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Long wavelength TCF-based fluorescence probe for the detection of Alkaline Phosphatase in live cells

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- 12 Keywords: Reaction-based fluorescent probe₁, Alkaline Phosphatase₂, Cell Imaging₃

13 Abstract

- 14 A long wavelength TCF-based fluorescent probe (TCF-ALP) was developed for the detection of
- alkaline phosphatase (ALP). ALP-mediated hydrolysis of the phosphate group of TCF-ALP resulted
- in a significant fluorescence 'turn on' (58-fold), which was accompanied by a colorimetric response
- from yellow to purple. **TCF-ALP** was cell-permeable, which allowed it to be used to image ALP in
- HeLa cells. Upon addition of bone morphogenic protein 2, **TCF-ALP** proved capable of imaging
- 19 endogenously stimulated ALP in myogenic murine C2C12 cells. Overall, TCF-ALP offers promise as
- an effective fluorescent/colorimetric probe for evaluating phosphatase activity in clinical assays or live
- 21 cell systems.

22

1 Introduction

- 23 Alkaline phosphatase (ALP) is an ubiquitous enzyme found in the majority of human tissues, where it
- catalyses the dephosphorylation of various substrates such as nucleic acids, proteins and other small
- molecules (Millán, 2006, Coleman, 1992). ALP also plays an important role in signal transduction and
- regulation of intracellular processes (cell growth, apoptosis and signal transduction pathways) (Julien
- et al., 2011). Abnormal levels of ALP in serum are an indicator of several diseases including bone
- disease (Garnero and Delmas, 1993), liver dysfunction (Rosen et al., 2016), breast and prostatic cancer
- 29 (Ritzke et al., 1998, Wymenga et al., 2001) and diabetes (Tibi et al., 1988). As a result, ALP is regarded
- as a key biomarker in medical diagnosis (Coleman, 1992, Ooi et al., 2007). Therefore, it is important
- 31 to develop a fast, reliable and selective detection system for monitoring ALP activity that is amenable
- 32 to clinical diagnostics.
- 33 There have been numerous approaches to determine ALP levels, including colorimetric (Yang et al.,
- 34 2016, Hu et al., 2017), chemiluminescent (Jiang and Wang, 2012), electrochemical (Zhang et al.,
- 35 2015b), surface-enhanced Raman methods (Ruan et al., 2006) and fluorescence (Cao et al., 2016, Fan

- 36 et al., 2016). This work focused on the development of fluorescent probes for the detection of
- 37 biologically relevant analytes (Sedgwick et al., 2017a, Sedgwick et al., 2018b, Wu et al., 2017,
- 38 Sedgwick et al., 2017b, Sedgwick et al., 2018a, Zhang et al., 2019). Fluorescence has many advantages
- 39 over other methods owing to its simplicity and high sensitivity/selectivity, providing rapid, non-
- 40 invasive, real-time detection (Wu et al., 2017). Whilst there have been many fluorophores developed
- 41 for assaying ALP activity such as organic dyes (Zhang et al., 2015a, Zhao et al., 2017), conjugated
- 42 polymers (Li et al., 2014), inorganic semiconductor dots (Qian et al., 2015), and noble metal clusters
- 43 (Sun et al., 2014), most require high probe concentrations and crucially rely on short wavelength
- 44 emission, thus limiting their applicability in biological systems. Therefore, ALP probes that operate at
- 45 long wavelengths are urgently required. Such probes should allow for deeper tissue penetration and be
- 46 subjected to less cell-based autofluorescence (Liu et al., 2017, Zhang et al., 2017, Tan et al., 2017).

