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1 **Whole-Cell Paper Strip Biosensors to Semi-quantify**
2 **Tetracycline Antibiotics in Environmental Matrices**

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22 **ABSTRACT**

23 A novel, low-cost, and portable paper strip biosensor was developed for the detection of
24 tetracycline antibiotics. *Escherichia coli*/pMTLacZ containing the tetracycline-mediated
25 regulatory gene used as recognition elements with β -galactosidase as the reporter protein
26 was designed and applied to cheap and portable Whatman filter paper as the carrier to
27 prepare this paper strip biosensor. The detection process was optimized by using EDTA
28 and polymyxin B as a sensitizer to improve the accuracy of detection for complicated
29 matrices. The paper strip biosensor was suitable for tetracycline concentrations in the
30 range of 75–10000 $\mu\text{g/L}$ in water and 75–7500 $\mu\text{g/L}$ in soil extracts. Detection limits of
31 5.23–17.1 $\mu\text{g/L}$ for water and 5.21–35.3 $\mu\text{g/kg}$ for the EDTA soil extracts were achieved at
32 a response time of 90 min. The standard deviation (SD) of detected values by the
33 biosensor paper strip compared to those determined by HPLC was between 13.4–59.6%
34 for tetracycline and 2.01–33.5% for oxytetracycline in water and was between
35 6.22–72.8% for tetracycline and 5.90–43.4% for oxytetracycline in soil. This suggests that
36 the paper strip biosensor was suitable for detecting both tetracycline and oxytetracycline
37 in water, and could provide a suitable detection for extractable oxytetracycline in soils.
38 Therefore, this biosensor provides a simple, economical, and portable piece of field kit for
39 on-site monitoring of tetracyclines in a variety of environmental samples, such as pond
40 water and agricultural soil that are susceptible to tetracycline pollution from feed additives
41 and fertilization with livestock manure.

42

43 **Keywords:** Paper strip; Tetracyclines; Detection; Semi-quantity; Water; Soil

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46

47 **1. Introduction**

48 Antibiotics are largely used in human medicine, animal husbandry, agriculture, and
49 aquaculture (Hoa et al. 2011; Teuber 2001). The rampant usage of antibiotics has led to
50 their ubiquitous occurrence in environmental compartments, including water, soil, and
51 sediment impacted by wastewater, sewage sludge, or livestock manure, etc (Berendonk et
52 al. 2015; Gothwal and Shashidhar 2015; Liu et al. 2017). Until now, over 30 types of
53 antibiotics, including those from the tetracycline, sulfonamide, macrolide, and quinolone
54 classes of antibiotics, have been detected with concentrations typically at microgram per
55 liter levels in pond waters (Limbu et al. 2018; Liu et al. 2017; Rico et al. 2017) and
56 microgram per kilogram levels in soils and sediment (Hu et al. 2010; Kumar et al. 2005;
57 Liu et al. 2009). However, in heavily impacted agricultural waters, concentrations have
58 been measured at the milligram per liter level (Bartelt-Hunt et al. 2011; Peak et al. 2007;
59 Zilles et al. 2005). One major concern is that environmental exposure to these antibiotics
60 can induce resistance in native bacteria, contributing to the development of the
61 environmental resistome, and resulting in lower effectiveness of antibiotics in the
62 treatment of bacterial infections.

63 Preventing the environmental introduction of antibiotics is an ideal way to reduce the
64 proliferation of antibiotic resistant bacteria and antibiotic resistance genes (Berendonk et
65 al. 2015; Gao et al. 2018; Rodriguez-Mozaz et al. 2015; Zhu et al. 2013). Among the
66 antibiotics, tetracyclines are essential in modern intensive agriculture production and
67 widely used in livestock and mariculture farming (Gu et al. 2020; Scarano et al. 2018).
68 Currently, there is a growing effort to reduce their input and impact (Du and Liu 2012; Hu
69 et al. 2010; Kumar et al. 2005; Liu et al. 2017; Thiele-Bruhn 2003). Quick, easy, and
70 cost-effective methods are urgently needed to monitor and support the management
71 practices for drug control during food production as well as following their introduction

72 into water and soil.

