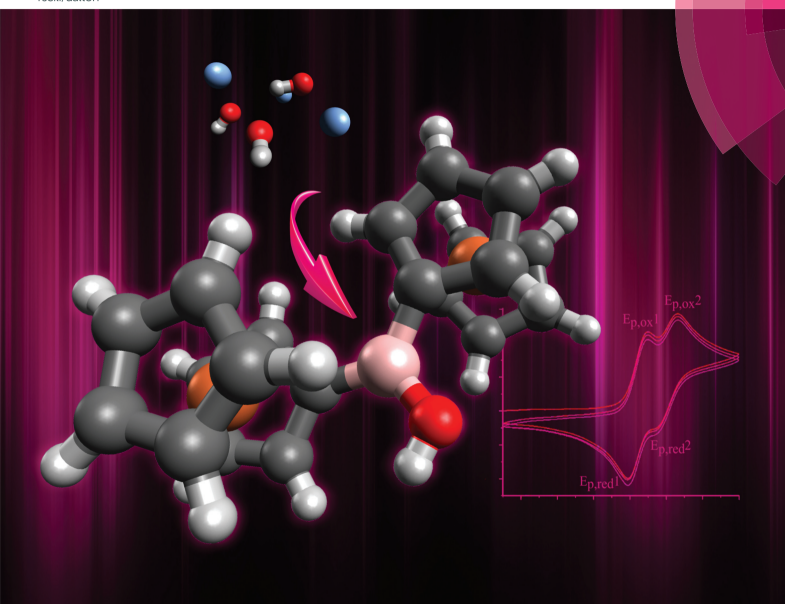
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#### PAPER

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Voltammetric characterisation of diferrocenylborinic acid in organic solution and in aqueous media when immobilised into a titanate nanosheet film†

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Diferrocenylborinic acid (Fc<sub>2</sub>BOH, **1**) has been synthesized in good yield *via* an improved synthetic path. Characterisation by nuclear magnetic resonance (NMR), mass spectrometry (HRMS), infrared spectroscopy (FTIR), X-ray crystallography, and by electrochemical methods reveal two one-electron oxidation processes for the two electronically coupled ferrocenyl moieties. The oxidation of **1** dissolved in organic media is contrasted to the oxidation of **1** in aqueous environments (by incorporation of **1** into a lamellar film of 2D titanate nanosheets on a glassy carbon electrode). Data from cyclic voltammetry and from square wave voltammetry suggest that the bridging boron can bind to nucleophiles (hydroxide, fluoride) upon oxidation of the ferrocenyl groups. A multi-pathway ECE reaction scheme is proposed. Potential applications in sensing are discussed.

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# 1. Introduction

Ferroceneboronic acid as an electrochemically active boronic acid has been used previously in electroanalytical sensing protocols. Paronic acid moieties  $[R^1B(OH)_2]$  can bind nucleophiles and in particular vicinal diols or other polyols And importantly to glucose. From this, a range of sensing applications have been developed, including applications with electrochemical detection. In contrast, the borinic acid moiety, a chemical relative of boronic acid, consists of two alkyl substituents and only one hydroxyl linked to the boron atom  $[R^1(R^2)B-OH]$ . It has been suggested that borinic acids

Organometallic complexes containing boron-based pendants have been proven to offer fertile ground for the exploration of fundamental structure and reactivity issues. 15 Trisferrocenylborane has been prepared and characterised. 16 Also the zwitter-ionic ferricenyl(III)tris(ferrocenyl(III))borate has been obtained and characterised by cyclic voltammetry. 17 Diferrocenylborinic acid (Fc<sub>2</sub>BOH, 1, Fig. 1), the simplest electroactive borinic acid, is an interesting material fundamentally and in conjunction with electrochemical sensing protocols. In contrast to trisferrocenylborane, it is hydrolytically stable and potentially useful as an analytical reagent in aqueous environments. Complex 1 was first prepared and reported by Post et al. in 1970, 18 but its electrochemical properties have not been investigated in further detail. In this report, we describe the electrochemical properties of 1 in organic solution and in aqueous environments.

In order to investigate water-based redox properties, waterinsoluble redox systems can often be immobilised into films at electrode surfaces and immersed into an aqueous solution. In

should have similar binding properties to nucleophilic analytes compared to those observed for boronic acids, <sup>14</sup> but little attention has been paid to borinic acids. Diferrocenylborinic acid is an interesting system to study as it has the additional benefit of providing a two-electron redox system, where not only the formal potential for electron transfer, but also the potential gap between the first and second electron transfer, are accessible parameters for analytical purposes.

