# **MATERIALS** CHEMISTRY

**FRONTIERS** 







## RESEARCH ARTICLE

**View Article Online** 



Cite this: Mater. Chem. Front., 2020, 4, 2370

Received 16th March 2020, Accepted 11th May 2020 DOI: 10.1039/d0gm00157k

rsc.li/frontiers-materials

## Conjugated molecules for colourimetric and fluorimetric sensing of sodium and potassium<sup>†</sup>

Zachary S. Parr p and Christian B. Nielsen \*\*

We present a series of conjugated molecules for colourimetric and fluorimetric sensing of alkali metal cations Na<sup>+</sup> and K<sup>+</sup>. The dye molecules are based on an electron-rich bithiophene core bridged by an oligoethylene glycol bridge capable of selectively binding either sodium or potassium. The molecules are further functionalised with electron-rich or electron-deficient moieties to modulate the band gap and thus shift the absorbance across the visible range. We observe distinct colour changes and high selectivity for alkali metal cations at millimolar concentrations. We moreover see highly selective fluorimetric responses with the molecules exhibiting either turn-on, turn-off or ratiometric sensing behaviour. We finally demonstrate a dual optical sensor capable of distinguishing between sodium and potassium in solution.

### Introduction

Colourimetric measurement of physical properties is commonplace, with every practicing chemist having at some point employed universal indicator paper as a simple method to measure pH. Commonly, pH paper contains a series of dyes, which exhibit colour change dependant on pH allowing for visual determination of acidity of a solution. In order to examine biological samples, dyes have been employed in microscopy for many years and recent advancements in fluorescent materials have been employed in confocal, fluorescence as well as two-photon microscopy with a number of selective fluorescent probes having been developed.<sup>1,2</sup>

Two of the most common intra- and extracellular biologically active cations are the sodium and potassium ions, associated with a range of biological processes including propagation of nerve impulses, muscle contraction, and regulation of blood pressure.3 A deviation from a narrow typical concentration range of 100-200 mM for Na<sup>+</sup> in serum and 4-5 mM for K<sup>+</sup> in the blood and 140 mM in the intracellular fluid for a healthy individual leads to serious health conditions,4 thus defining a concentration range on the order of 1-100 mM where accurate and selective measurement of sodium and potassium ions is desirable. Alkali metal ion sensing has previously been carried out electrochemically, with either ion-selective membranes on organic electrochemical transistors<sup>5</sup> or with crown ether functionalised polymers<sup>6</sup> and commercially with ion-selective electrodes. Optical sensors are generally characterised by a

Materials Research Institute and School of Biological and Chemical Sciences, Queen Mary University of London, Mile End Road, London E1 4NS, UK. E-mail: c.b.nielsen@amul.ac.uk

† Electronic supplementary information (ESI) available. See DOI: 10.1039/d0qm00157k

slightly simpler operation over ion-selective electrochemical sensors which require careful calibration, temperature and pH control.7

Typically, solution phase optical sensors for Na<sup>+</sup> and K<sup>+</sup> and other metal ions have been based on size-selective heteroatomcontaining macrocycles such as the crown ether, 8,9 azacrown, 10,11 cryptand<sup>12</sup> or calixarene<sup>13,14</sup> as the binding unit. The binding unit is covalently bonded to a chromophore or fluorophore which exhibits a change in absorbance or fluorescence properties as a result of the binding event. The attractiveness of optical sensors is their relative ease of application, high selectivity and high concentration range for quantification. Further, ratiometric sensing, whereby a change in analyte concentration results in a decrease of one optical transition and an increase of a second transition, provide a more robust measurement of concentration. The alternative being an on/off sensor where only depletion or enhancement of an optical event is observed upon change in analyte concentration. Ratiometric sensors eliminate some deviceto-device variation and background noise due to the self-referencing measurement of the probe. Vanishingly few Na and K sensors show ratiometric colourimetric detection in the visible range of the spectrum and even fewer fluorescent materials show ratiometric detection.15-17 Furthermore, if absorbance and emission can be pushed into the 650-950 nm range or the near infrared over 1100 nm then biological sensing becomes more relevant due to the transparency window of tissue18 as well as being beyond autofluorescence emission wavelength for almost all common endogenous fluorophores. 19 It is moreover advantageous for excitation wavelengths to be red-shifted to mitigate the effect of high energy photon damage to cells.