47 2 **Results and Discussion**

48 2.1 **Chemistry**

- 49 Here we report a TCF-based fluorescent probe that allows for the detection of ALP and/or ACP. As
- 50 shown in **Scheme 1**, this probe (**TCF-ALP**) is based on the conjugation of 2-dicyanomethylene-3-
- 51 cyano-4,5,5-trimethyl-2,5-dihydrofuran (TCF) to an electron-donating moiety, a phosphorylated
- 52 phenol; this affords an internal charge transfer (ICT) donor- π -acceptor (D- π -A) system whose
- 53 fluorescence properties vary dramatically following ALP-mediated phosphate group cleavage
- 54 (Gopalan et al., 2004, Liao et al., 2006, Lord et al., 2008, Bouffard et al., 2008, Jin et al., 2010, Teng
- 55 et al., 2018, Sedgwick et al., 2017b). TCF-ALP was synthesised in four steps with an overall yield of
- 56 27% (Scheme 2). In brief, 3-hydroxy-3-methyl-2-butanone, malononitrile and NaOEt were heated at
- 57 reflux in EtOH for 1 h. The resultant precipitate TCF (1) was then treated with a mixture of piperidine
- (cat.) and 4-hydroxybenzaldehyde in EtOH to afford intermediate 2 (TCF-OH). Intermediate 2 was 58
- 59 then treated with diethylchlorophosphate, DMAP (cat.) and NEt₃ in THF to give the phosphonate ester
- 60 3. Hydrolysis using trimethylsilyl iodide in dichloromethane (DCM) afforded TCF-ALP as a
- 61 crystalline solid (Et₂O).

62

Spectroscopic studies of TCF-ALP 2.2

- 63 UV-Vis and fluorescence spectroscopic titrations of TCF-ALP were performed in 50 mM Tris-HCl
- 64 buffer in the absence and presence of ALP from porcine kidney. In the absence of ALP, TCF-ALP
- was found to have no UV absorption features above ~550 nm; however, upon addition of ALP a 65
- bathochromic shift in the UV absorption maximum was observed (from 440 to 580 nm), which was 66
- 67 accompanied by a change in colour from yellow to purple (Figure S1). ALP-mediated hydrolysis of
- **TCF-ALP** to form highly fluorescent phenol (2), was confirmed by ³¹P NMR studies and HRMS (See 68
- Figure S1 S4). The effect of pH on the rate of ALP mediated hydrolysis of TCF-ALP was evaluated. 69
- 70 It was found that incubation with 0.8 U/mL of ALP at pH 9.2 resulted in the largest fluorescence
- 71 response (Figure S5). Consequently, all in vitro experiments to determine ALP activity were carried
- 72 out in 50 mM Tris-HCl buffer at pH 9.2.
- 73 The kinetics of ALP towards TCF-ALP were determined via fluorescence spectroscopy (Figure S6
- 74 and S7), with the resultant fluorescence data analysed using the Michaelis-Menten equation (Figure
- 75 **S8**). This revealed a K_m of 35.81 \pm 2.63 μ M and a V_{max} of 3029 \pm 157.3 min⁻¹ for hydrolysis of **TCF**-
- 76 **ALP** by ALP at pH 9.2 (see SI for details). **TCF-ALP** was then incubated with various concentrations
- 77 of ALP (0.0 - 0.2 U/mL) for 15 minutes to evaluate its ability to monitor ALP activity. As shown in
- 78 **Figure 1**, a significant fluorescence response was observed in the presence of ALP (58-fold) with a
- 79 limit of detection (LOD) calculated as 0.12 mU/mL (**Figure S9**). This sensitivity is comparable to other