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Fig. 1 Synthesis of Fc<sub>2</sub>BOH (1); (a) in toluene, excess of n-propanol, molecular sieve 4 Å, 1 h azeotropic dehydration; (b) in THF under inert atmosphere, 1 mol. eq. of t-BuLi, 0 °C, 15 min; (c) in THF under inert atmosphere, 0 °C, 15 min, H<sub>2</sub>O quenching, further extraction and purification according to experimental details in section 2.3.

this way, metal complexes such as diferrocenylborinic acid 1 can be investigated for potential applications in electroanalysis in aqueous electrolyte environments. A versatile way of immobilising hydrophobic metal complexes at the surface of glassy carbon electrodes can be based on a 2D titanate nanosheet host, 19 which is able to bind the redox active molecule from organic solvents. Nanomaterials such as titanate nanosheets have been widely investigated due to their catalytic, semiconducting, photo- and electrochemical properties and for their advanced, next-generation use nanoelectronics. 20,21 2D titanate nanosheets can be obtained as a colloidal solution by delaminating a layered titanate precursor.<sup>22</sup> In previous studies, it has been shown that such titanate nanosheets can be beneficial as an additive to polymer composite materials to improve their mechanical properties.<sup>23</sup> A report on potential applications of multi-layer titanate nanosheet film deposits in electroanalysis has also appeared.<sup>24</sup> One of the first studies with modified electrodes based on titanate nanosheets with an embedded electrochemical probe (ferroceneboronic acid) was reported recently.<sup>25</sup> It was shown that the ferroceneboronic redox signal in a titanate nanosheet film immersed in aqueous Na2SO4 was responsive for example to the anions in the electrolyte and to fructose binding. We therefore decided to explore the reactivity of diferrocenylborinic acid 1 under similar conditions in aqueous environments and exposed to nucleophiles such as fluoride.

In this study, diferrocenylborinic acid 1 is synthesized via an improved synthetic pathway (see Fig. 1), spectrally and structurally characterised, then utilized in a film modified electrode based on titanate nanosheets on a glassy carbon surface. The type of redox response and the potential for sensing applications are evaluated.

#### 2. **Experimental details**

#### 2.1. Chemical reagents

All chemicals were of reagent grade and were used without further purification unless otherwise noted. Ferrocene was purchased from Fluka, t-BuLi, tetrabutylammonium hexafluorophosphate (TBAH), sodium sulphate, and sodium

fluoride from Sigma Aldrich, ferroceneboronic acid from TCI Europe, and all solvents were purchased from VWR International or Sigma Aldrich. Titanate nanosheet material was synthesized as a colloidal solution (2.56 g dm<sup>-3</sup>) as described previously by Sasaki<sup>26</sup> and by Harito et al.<sup>23</sup> Solutions were prepared under ambient condition in volumetric flasks with ultrapure water from ELGA Purelab Classic system (resistivity of 18.2 M $\Omega$  cm at 22 °C).

#### 2.2. Instrumentation

<sup>1</sup>H and <sup>13</sup>C NMR spectra were recorded on a 300 MHz Bruker Avance spectrometer. 11B NMR spectra were recorded on a 500 MHz Agilent ProPulse spectrometer. All measurements were performed in chloroform- $d_1$  (CDCl<sub>3</sub>) at 303 K. The chemical shifts are reported in ppm on the  $\delta$  scale using signal of deuterated solvent residue as a reference. HRMS spectra were recorded on Agilent 6224 Accurate-Mass TOF LC-MS system with ESI + (MMI) ionisation. FTIR spectra were collected on Bruker Alpha FT-IR spectrometer (Platinum ATR). X-ray intensity data for 1 were collected at 150(2) K on Rigaku Xcalibur, EosS2 using monochromated graphite Mo-K $\alpha$  radiation ( $\lambda$  = 0.71073 Å). A symmetry-related (multi-scan) absorption correction and an analytical numeric absorption correction had been applied. The structure was solved with SHELXT and refined by a full-matrix least-squares procedure based on  $F^2$ (SHELXL-2018/3<sup>27</sup>). All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were placed in calculated positions and refined using a riding model except for the OH hydrogen atoms which had been located in the difference Fourier map and were refined with bond length restraints. Additional programmes used for analysing data and their graphical manipulation included: SHELXle<sup>28</sup> and ORTEP 3 for Windows.<sup>29</sup> The cif file of the structure has been deposited in the Cambridge Structural Database, deposition number CCDC 1894130.†