Conjugated small molecules have been explored extensively as active materials in organic electronic applications from

field-effect transistors to photovoltaics, light-emitting diodes and electrochromic devices and more recently biological sensing.<sup>20-24</sup> Common to these materials is the synthetic flexibility that allows for inclusion of various structural moieties to control solubility, solid state morphology, colour, selectivity to biological analytes, and electronic properties, and is one of the standout advantages of organic electronic materials. Recently, it was demonstrated that a conjugated polymer which included either a 17-crown-5 or a 20-crown-6 ether moiety was capable of ratiometric detection of Na<sup>+</sup> and K<sup>+</sup> in the visible range in both solution and solid state. <sup>25,26</sup> Upon binding of an alkali metal ion, a backbone twist occurred breaking conjugation and thereby resulting in ratiometric colour change in the visible spectrum. This concept has more recently been extended to small conjugated molecules showing ratiometric colourimetric response to Na<sup>+</sup> ions.<sup>27</sup> Here, we present a series of related small conjugated molecules and show how rational molecular design allows for both colourimetric and fluorimetric sensing of sodium and potassium. While all colourimetric responses are ratiometric in nature, we show how different structural motifs can be used to alter fluorimetric responses from ratiometric to turn-off and turn-on. As a proof-of-concept, we furthermore show how access to optical sensor molecules with spectral complementarity allows for mixtures with dual Na<sup>+</sup>/K<sup>+</sup> sensing capabilities.

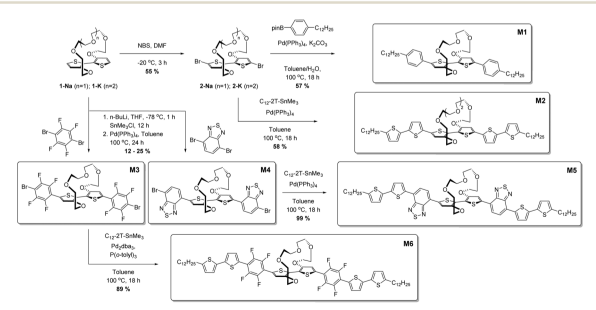
### **Synthesis**

In an effort to create soluble donor-acceptor small molecule ion sensors, we initially synthesised 2,2'-bithiophene tethered at the 3 and 3' positions by a polyether bridge, 28 followed by a series of cross coupling reactions to afford the optical ion sensing materials; full experimental details are available in the ESI† and described in brief here and in Scheme 1. The sodium-binding unit 1-Na was brominated to afford 2-Na, after which Suzuki coupling with 4-(n-dodecyl)phenylboronic acid

pinacol ester afforded M1 in 57% yield as a yellow solid. To afford a second all-donor potassium sensor, 2-K was reacted under Stille conditions with 5-dodecyl-5'-trimethylstannyl-2,2'bithiophene to afford M2 as an orange solid in 58% yield. We then developed a series of donor-acceptor materials by introducing either 2,1,3-benzothiadiazole (BT) or 2,3,5,6-tetrafluorophenylene moieties flanking the bithiophene core through stannylation of **1-Na** in situ followed by reaction with excess acceptor unit under Stille conditions. Reaction with 4,7-dibromo-2,1,3-benzothiadiazole afforded M3 in a yield of 12% as a purple solid, while reaction with 1,4-dibromo-2,3,5,6-tetrafluorphenylene gave M4 in a yield of 25% as a yellow solid. M3 and M4 were further reacted under Stille conditions with 5-dodecyl-5'-trimethylstannyl-2,2'-bithiophene to afford molecules M5 and M6 as blue and orange powders respectively in good yields of 99% and 89% respectively. Thus, a total of six uniquely coloured materials were synthesised as sodium and potassium ion-selective chromo- and fluoroionophores.