- 80 fluorescent probes found in literature (**Table S3**). Although serum alkaline phosphatase levels vary
- 81 with age in normal individuals (Lowe et al., 2018), it is widely accepted that serum ALP levels in
- 82 healthy adults lies between 39 – 117 U/mL (Sahran et al., 2018, Saif et al., 2005). This suggests that
- 83 **TCF-ALP** is capable of detecting ALP in human serum, and therefore could be used in clinical assays.
- 84 Inhibition studies were carried out in the presence of sodium orthovanadate (Na₃VO₄), which is known
- 85 to be a strong inhibitor of ALP activity. Addition of Na₃VO₄ resulted in a decrease in the fluorescence
- response in the TCF-ALP hydrolysis assay (see Figure S10) (Swarup et al., 1982). These inhibition 86
- 87 studies enabled an IC₅₀ of 6.23 µM to be calculated (**Figure S11**), which is similar in value to other
- 88 ALP substrates that have been reported in the literature (Tan et al., 2017, Zhang et al., 2015a).
- 89 The selectivity of **TCF-ALP** towards other biologically relevant enzymes (at their optimal pH values)
- 90 was then determined (Figure 2 and S12), with TCF-ALP displaying high substrate selectivity for ALP
- 91 over other common hydrolytic enzymes (e.g. trypsin, porcine liver esterase) or non-specific binding
- proteins (e.g. bovine serum albumin (BSA)). Interestingly, TCF-ALP produced a fluorescence 92
- 93 response when treated with acid phosphatase (ACP). The detection of this enzyme is of significance
- 94 since it is a tumour biomarker for metastatic prostate cancer (Makarov et al., 2009). Normal levels of
- ACP in serum range from 3.0 4.7 U/mL, and elevated ACP levels can be indicative of a variety of 95
- 96 other diseases (Bull et al., 2002). Furthermore, TCF-ALP proved capable of detecting ACP (25-fold
- 97 fluorescence enhancement) and ALP (38-fold enhancement) at a physiological pH of 7.1 (Figure S13
- 98 and S14). Kinetic determination of ALP and ACP towards TCF-ALP at pH 7.1 was conducted, and
- 99 the resultant K_m and V_{max} were compared (see SI 2.1 and Figures S15-S18). It was found that ALP has
- a smaller K_m value in comparison to ACP (0.38 \pm 0.042 μM and 99.22 \pm 13.16 μM respectively) and 100
- a lower V_{max} (208 ± 3.81 min⁻¹ and 1962 ± 223.6 min⁻¹ respectively). Hence, ALP has higher affinity 101
- 102 towards TCF-ALP compared to ACP, thus is more selective towards ALP at physiological pH.
- 103 According to current standards, determination of ALP and ACP is undertaken at the phosphatase's
- 104 optimum pH. For example, the Centers for Disease Control and Prevention (CDC) procedure for ALP
- 105 determination is carried out in 2-amino-2-methyl-1-propanol (AMP) buffer at pH 10.3 ((CDC), 2012).
- 106 This is in accordance with other literature sources (Guo et al., 2018, Di Lorenzo et al., 1991, Radio et
- al., 2006, Pandurangan and Kim, 2015). Likewise, ACP determination is carried out at pH 4-6 (Myers 107
- 108 and Widlanski, 1993, Boivin and Galand, 1986, LI et al., 1984). Following these observations, further
- 109 studies were conducted to determine selectivity at pH 5.0 and 9.2 (Figures S19 - S22). Results showed
- 110 that TCF-ALP acts selectivity towards ACP at acidic pH, and ALP at alkaline pH. Therefore, TCF-
- 111 **ALP** can be used to selectively detect ALP/ACP in clinical assays, or live cell systems (provided the
- 112 buffer solution is optimal for the phosphatase under study).

2.3 Imaging of ALP in living cells

113

- Prior to exploring whether **TCF-ALP** could be used to image ALP activity levels in live cells, the 114
- 115 cytotoxicity of TCF-ALP was assessed using a MTT assay (Figure S23). Negligible cell toxicity was
- 116 observed for TCF-ALP concentrations between $0-5 \mu M$, and cell viability was only slightly reduced
- 117 (91%) when incubated with 10 μM TCF-ALP, indicating good biocompatibility.
- 118 TCF-ALP proved cell permeable to HeLa cells that express ALP and provided a clear 'turn on'
- 119 response (Figure 3). In contrast, pre-treatment of HeLa cells with 5 mM Na₃VO₄ prior to incubation
- 120 with TCF-ALP resulted in minimal 'turn on'. This was taken as evidence that the increase in TCF-
- 121 **ALP** fluorescence levels seen for HeLa cells in the absence of Na₃VO₄ is due to ALP activity. We thus
- 122 conclude **TCF-ALP** is a probe that allows for the selective cellular imaging of ALP activity.

- Bone morphogenetic protein 2 (BMP-2) is capable of inducing osteoblast differentiation into a variety
- of cell types (Guo et al., 2014, Wang et al., 2015) via pathways that result in increased ALP mRNA
- expression, leading to increased ALP activity (Kim et al., 2004). Treatment of myogenic murine C2C12
- cells with **TCF-ALP** resulted in a low fluorescence intensity (low ALP levels) being observed (**Figure**
- 4); however, pre-treatment of these cells with BMP-2 (300 ng/mL, 3 days) resulted in a significant
- increase in **TCF-ALP**-derived fluorescence intensity (high ALP levels). Once again, pre-incubation
- with 5 mM Na₃VO₄ led to no fluorescence response being observed in the cells treated with **TCF-ALP**
- (with or without BMP-2). This provided support for the notion that **TCF-ALP** is capable of imaging
- endogenous ALP activity induced by BMP-2.