Spectroelectrochemical data were recorded on an Autolab PGSTAT30 (Metrohm-Autolab, Netherlands) combined with Avantes spectrometer AvaSpec-ULS2048XL-EVO (Avantes, Netherlands) and optically transparent thin-layer spectroelectrochemical (OTTLE) cell with Ag wire as reference electrode and Pt grid as both working and counter electrode, respectively, and 1 mm optical path length. Electrochemical experiments were performed using three-electrode system controlled with a microAutolab III system (Metrohm-Autolab, Netherlands) in voltammetry mode. The scan rate was maintained at 100 mV s<sup>-1</sup> or as otherwise noted. The platinum wire and KCl-saturated calomel electrode (SCE) were used as counter and reference electrodes, respectively. The working electrode was a 3.0 mm-diameter glassy carbon electrode (GCE) alternatively modified with TiO<sub>2</sub> nanosheet film (or in

some cases a 1 mm diameter Pt disk electrode). Working elec-

trodes were polished with alumina slurry (50 nm diameter) and washed with ultrapure water before modification with tita-

nate nanosheet films (see below for details).

#### 2.3. Syntheses

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Ferroceneboronic acid (0.38 g, 1.66 mmol) was dissolved in toluene (10 mL), a few drops of H<sub>2</sub>O and 1-propanol (1 mL) were added and the mixture was stirred with a 4 Å molecular sieve for 30 minutes at room temperature. All volatile solvents were then evaporated under continuous vacuum for 5 minutes to give a red oily intermediate containing mostly ferroceneboronic acid 1-propanol ester. The material was then dissolved in dry THF (3 mL) under inert atmosphere (flushed with N<sub>2</sub>) and stirred on an ice bath for further synthesis. Next, ferrocene (0.8 g, 4.3 mmol) was weighed into the Schlenk flask, flushed with N<sub>2</sub> (inert atmosphere) and dissolved in dry THF (5 mL), maintaining 0 °C on an ice bath. t-BuLi (2.7 mL, 4.2 mmol) was added dropwise to the stirred solution of ferrocene. The formation of FcLi was accompanied by darkening from orange to red colour of solution. After 15 minutes of stirring, the batch of ferroceneboronic acid 1-propanol ester was transferred to the FcLi solution and stirred together for a further 15 minutes. The colour of the solution turned to dark red. Finally, the reaction was quenched with H<sub>2</sub>O (5 mL), brine (3 mL) was added, and the mixture was warmed to room temperature (22 °C). The crude product was extracted with Et<sub>2</sub>O (3  $\times$ 15 mL), organic layers were combined, dried using CaCl2 and evaporated to dryness under reduced pressure to give a dark red solid. This solid was slurried with hexane and placed on silica gel column (2 cm × 20 cm). The unreacted and quenched ferrocene was eluted with hexane or petroleum spirit at 40-60 °C (0.4 g, 2.15 mmol, recovered for further synthesis). The red band containing mainly Fc3B (2) was eluted with benzene (0.16 g, 0.28 mmol, 17% in respect to boron equivalent in reactants; note that Fc3B is easily hydrolysed to give Fc<sub>2</sub>BOH (1) in CDCl<sub>3</sub> after 50 h or directly on the silica gel columns to give Fc<sub>2</sub>BOH). The desired Fc<sub>2</sub>BOH (1) was eluted as last orange-brown band with benzene and as main band slowly eluted with toluene as a brownish/ochre solid (0.25 g, 0.63 mmol, 38% in respect to boron equivalent in reactants).

Note –  $Fc_2BOH$  (small amounts *e.g.* 10 mg of  $Fc_3B/Fc_2BOH$  mixture) can be further purified by a Pasteur pipette flash LC (silica gel/hexane). Impurities stay either on the column or they are eluted with hexane (FcH). The pure  $Fc_2BOH$  is then

slowly eluted with benzene or toluene. The rest of less pure Fc<sub>2</sub>BOH is then eluted with diethylether.