#### Results and discussion

Initial characterisation was carried out using UV-vis spectroscopy. The molecules were dissolved in chloroform or tetrahydrofuran (THF) and titrated with a solution of NaBPh4 or KBPh4. Chloroform and THF were chosen to emphasise the behaviour of these novel sensor molecules in solvents miscible and immiscible with water. In the ion-free (unbound) form, the molecules absorb across most of the visible spectrum appearing from yellow to orange through to blue and purple in solution (Fig. 1a). Upon binding with Na<sup>+</sup> (M1, M3, M4, M5, and M6) or K<sup>+</sup> (M2) in solution, there is a quenching of the dominant optical absorption band and a simultaneous growth of a new blue-shifted (29-88 nm) absorption feature from the alkali metal ion complex, consistent with previous reports for this binding unit (Fig. 1c). 25-27 This blue-shift in absorbance is rationalised from a twist in the central S-C-C-S



Scheme 1 Synthetic scheme outlining the modular synthesis of molecular ion sensors M1-M6 via stepwise palladium-mediated cross-coupling reactions;  $C_{12}$ -2T-SnMe<sub>3</sub> = 5-dodecyl-5'-trimethylstannyl-2,2'-bithiophene.

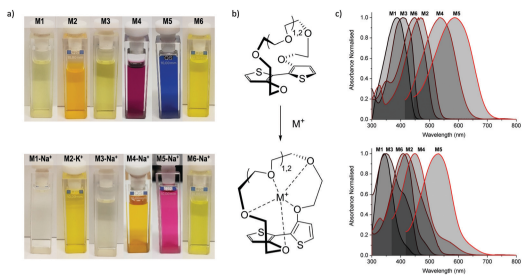


Fig. 1 (a) Photographs of the molecules (top) and their alkali metal ion complexes (bottom) in chloroform solution; (b) illustration of backbone twist of crown ether tethered bithiophene sensing unit adapted from ref. 26; (c) collated UV-vis absorption spectra for the molecules (top) and their alkali metal ion complexes (bottom) in chloroform solution at approximately  $10^{-5}$  M concentration at room temperature.

dihedral angle of the ion selective bithiophene core upon Na $^{+}$  or K $^{+}$  binding. The conjugation along the backbone is disrupted as the dihedral twists as much 70 $^{\circ}$  as has been demonstrated by previous crystal structures and theoretical calculations (Fig. 1b). Representative titration plots can be seen in Fig. 2 for **M2** and **M5** with the remaining plots depicted in Fig. S1–S12 in the ESI $^{\dagger}$  and all data summarised in Table 1.

The molecules can be divided into three categories. **M1** and **M2** are all-donor molecules comprising only phenylene and/or thiophene building blocks. In solution, **M1** is pale yellow, turning colourless upon titration with Na<sup>+</sup> caused by a 44 nm blue-shift (Fig. 1a). The increased conjugation length of the sexithiophene backbone in **M2** causes a red-shift of the absorption maximum ( $\lambda_{max}$ ) from 397 nm for **M1** to 468 nm for **M2**. When titrated with KBPh<sub>4</sub>, the  $\lambda_{max}$  blue-shifts 49 nm as depicted in

Fig. 2a. The K<sup>+</sup> sensing unit (1-K) was employed for M2 in order to have complementary Na<sup>+</sup> and K<sup>+</sup> molecular sensors available. The second set of molecules (M3 and M6) are weak donor–acceptor systems containing electron-deficient tetrafluorophenylene units with additional extension of conjugation in M6 with flanking bithiophene moieties. M3 displays a  $\lambda_{\rm max}$  at 412 nm with a significant 65 nm blue-shift upon ion-sensing to form the colourless M3-Na<sup>+</sup> complex. The  $\pi$ -extended M6 also appears yellow with a  $\lambda_{\rm max}$  of 453 nm in solution (Fig. 1a); however, it displays the smallest blue-shift of 29 nm when undergoing complexation to form M6-Na<sup>+</sup>. The third set of molecules are stronger donor–acceptor systems where the central 1-Na core is flanked by highly electron-deficient 2,1,3-benzothiadiazole groups, thus affording M4 and its bithiophene-extended derivative M5. These red-shifted materials achieved the

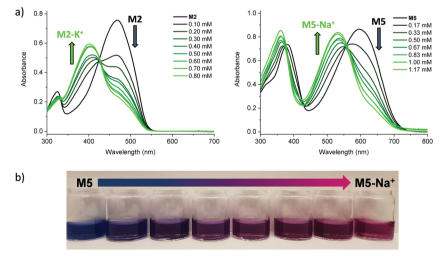


Fig. 2 (a) Representative UV-vis absorption spectra in tetrahydrofuran solution for M2 upon titration with aliquots of 0.002 M KBPh<sub>4</sub> in tetrahydrofuran (left) and for M5 upon titration with aliquots of 0.005 M NaBPh<sub>4</sub> in tetrahydrofuran (right); (b) photographs of M5 titration series with NaBPh<sub>4</sub> in tetrahydrofuran. All data recorded at room temperature.