73 Currently, numerous antibiotic detection methods have been developed, most of
74 which being chemical analytical methods utilizing high-performance liquid
75 chromatography and mass spectrometry for the extraction and separation of antibiotics
76 from complex environmental samples. These traditional methods require complicated,
77 time-consuming, reagent heavy processing and expensive instrumentation that relies upon
78 the user's expertise to interpret results (Batt and Aga 2005; Hamscher et al. 2002).
79 Recently, paper test strips have been developed for the detection of bioactive contaminants
80 like antibiotics. Most of these paper sensors are based on aptamers or monoclonal
81 antibodies as recognition elements with nanofibers or gold nanoparticles acting as
82 transducers (Abbas et al. 2013; Ferreira et al. 2015; Liana et al. 2012; Ornatska et al.
83 2011). Nevertheless, some deficiencies of these sensors have been recognized and include
84 time-consuming, expensive, cumbersome fabrication (Gullapalli et al. 2010) or
85 insufficient sensitivity due to shifting ion level, pH, temperature, or light interference
86 (Ahmed et al. 2014; Chaiyo et al. 2015; Hossain et al. 2009; Quesada-González and
87 Merkoçi 2015). All these factors have impeded the practical application of existing sensors
88 for antibiotic detection in soils and water.

89 Whole-cell biosensors provide a self-contained portable sensing system based on
90 genetically engineered whole cells that physically are adsorbed on filter-paper strips for
91 on-site semiquantitative visual monitoring of N-acylhomoserine lactones (AHLs) agonists
92 in a test sample (Struss et al. 2010). This paper strip biosensor could serve as a simple and
93 economical portable piece of field kit for on-site monitoring of AHLs in various types of
94 environmental samples. So far, the efficacy of these antibiotic biosensors have only been
95 demonstrated in preliminary research for the detection of antibiotics in water and soils
96 under laboratory conditions (Ma et al. 2020). Studies on antibiotic colorimetric strips

97 using whole-cell biosensors for on-site semiquantitative visual testing are not available.

98 In this study, we reported the development of a self-contained sensing system
99 deployed on a paper strip for the detection of tetracyclines in environmental samples. This
100 sensing system was based on genetically engineered bacterial cells that were directly dried
101 on filter paper strips. These bacterial sensing cells employ β -galactosidase as the reporter
102 protein, which can serve visual detections for antibiotics by using a chromogenic enzyme
103 substrate (X-gal). The sensing system was validated by application in the detection of
104 tetracyclines in water and soil. Paper strip biosensors allowed for visual, fast, convenient,
105 and dose-dependent monitoring of tetracyclines in tested samples, thus demonstrating their
106 value as a portable tool for on-site analysis of environmental samples.

107

108 **2. Methods**

109 *2.1 Chemicals*

110 All chemicals used in this study are described in Supplemental [Text S1](#) and [Table S1](#)
111 of Supporting Information ([SI](#)).

112 *2.2 Plasmids construction*

113 The gene sequences corresponding to the pMT fragment from the
114 tetracycline-mediated regulatory system of *Staphylococcus rostri* strain RST11:Tn916 and
115 transposon Tn10 were isolated from plasmid pMTmCherry using polymerase chain
116 reaction (PCR) with primers ([Table S2](#), [Table S3](#)). The *lacZ* fragment encoding
117 β -galactosidase was PCR-amplified by the primers ([Table S3](#)) from the pUC19 plasmid
118 ([Table S2](#)). The pMTLacZ plasmid was prepared by recombining the *lacZ* fragment and
119 pMT fragment using Trelief™ SoSoo cloning kit. The recombined plasmids were
120 subsequently transformed into competent *Escherichia coli* BL21 cells ([Table S2](#))
121 according to molecular cloning protocols and then verified using highly specific primers

122 (Table S4) (Sambrook et al. 1989). The transformed cells were sifted via culturing on
123 Lysogeny broth (LB) agar plates containing 100 mg/L ampicillin and 20 mg/L X-gal at
124 37 °C overnight for screening *E. coli* BL21/pMTLacZ.

125 2.3 Fabrication of paper strips

126 Paper strips were prepared as described by Struss *et al.* with some modifications
127 (Struss et al. 2010). Biosensor cells were cultured for about 4–5 hours at 37 °C and at 150
128 r/min in the LB medium (containing 100 mg/L ampicillin), resulting in an OD₆₀₀ of
129 0.450–0.500. Cells were centrifuged at 4,000 × *g* for 10 min and resuspended in 1/4
130 lysogeny broth with 10% lactose and polymyxin B. The biosensor cell suspension (50 μL)
131 was spotted on Whatman filter paper strips (1 × 4 cm), dried for 10 min at room
132 temperature and then subsequently dried by vacuum freeze-drying. These paper strips
133 were stored at -20 °C for further study.