Diferrocenylborinic acid (1). <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 4.15 (s, 10H Cp), 4.53 (2dd, 8H CpB), 4.92 (br s, OH); <sup>13</sup>C NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 68.73 (Cp), 72.41 (CpB), 73.66 (CpB), n.o. (C<sub>ipso</sub>); <sup>11</sup>B NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 46.05; HRMS (CH<sub>3</sub>CN): m/z = 398.0229 [M]<sup>+</sup>; calcd mass C<sub>20</sub>H<sub>19</sub>BFe<sub>2</sub>O = 398.0227; FTIR:  $\nu$  (cm<sup>-1</sup>) = 3596 (m), 3091 (w), 1771 (w), 1650 (w), 1452 (s), 1407 (m), 1375 (s), 1277 (s), 1149 (w), 1102 (s), 1043 (s), 998 (s), 925 (m), 817 (s), 752 (m), 688 (s), 569 (m), 536 (m), 476 (s), 464 (s).

Trisferrocenylborane (2). <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 4.22 (s, 15H Cp), 4.62 (t, 6H CpB), 4.89 (t, 6H CpB); <sup>13</sup>C NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 69.00 (Cp), 72.43 (CpB), 77.15 (CpB), n.o. (C<sub>ipso</sub>); <sup>11</sup>B NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 62.73; HRMS (CH<sub>3</sub>CN): m/z = 566.0248 [M]<sup>+</sup>; calcd mass C<sub>30</sub>H<sub>27</sub>BFe<sub>3</sub> = 566.0256; FTIR:  $\nu$  (cm<sup>-1</sup>) = 3089 (w), 1690 (w), 1427 (s), 1375 (s), 1352 (w), 1250 (s), 1187 (w), 1105 (s), 1052 (s), 999 (s), 814 (s), 759 (m), 697 (m), 485 (s), 468 (s).

Ferroceneboronic acid 1-propanol ester (3). <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 0.94 (t, 3H CH<sub>3</sub>), 1.00 (t, 3H CH<sub>3</sub>), 1.58 (sextet, 2H CH<sub>2</sub>), 1.68 (sextet, 2H CH<sub>2</sub>), 3.61 (m, 2H CH<sub>2</sub>O), 4.00 (m, 2H CH<sub>2</sub>O), 4.13 (s, 5H Cp), 4.37 (t, 2H CpB), 4.42 (t, 2H CpB); <sup>11</sup>B NMR (CDCl<sub>3</sub>):  $\delta$  (ppm) = 30.33.

#### 2.4. Procedure for electrode preparation

An aliquot of 5  $\mu$ L titanate nanosheet colloidal solution (2.56 g dm<sup>-3</sup> in water) was deposited onto the glassy carbon electrode (GCE) surface by drop casting. The solution was dried at ambient temperature forming a transparent thin film of titanate nanosheets. Diferrocenylborinic acid (1) was immobilised into the titanate film by incubation of the modified electrode in 1 mM solution of Fc<sub>2</sub>BOH in chloroform overnight in a fridge/dark or at room temperature with ambient light conditions with the same results. The modified electrode was then potential-cycled in aqueous electrolyte solution (0.1 M Na<sub>2</sub>SO<sub>4</sub> or 0.1 M NaF) until the stabilisation of the voltammetric response (typically from one to twelve potential cycles).

# Results and discussion

# 3.1. Synthesis and characterisation of diferrocenylborinic acid

The synthesis of diferrocenylborinic acid (1) was based on previously published procedures developed by Post *et al.*<sup>18</sup> and by Rebiere<sup>30</sup> (see Fig. 1). Ferroceneboronic acid or its anhydride or ester were used as the boron donor. Using only crude commercial ferroceneboronic acid (as purchased containing FcB(OH)<sub>2</sub> and its anhydride boroxine (FcBO)<sub>3</sub>) gave the desired product with 15–20% isolated yield of 1. When employing the more reactive ferroceneboronic acid 1-propanol ester (3), the yield is improved to 38% (in respect to boron equivalent in reactants) with further product (1) being recovered from the hydrolysis of side product trisferrocenylborane (2).

Fc<sub>2</sub>BOH (1) was structurally characterised using <sup>1</sup>H, <sup>11</sup>B and <sup>13</sup>C NMR (ESI Fig. S1–S3, see ESI†), by HRMS (Fig. S9†), by

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FTIR (Fig. S11 and S12†), by spectroelectrochemistry (Fig. S15 and S16†), and by single crystal X-ray crystallography (Fig. S17†). In <sup>11</sup>B NMR a characteristic peak is observed at 46.0 ppm consistent with sp<sup>2</sup>-boron, which compares to the <sup>11</sup>B chemical shift for ferroceneboronic acid of 31.0 ppm.2 For further details and data, see ESI.† Fc<sub>3</sub>B (2) was also characterised (see ESI Fig. S4-S6, S10 and S13†). Ferroceneboronic acid 1-propanol ester (3) was identified only as an intermediate, as a part of the reaction mixture with (FcBO)<sub>3</sub> (see NMR spectra ESI Fig. S7 and S8†).