Table 1 Summarised spectrophotometric data and sensing parameters for sensor molecules M1-M6. Dissociation constants (K<sub>d</sub>), limit-of-detection (LOD) and limit-of-quantification (LOQ) reported for Na<sup>+</sup> for M1, M3-M6 and for K<sup>+</sup> for M2

	$\lambda_{\max}^a M (nm)$	$\lambda_{\max} M-Na^{+a} (nm)$	$\lambda_{\max} M-K^{+a} (nm)$	$\Delta \lambda_{max}^{a} (nm)$	$\varepsilon  \mathrm{M}^b  (10^3  \mathrm{cm}^{-1}  \mathrm{M}^{-1})$	$K_{\rm d}^{\ a}  ({\rm mM})$	$LOD^{a}$ (nM)	$LOQ^{a}$ (nM)
M1	397	338	_	59	32.6	1.45	70	220
<b>M</b> 2	468	_	400	68	72.8	0.13	90	300
<b>M</b> 3	412	347	_	65	31.6	34.4	1100	3700
<b>M4</b>	540	452	_	88	43.0	6.81	240	790
M5	595	531	_	64	83.5	0.33	180	590
<b>M</b> 6	453	424	_	29	123.0	18.2	590	1960

<sup>&</sup>lt;sup>a</sup> Data obtained from tetrahydrofuran solution at room temperature. <sup>b</sup> Extinction coefficients (ɛ) determined in chloroform solution at room temperature.

highest visual contrast of the series upon sensing of sodium ions. In solution, M4 is purple with a  $\lambda_{max}$  of 540 nm; the colour changes from purple to yellow when the M4-Na<sup>+</sup> complex forms (Fig. 1a). This molecule displayed the largest blue-shift of 88 nm upon complexation and displayed two isosbestic points at 481 nm and 392 nm indicating a direct conversion from M4 to M4-Na<sup>+</sup> without interfering side-reactions. M5 is blue when ionfree and when complexed with Na<sup>+</sup> has a magenta hue (Fig. 1a). The shift in  $\lambda_{\text{max}}$  was 64 nm from 595 nm to 531 nm upon Na<sup>+</sup> sensing as depicted in Fig. 2a. It should be noted that the presence of sodium ions can be detected visually at concentrations below 1 mM using a  $10^{-5}$  M solution of M5 as illustrated in Fig. 2b.

The affinities of the sensor molecules for Na<sup>+</sup> and K<sup>+</sup> were investigated at room temperature and quantified using the Benesi-Hildebrand method to afford the dissociation constants,  $K_{\rm d}$ , listed in Table 1 with further details available in the ESI<sup>†</sup> (Fig. S12-S23).<sup>29,30</sup> The limit of detection (LOD) and limit of quantification (LOQ) were quantified from the absorbance of the complex (M-Na<sup>+</sup> or M-K<sup>+</sup>) relative to the absorbance of the ion-free sensor molecule (M) at their respective absorption maxima during titration experiments (Table 1 and Fig. S24-S34, ESI†). We observe

slightly lower K<sub>d</sub> values in chloroform than in THF, indicating stronger affinities, likely due to the ability of THF to coordinate ions in solution thus competing with the binding of the molecule.31 Deviations from linearity in the Benesi-Hildebrand and LOD/LOQ plots obtained in chloroform lead us to focus mainly on the alkali metal ion affinities in THF in the following. We observe that the dissociation constant drops markedly as the conjugation length of the sensor molecule increases with  $K_d$  of **M6** (18.2 mM) lower than that of **M3** (34.4 mM) and similarly  $K_d$ of M5 (0.33 mM) lower than that of M4 (6.81 mM). We hypothesise that this is caused by the fact that breaking of the  $\pi$ -conjugation upon complexation is more unfavourable in a shorter  $\pi$ -conjugated molecule than in a longer molecule as the highest occupied molecular orbital would be significantly more localised in the first instance. Interestingly, the fluorinated molecules, M6 and M3, exhibit the highest  $K_d$  values followed by the brominated sensor molecule M4, which are all significantly higher than that of the non-halogenated molecules. It is likely that the halogen lone pairs and the partial negative charge on the polarised C-X bonds compete as binding sites for the alkali metal ions and therefore reduce the binding efficiency of

