding enhancements, of up to 3 orders of magnitude, in the oxidized state. This is directly reflected in large cathodic voltammetric shifts in the presence of anions, which persist even in the competitive CH3CN/H2O 9:1 solvent system, with a noteworthy selectivity in the presence of halides, in particular bromide. Impressively, the high sensitivity of these sensors compares very favorably with the halides, in particular bromide. Highly electronically tunable and not only pave the way toward the further exploitation of ChB in anion sensor design and redox switches but also present fundamental insights into the nature of ChB interactions.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c02924.

Experimental details and materials, additional data, and figures (PDF)

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