Orange-red prism-shaped crystals of 1 for X-ray crystallography (needles growing from glass wall into solution) were slowly grown from saturated solutions in CDCl3 or toluene (50 mg cm<sup>-3</sup>, 1 week or to dryness). Single crystal X-ray crystallography analysis revealed that, in the monoclinic solid state, 1 exists as disordered Fc2BOH structure in the ratio 1:1 in space group  $P2_1/c$  (Fig. 2).

Both ferrocenyl moieties in Fc<sub>2</sub>BOH occupy places similarly to those observed in structurally related bridging borinic acid derivatives.31 The substituted cyclopentadienide rings are fairly co-planar. The B-Cipso bond distances are 1.563(4) Å and 1.571(4) Å. The B-O bond length is 1.375(4) Å, which is in line with bond length data for other related borinic acids as shown in Table 1. The B-O bond length is essentially identical to that for ferroceneboronic acid. 32

## 3.2. Voltammetric characterisation of diferrocenylborinic acid in solution

The electrochemical behaviour of 1 in solution is dependent on the type of electrolyte and the experimental conditions. For exploratory electrochemical characterisation, two non-aqueous solvents were used - acetonitrile and dichloromethane (with 50 mM TBAH as background electrolyte). Fig. 3A shows cyclic voltammetry data for 1 dissolved in acetonitrile. A separated pair of redox processes (see  $E_{p,ox}1$  and  $E_{p,ox}2$ ) consistent with the stepwise oxidation of the two ferrocenyl substituents can be seen with a gap in potentials  $\Delta E_{\rm p,ox}$  = 160 mV. This observation is consistent with the oxidation of the first ferrocenyl unit causing a pull in electron density in the direction of the oxidised ferrocenyl moiety and a resulting shift in the oxidation of the second ferrocenyl moiety to higher potentials. Data from spectroelectrochemistry in Fig. 3D confirm an isosbestic point associated with predominantly two stable pro-

Table 1 Comparison of selected B-C and B-O bond lengths for related borinic acids

Compound	B-C (R <sub>1</sub> and R <sub>2</sub> ) bond lengths (Å)	B–O bond length (Å)	
Fc <sub>2</sub> BOH <sup>a</sup>	1.563(4)/1.571(4)	1.375(4)	
1,2-FcBOH <sup>32</sup>	1.539/1.550	1.379	
1,2-FcBOMe <sup>32</sup>	1.556/1.556	1.361	
Diazulenylborinic acid <sup>33</sup>	1.568/1.571	1.348	
Bis(pentafluorophenyl)borinic acid <sup>34</sup>	1.615/1.614	1.528	
Dimesitylborinic acid <sup>35</sup>	1.574/1.588	1.365	
Mesityl(2,4,6-trimethoxyphenyl) borinic acid <sup>36</sup>	1.570/1.580	1.368	
10 <i>H</i> -Dibenzo[ $b$ , $e$ ][1,4]oxaborinin-10-ol (computed values) <sup>37</sup>	1.547/1.547	1.354	
5-Methyldibenzo[ $b$ , $e$ ][1,4]azaborinin-10(5 $H$ )-ol (computed values) <sup>37</sup>	1.543/1.541	1.360	
<sup>a</sup> This study			

This study.

ducts, e.g. the Fe(II)Fe(II) starting material and the Fe(III)Fe(III) product. The shift of the main absorbance peak upon oxidation from 220 nm to 260 nm is consistent with published observations for similar materials.<sup>38</sup> In dichloromethane (Fig. 3B), the same step-by-step oxidation of both ferrocenyl moieties is also observed (see scheme in Fig. 3F). A scan rate of approximately 0.4 V s<sup>-1</sup> or higher is required to avoid the formation of new products (see new reduction peaks at 0.10 and -0.14 V vs. SCE, denoted by \*, and ESI Fig. S14† for further detail). During lower scan rates in dichloromethane more complex changes in behaviour of 1 occur, which are tentatively assigned here to the formation of products with higher electron density on the boron, e.g. deprotonation and nucleophile adduct formation possibly leading to boroxines. The additional complexity is confirmed by the absence of an isosbestic point in data from spectroelectrochemical measurements (Fig. 3E).