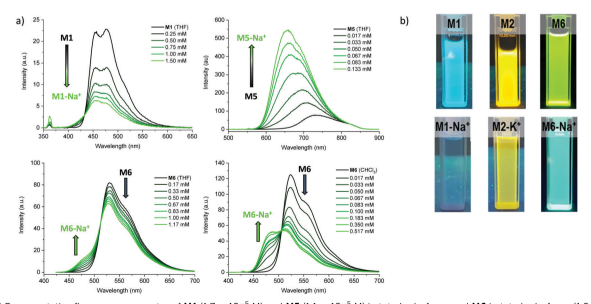


Fig. 3 (a) Representative fluorescence spectra of M1 ( $1.7 \times 10^{-5}$  M) and M5 ( $1.1 \times 10^{-5}$  M) in tetrahydrofuran and M6 in tetrahydrofuran ( $1.0 \times 10^{-5}$  M) and chloroform  $(2.3 \times 10^{-5} \text{ M})$  upon titration with aliquots of NaBPh<sub>4</sub>; (b) photographs of representative unbound sensor molecules and their alkali metal ion complexes in chloroform solution upon excitation at 365 nm. All data recorded at room temperature

the crown ether moiety.<sup>32</sup> For all the sensor molecules, the alkali metal ion binding is at least an order of magnitude weaker than for the parent crown ethers 15-crown-5 ( $K_{\rm d}<0.1$  mM) and 18-crown-6 ( $K_{\rm d}<0.01$  mM).<sup>33</sup> This allows for a reversible process as the unbound state can be regenerated upon addition of either 15-crown-5 or 18-crown-6. Characteristic of  $\pi$ -conjugated materials, the sensor molecules show very high extinction coefficients of 32 600–123 000 M<sup>-1</sup> cm<sup>-1</sup> with the longer molecules M2, M5 and M6 showing the highest values as expected. Sensor molecules M2, M4, M5 and M6 are highly bench stable having

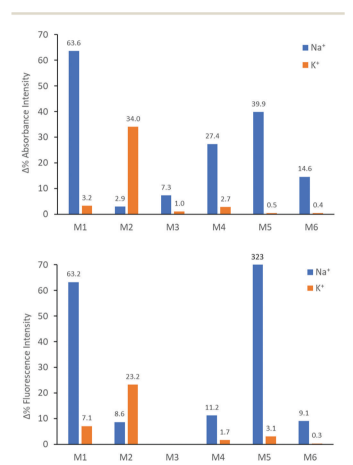


Fig. 4 Colorimetric (top) and fluorimetric (bottom) selectivity of the sensor molecules for Na<sup>+</sup> and K<sup>+</sup> (added as BPh<sub>4</sub> salts at 0.33  $\mu$ M concentration) measured at 10<sup>-5</sup> M sensor molecule concentration in tetrahydrofuran at room temperature. The change in emission is 323% for **M5** upon addition of Na<sup>+</sup>, the axis was cropped for clarity.

been stored as solids for six months under ambient conditions without any change in UV-vis spectra, while small changes are seen for M1 and M3 possible due to oxidation and/or aggregation (Fig. S35–S40, ESI†).