132 **3 Conclusions**

- In summary, a long wavelength TCF-based fluorescent probe (**TCF-ALP**) has been prepared with the
- goal of detecting ALP activity. ALP hydrolyses the phosphate group of **TCF-ALP** resulted in a
- significant 'turn on' fluorescence response (58-fold) within 15 minutes. These spectroscopic changes
- were accompanied by a colorimetric change from yellow to purple. This enables **TCF-ALP** to be used
- as a simple assay for the evaluation of ALP activity. Further analysis revealed that **TCF-ALP** could
- also be used as a probe for detecting ACP activity. **TCF-ALP** was shown to be cell permeable, enabling
- its use as a fluorescent probe for monitoring ALP levels in HeLa cells. **TCF-ALP** also proved capable
- of imaging endogenously stimulated ALP produced in myogenic murine C2C12 cells through the
- addition of bone morphogenetic protein 2. We thus suggest that **TCF-ALP** offers promise as a tool for
- measuring ALP and ACP activity levels in clinical assays or in live cell systems.

143 **4 Conflict of Interest**

- The authors declare that the research was conducted in the absence of any commercial or financial
- relationships that could be construed as a potential conflict of interest.

146 **5 Author Contributions**

- 147 LG and ACS carried out synthetic and spectroscopic experiments and co-wrote the manuscript with
- TDJ and JLS. JEG and GTW carried out background experiments. GK carried out cellular imaging
- experiments. JPL carried out the ³¹P NMR titrations. J-YM and ATAJ are supervisors of LG and GTW.
- SDB, JY, JLS and TDJ both conceived the idea and helped with the manuscript.

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159

160

161 **8 Supplementary Material**

- Data supporting this study are provided as supplementary information accompanying this paper, which
- is available free of charge.

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- 297 10 Captions for Figures and Schemes
- 298 **Scheme 1** A TCF-based fluorescence probe (**TCF-ALP**) for the detection of alkaline phosphatase.
- 299 **Scheme 2** Synthesis of **TCF-ALP.**
- 300 **Figure 1** Fluorescence spectra of **TCF-ALP** (10 μM) produced via the addition of alkaline
- phosphatase (ALP; 0 0.2 U/mL) in 50 mM Tris-HCl buffer, pH = 9.2 at 25 °C. λ_{ex} = 542-15 nm. All
- measurements were made 15 min after the addition of ALP.
- Figure 2 Fluorescence spectra of TCF-ALP (10 μM) recorded in the presence of trypsin (0.8 BAEE
- 304 U/mL), porcine liver esterase, protease from *Streptomyces griseus*, proteinase K, bovine serum
- albumin (0.1 mg/mL), acid phosphatase (50 mM Tris-HCl, pH = 5.0) and alkaline phosphatase (50
- 306 mM Tris-HCl, pH = 9.2). All enzymes were standardised to 0.8 U/mL in Tris-HCl buffer pH 7.1 unless
- otherwise stated. $\lambda_{ex} = 542-15$ nm/ $\lambda_{em} = 606$ nm. Fluorescence measurements were made 30 min after
- adding the enzyme in question.
- Figure 3 HeLa cells incubated under the following conditions: (a) No treatment. (b) TCF-ALP (10
- 310 μM, 30 min). (c) Pre-treated with Na₃VO₄ (5 mM, 30 min), followed by the addition of **TCF-ALP** (10
- 311 μM, 30 min). (d) Pretreated with Na₃VO₄ (0.5 mM, 30 min) and **TCF-ALP** (10 μM, 30 min). Cells
- were washed with DPBS before their fluorescence images were acquired using a confocal microscope.
- 313 Top half: fluorescence images, bottom half: fluorescence images merged with its corresponding DIC
- image. Ex. 559 nm/ em. 575-675 nm. Scale bar : 20 μm. DIC differential interference contrast.
- Figure 4- TCF-ALP in C2C12 cell. C2C12 cells were treated with 300 ng/mL BMP-2 for 3 days and
- then pretreated with 5 mM levamisole for 30 min and stained with 10 µM probe for 30 min. After
- washing with DPBS, fluorescence images were acquired by confocal microscopy. (a) only probe, (b)
- 318 levamisole + probe, (c) BMP-2 + probe (d) BMP-2 + levamisole + probe. Top: fluorescence images,
- bottom: merged with DIC image. Ex. 559 nm/ em. 575-675 nm. Scale bar: 20 μm. DIC differential
- interference contrast.