Data from cyclic voltammetry experiments are summarised in Table 2. The first oxidation in diferrocenylborinic acid occurs at 0.445 V vs. SCE in both acetonitrile and in dichloromethane. However, in the aqueous environment the oxidation occurs at a slightly lower potential of 0.265 V vs. SCE, which may be linked to the more polar solvent environment stabilising the charge. The separation of the first and second redox process,  $\Delta E_{p,ox} = \Delta (E_{mid}1 - E_{mid}2)$ , also decreases when going from organic to aqueous conditions. This may also suggest a

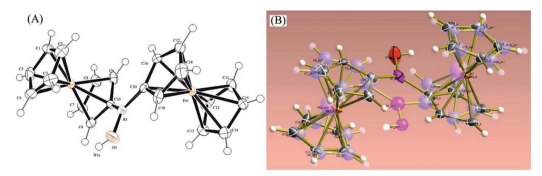
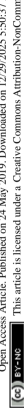


Fig. 2 Molecular structure based on X-ray diffraction of 1 (A) and overlaid with both disordered structures of 1 (B).

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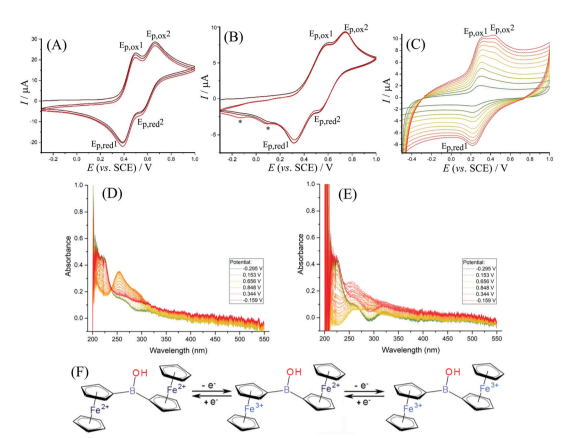


Fig. 3 (A) Cyclic voltammograms (scan rate 100 mV s<sup>-1</sup>; 3 mm diameter glassy carbon electrode) for the oxidation of approx. 1 mM Fc<sub>2</sub>BOH in acetonitrile with 50 mM TBAH. (B) Cyclic voltammograms (scan rate 500 mV s<sup>-1</sup>; 1 mm diameter platinum electrode) for the oxidation of approx. 1 mM Fc<sub>2</sub>BOH in dichloromethane with 50 mM TBAH. (C) Cyclic voltammograms (scan rate 0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9 and 1.0 V s<sup>-1</sup>; 3 mm diameter glassy carbon electrode) for the oxidation of Fc<sub>2</sub>BOH immobilised into a film of titanate nanosheets and immersed in 0.1 M Na<sub>2</sub>SO<sub>4</sub>. (D) Spectroelectrochemistry as a function of applied potential in acetonitrile and (E) in dichloromethane. (F) Proposed mechanism based on two consecutive one-electron transfer reactions.

Table 2 Comparison of redox potentials for oxidation of 1 in different organic solutions and in titanate environments (from cyclic voltammetry, see Fig. 3)

	Potentials					
Environment	Diferrocenyl redox pair	$E_{ m p,ox}$ 1 and $E_{ m p,ox}$ 2 ( $ u$ s. SCE)/mV	$E_{ m p,red}$ 1 and $E_{ m p,red}$ 2 ( $ u$ s. SCE)/mV	$E_{\text{mid}} = \frac{1}{2}(E_{\text{ox}} + E_{\text{red}})$ (vs. SCE)/mV	$\Delta (E_{\rm mid}1 - E_{\rm mid}2)/{\rm mV}$	
Acetonitrile <sup>a</sup>	1	500	390	445	160	
	2	660	550	605		
Dichloromethane <sup>a</sup>	1	570	320	445	190	
	2	740	530	635		
Embedded in titanate composite	1	310	220	265	120	
immersed in 0.1 M aqueous Na <sub>2</sub> SO <sub>4</sub>	2	430	n.o.	n.o.		

<sup>&</sup>lt;sup>a</sup> Data from cyclic voltammograms in solution.

solvent polarity effect and better compensation of the first Fe(III) ferrocenyl charge in the more polar aqueous environment.