We subsequently explored the emissive properties of the sensor molecules by exciting at the isosbestic point identified from the UV-vis titration experiments (Fig. 3 and Fig. S41-S50, ESI†). In line with the colourimetric studies, the sensor molecules were examined in both tetrahydrofuran and chloroform with our main focus devoted to the behaviour in tetrahydrofuran due to more reliable extraction of sensing parameters and tetrahydrofuran generally solvating alkali metal salts better than chloroform. Upon titration with NaBPh<sub>4</sub> in tetrahydrofuran, M1 exhibited a decrease in fluorescence intensity (turn-off), with a slight shift in emission wavelength ( $\lambda_{em}$ ) from 476 nm to 459 nm. With KBPh4, M2 showed a similar decrease in fluorescence intensity with a slight broadening and blue-shift of the emission band. The second set of sensor molecules, M4 and M5, showed a significant fluorescence enhancement (turn-on) upon complexation with Na<sup>+</sup>. This turn-on fluorimetric response also had the advantage of being significantly shifted towards the red compared to the other materials, with the  $\lambda_{em}$  being 712 nm and 733 nm for the ion-free sensor molecules in solution and 650 nm and 661 nm upon Na<sup>+</sup> complexation for **M4** and **M5**, respectively. The molecules behave similarly in chloroform exhibiting strong fluorescence turn-on (Fig. S46 and S48, ESI†). It is likely that there is significant charge transfer character from donor to acceptor moiety in these molecules, which in the ion-free form quenches the emission. Complexation with sodium, on the other hand, is expected to disrupt effective charge transfer and thus enhance emission. While M3 did not fluoresce, M6 showed a ratiometric fluorimetric response to titration with sodium ions. Upon titration with NaBPh4 in THF, a decrease in the emission peak at  $\lambda_{em}$  530 nm occurred with a concurrent increase in intensity of a second fluorescence feature at 480 nm; the ratiometric response is even clearer in chloroform as illustrated in Fig. 3. The selectivity of all the molecules to Na<sup>+</sup> and K<sup>+</sup> examined by colourimetry and fluorimetry is summarized in Fig. 4 (details in Fig. S64-S73, ESI†). As expected from the crown ether ring size, M1, M3, M4, M5 and M6 were all highly selective to Na<sup>+</sup> with M2 being moderately selective to K<sup>+</sup>. The Stokes shifts (Table 2) for the unbound molecules were on the order of 3000-4000 cm<sup>-1</sup> with the shorter molecules M1 and M4 showing the larger Stokes shifts and the longer and more conformationally locked molecules showing

Table 2 Summarised fluorescence data for sensor molecules M1-M6

	$\lambda_{\mathrm{em}} \mathbf{M}^{a} (\mathrm{nm})$	$\lambda_{\rm em} \ { m M-Na}^{+a} \ ({ m nm})$	$\lambda_{\rm em} \ { m M-K}^{+a} \ ({ m nm})$	Stokes shift $M^a$ (cm <sup>-1</sup> )	Stokes shift M-Na <sup>+</sup> /M-K <sup>+ a</sup> (cm <sup>-1</sup> )	$\Phi^b$ M	$\Phi^b$ M-Na $^+$ /M-K $^+$
M1	476	459	_	4180	7800	0.06	0.01
M2	546	_	542	3050	6550	0.12	0.12
M4	712	650	_	4470	6740	0.005	0.02
M5	733	661	_	3160	3700	0.006	0.02
M6	524	528	_	2990	4650	0.36	0.07

<sup>&</sup>lt;sup>a</sup> Data obtained from tetrahydrofuran solution at room temperature at approximately  $10^{-5}$  M concentration (see Fig. S41–S50, ESI) with excitation wavelengths 362 nm (M1), 426 nm (M2), 481 nm (M4), 556 nm (M5) and 427 nm (M6). <sup>b</sup> Quantum yields ( $\Phi$ ) determined in either dichloromethane (M2) or chloroform (M1, M4–M6) solution at room temperature (see Fig. S51–S63, ESI).

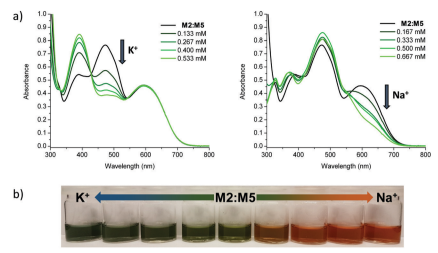


Fig. 5 (a) UV-vis absorption spectra of the M2:M5 blend at approximately  $3 \times 10^{-5}$  M in tetrahydrofuran titrated with aliquots of KBPh<sub>4</sub> (left) and NaBPh<sub>4</sub> (right) in tetrahydrofuran; (b) photographs of M2:M5 titration series with KBPh4 and NaBPh4 in tetrahydrofuran. All data recorded at room temperature.