### 3.3. Voltammetric characterisation of diferrocenylborinic acid immobilised in a titanate nanosheet host

In aqueous electrolyte, 1 is essentially insoluble, thus immobilisation onto the electrode surface was necessary. Titanate nanosheet material has been shown recently to provide a good

host for ferroceneboronic acid<sup>25</sup> and for a range of other redox active and hydrophobic redox systems. 19 Therefore, the glassy carbon electrode is firstly covered with a thin (ca. 5 µm) layer of 2D titanate nanosheets according to the previously published procedure. Following this, diferrocenylborinic acid 1 was immobilised by absorption from a chloroform solution (1 mM) into the titanate nanosheet film by immersion for 24 h. Cyclic voltammetry measurements were first performed

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in 0.1 M Na<sub>2</sub>SO<sub>4</sub> as a background electrolyte, as this provided stable and reproducible voltammetric responses for both diferrocenylborinic and ferroceneboronic acid.<sup>25</sup> The composite electrode containing 1 immobilised in a titanate nanosheet host was less suitable for work in organic media due to loss of the redox system into the surrounding solution.

As in the case of data for diferrocenylborinic acid 1 in homogeneous organic phase, a split peak voltammetric response is observed in cyclic voltammograms in aqueous environments (Fig. 3C). Two distinct peaks for the oxidation of the ferrocenyl moieties are seen and the separation of the midpoint potentials can be estimated as  $\Delta E_{\text{mid}} = 120 \text{ mV}$  (Table 2). The second oxidation step shows a peak response during the forward potential scan, but not during the backward potential scan. This observation can be explained by the occurrence of a follow-on chemical reaction step. The second oxidation peak during the forward potential scan is also diminished at lower scan rates (below 0.7 V s<sup>-1</sup>). The behaviour is reminiscent of that for an ECE scheme with a short-lived intermediate that exists after the first one-electron oxidation. In order to further explore this reaction, square wave voltammetry is considered.

Fig. 4A shows data for square wave voltammetry during oxidation and reduction of diferrocenylborinic acid 1 immobilized in titanate layer on the electrode. Data are consistent with those obtained for cyclic voltammetry (Fig. 3C) and indicate consecutive two-electron oxidation as long as the time for the measurement is short. The second oxidation peak at 0.4 V vs. SCE appears at a scan rate of approximately 0.5 V s<sup>-1</sup> or higher consistent with a short-lived Fe3+-Fe2+ intermediate.

The scheme in Fig. 4B shows a tentatively assigned reaction scheme with the oxidation to the Fe<sup>3+</sup>-Fe<sup>3+</sup> intermediate only at higher scan rates (shorter times). During the reduction, there is no clear evidence for any Fe<sup>2+</sup>-Fe<sup>2+</sup> intermediate being formed.

The oxidation reactions tentatively assumed in Fig. 4B includes the slow binding of a nucleophile (hydroxide). At short time scale, both the first and the second oxidation are observed before the nucleophile reacts (consistent with an EEC reaction type). At slower time scales, however, the first oxidation is rapidly followed by a chemical reaction step then by an immediate oxidation (due to the oxidation potential for the hydroxide adduct being lower, similar potential for the oxidation of the second Fe<sup>2+</sup> centre as in the first oxidation step is assumed). The process can be considered as an ECE type reaction. Very similar reaction sequences have been reported for ferroceneboronic acid. 13,25

Measurements were performed in aqueous 0.1 M NaF to investigate the electrochemically driven binding of fluoride as nucleophile to the borinic acid. Fig. 5 shows data for square wave voltammetry measurements based on diferrocenylborinic acid 1 immobilised in a titanate nanosheet film deposit on a glassy carbon electrode. Comparing the data obtained in Na<sub>2</sub>SO<sub>4</sub> (Fig. 4) and in NaF electrolyte, the binding of fluoride at [Fc<sub>2</sub>BOH]<sup>2+</sup> (EEC pathway during faster scan rates in Fig. 5B) appears to occur at a lower scan rate and seems slower compared to the tentative addition of hydroxide in Na2SO4 electrolyte. A more dramatic change is observed for the reduction where a new  $Fe^{2+}$ – $Fe^{2+}$  intermediate is detected at 0.0 V vs. SCE