lower values.<sup>34</sup> The metal ion complexes have significantly larger Stokes shifts with values approaching 8000 cm<sup>-1</sup>, again with the shorter sensor molecules with more conformational freedom having the larger values.35,36 Quantum yields (Table 2) for the unbound molecules were lowest for the bromine-containing molecules M3 (no fluorescence detected) and M4, while strong donor-acceptor character (M4 and M5) was also seen to promote non-radiative emission. The longer molecules M2 and M6 showed relatively high quantum yields suggesting that the bulky tethered crown ether effectively prevents intermolecular interactions and the associated non-radiative processes.

Having confirmed alkali metal ion selectivity and the ability of the sensor molecules to give visual responses to Na<sup>+</sup> and K<sup>+</sup>, the two most complementary sodium and potassium sensor molecules in terms of spectral response were investigated as a blend. Employing M2 as the K+ selective chromophore and M5 as the Na<sup>+</sup> selective chromophore, a 1:1 blend of the two materials was used to create a colorimetric sensor for sodium and potassium ions (Fig. 5). The two molecules have similar molar extinction coefficients and exhibit a large separation in  $\lambda_{max}$  for both unbound and bound species in solution and thus make for an optimal dual sensor system. The blended M2:M5 solution was greenish when ion-free and turned blue upon addition of KBPh4 and orange/red upon addition of NaBPh4.

#### Conclusions

In conclusion, we have shown the design and modular synthesis of six novel ratiometric sensor molecules, capable of selective colourimetric and fluorimetric sensing of sodium and potassium cations. By rational design and choice of peripheral  $\pi$ -conjugated donor or acceptor moieties, we have created ion-selective sensor molecules with colours covering the entire visible spectrum in solution. This ability to navigate the colour palette was demonstrated with relatively short  $\pi$ -conjugated sensor molecules (M1 and M3) that go from coloured to colourless upon sensing and

longer  $\pi$ -conjugated sensors such as M5 that exhibit a very clear colour change from blue to magenta. Spectral complementarity was furthermore utilised to create a dual sodium and potassium sensor that turned blue in the presence of potassium and orange/ red in the presence of sodium cations. Effective alkali metal ion binding - realised by a tethered crown ether that can be varied in size to host either sodium or potassium ions - resulted in low dissociation constants, in particular for the non-halogenated sensor molecules with extended  $\pi$ -conjugation, M2 and M5. The ratiometric colourimetric response, due to the breaking of  $\pi$ -conjugation upon ion complexation, has furthermore enabled low detection and quantification limits in the 100 nM range well within a typical physiological concentration range. We further examined the potential of these molecules as fluorimetric sensors. Depending on the peripheral  $\pi$ -conjugated segments, the molecular sensors exhibited either turn-on, turn-off or ratiometric fluorimetric sensing of Na+ and K+ in both tetrahydrofuran and chloroform. The all-donor materials M1 and M2 displayed turn-off fluorimetric responses, while the strong donor-acceptor systems with 2,1,3benzothiadiazole (M4 and M5) exhibited turn-on fluorimetric responses. M6, on the other hand, with weak donor-acceptor character owing to the weakly electron-withdrawing tetrafluorophenylene moiety, showed clear ratiometric fluorimetric response. We believe that these efforts to understand structure-property relations in  $\pi$ -conjugated sensor molecules will help pave the way for the development of efficient molecular sensors. In this context, we have laid out several molecular design strategies including all-donor and weak and strong donor-acceptor type π-conjugated systems for selective colorimetric and fluorimetric sensing of alkali metal cations at biological relevant concentrations. While this proof-ofprinciple study was carried out with peripheral alkyl chains impairing good solubility and thus sensing behaviour in tetrahydrofuran and chloroform, recent work has demonstrated aqueous activity with  $\pi$ -conjugated molecules bearing ethylene glycol-based chains.37 Further biological relevance can thus be sought through further synthetic design in particular through substituting the peripheral alkyl chains with more polar counterparts.

### Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the Academy of Medical Sciences & Wellcome Trust (SBF002/1158) and the Materials Research Institute for financial support.

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