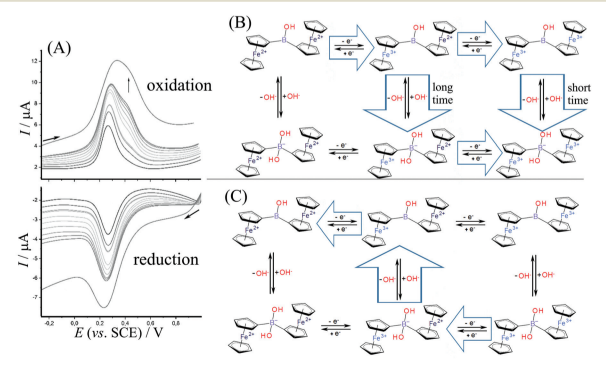


Fig. 4 (A) Redox properties of 1 immobilized in a titanate nanosheet film immersed in aqueous 0.1 M Na₂SO₄. Square wave voltammograms (modulation amplitude 0.02 V, frequency 25 Hz, scan rate 0.12, 0.25, 0.37, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9, 1.0, 2.0 V s<sup>-1</sup>), (B) reaction scheme for oxidation with a short-lived Fe $^{3+}$ /Fe $^{2+}$  species only detected at scan rates higher than 0.4 V s $^{-1}$ . (C) Reaction scheme for reduction via a single reaction pathway.

**Paper** 

(A)

Oxidation

(vii)

(vi)

(vii)

(vii)

(vii)

(viii)

(viii

Fig. 5 (A) Square wave voltammetry data for diferrocenylborinic acid 1 immobilised in a titanate nanosheet film on a glassy carbon immersed in aqueous 0.1 M NaF. Traces (i)–(iv) modulation amplitude 0.02 V, frequency 25 Hz, step 5–60 mV, scan rate 0.12, 0.25, 0.5, 1.5 V s<sup>-1</sup>; traces (v)–(vii) modulation amplitude 0.02 V, frequency 100 Hz, step 20–60 mV, scan rate 2.0, 4.0, 6.0 V s<sup>-1</sup>. (B) Reaction scheme for oxidation with a short-lived Fe<sup>3+</sup>–Fe<sup>2+</sup> species only detected at scan rates higher than 0.1 V s<sup>-1</sup>. (C) Reaction scheme for reduction *via* a dual reaction pathway with a Fe<sup>2+</sup>–Fe<sup>2+</sup> reaction intermediate.

(see \*). This suggests that there are two distinct pathways for reduction and that the corresponding Fe<sup>3+</sup>-Fe<sup>2+</sup> intermediate is more long-lived with fluoride.

The implications of these data in terms of potential applications of diferrocenylborinic acid 1 are limited. The two sequential oxidation processes for the ferrocenyl moities are clearly observed, particularly in dry organic solvents new reactions occur that are currently unresolved. However, diferrocenylborinic acid is water stable and can be reversibly switched between Fe<sup>2+</sup>-Fe<sup>2+</sup> to Fe<sup>3+</sup>-Fe<sup>2+</sup> to Fe<sup>3+</sup>-Fe<sup>3+</sup> state with a change in reactivity to nucleophiles. Here, only tentatively assigned reactions with fluoride are reported and more work will be required to explore reactivity towards other types of nucleophiles as well as reactivity in other types of immobilisation media. Diferrocenylborinic acid could be useful in electroanalytical detection protocols in cases where nucleophiles can bind more strongly or more selectively. More work in particular with saccharides (e.g. glucose) for sensing applications will be required.

# Summary and conclusion

Diferrocenylborinic acid (Fc<sub>2</sub>BOH) was synthesized *via* an improved synthetic path and structurally characterised using NMR, HRMS, FTIR and X-Ray. Redox properties of Fc<sub>2</sub>BOH were investigated in both homogeneous and heterogeneous

phase and redox activity of both ferrocenyl cores has been observed. A 2D-titanate nanosheet film was employed to host diferrocenylborinic acid redox probe molecules on a glassy carbon electrode immersed in aqueous electrolyte. In the presence of disodium sulfate, a stable reversible voltammetric response was observed. In the presence of sodium fluoride, "bound" and "unbound" states of the diferrocenylborinic acid were observed at higher scan rates (>0.1 V s $^{-1}$ ). In comparison with the response of the related ferroceneboronic acid, diferrocenylborinic acid appears more stable during voltammetric cycling in the presence of aqueous electrolyte, but on the other hand, Fc<sub>2</sub>BOH in this titanate nanosheet film appears to be less sensitive towards binding of nucleophilic analytes such as fluoride.

# Conflicts of interest

There are no conflicts of interest to declare.

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rying out HRMS analysis.

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