134 2.4 Tetracycline analysis in water

135 Standard solutions of six tetracyclines were prepared individually at fourteen
136 concentration levels (0, 10, 25, 50, 75, 100, 250, 500, 750, 1000, 2500, 5000, 7500 and
137 10000 μg/L) in sterilized Milli-Q water. Each tetracycline solution (100 μL) or
138 environmental water sample was mixed with 900 μL LB medium (containing 100 μg/mL
139 ampicillin) in a polyethylene tube. Previously prepared paper strips were inserted in these
140 culture tubes and incubated at 37 °C without shaking for 90 min. Paper strips were then
141 taken out of the culture tubes and prevented from drying. After, 10 μL of X-gal substrate
142 solution (50 g/L) in DMF was added on the biosensor cell spot. The paper strip was
143 shielded from light at 37 °C for 90 min for color development. A Sony α7 III digital
144 camera (Sony, Tokyo, Japan) with Sony shots (20 mm F1.8, Tokyo, Japan) was used for
145 taking RAW images of the strips (Struss et al. 2010).

146 The color intensities, rather than the size of the blue area, an artifact of differences in

147 the dispersive size of the X-gal color developing agent, were measured using the software
148 ImageJ (National Institutes of Health, Bethesda, Maryland, US)
149 (<https://imagej.net/Downloads>) upon acquired digital images above and used to determine
150 tetracycline concentrations. The measurement settings of the images were set to mean gray
151 value in ImageJ that converts 100% white as 255 and 100% black as zero. A rectangle
152 section (1 × 1 mm area) in each image was drawn around a spot on the strip and measured
153 using the selection tool in ImageJ software. A background measurement was implemented
154 of the same size on the bare paper strip, to normalize for slightly different color intensities
155 due to changing illumination while these pictures were obtained in the field. The paper
156 strip biosensor-based calibration was confirmed with the tetracycline concentration
157 quantified by high-performance liquid chromatography (HPLC) (Fig. S1). The limit of
158 detection (LOD) of a biosensor is usually calculated as the concentration at which
159 biosensor signal to noise ratio is above 3. Therefore, while measuring the pixel density
160 changes in samples devoid of tetracyclines (buffer only), biosensor noise was determined
161 and used for the calculation of the LOD for the paper strip biosensor (S/N=3). The
162 linearity ranges of the biosensors paper strip were calculated according to the r values of
163 the standard curves greater than 0.800 (Burrows and Watson 2015). The obtained standard
164 curves were then used for analyses of tetracyclines in water samples (W₁₋₂₀).

165 Twenty water samples were obtained from Nanjing fishponds (Physicochemical
166 properties of the water samples are provided in Table S5 of SI). The sampling sites of
167 water samples were labeled on the map of Nanjing and these samples were mainly
168 collected around Xuanwu lake and Yueya lake of Nanjing (Fig. S4). A random dose of
169 either tetracycline (W₁₋₁₀) and oxytetracycline (W₁₀₋₂₀) was added to the twenty water
170 samples (W₁–W₂₀), which was used to simulate polluted pond water from the input of feed
171 additives used in aquaculture. These samples were aged for 30 days under outdoor

172 conditions prior to analysis. HPLC (LC-20AT, Shimadzu Co., Kyoto, Japan, Detailed
173 process in SI of Fig. S2) was first used to screen and quantify tetracycline concentrations.
174 The estimated concentration of tetracyclines was detected by paper strip and calculated
175 using the fourteen-point paper strip calibration described above.

176 *2.5 Tetracyclines analysis in soil*

177 A volume of 100 μ L of each of the soil matrix calibrants (see Supplemental Text S2
178 for the preparation process) was added to 900 μ L of LB medium in culture tubes in
179 triplicate. The detection procedure was performed as described for water. A dose-response
180 curve using standard tetracyclines solutions prepared in soil extracts was obtained by
181 paper strip biosensors in each analytical run as well as HPLC as a reference for
182 comparison.

183 Twenty tetracycline-contaminated Inceptisol samples were obtained from a test field
184 at Nanjing Agricultural University (Nanjing, China). The physicochemical properties of
185 Inceptisol are given in Table S6. Ten soil samples (S₁₋₁₀) were obtained from the
186 tetracycline test field which received tetracycline exposures for at least one year, and ten
187 soil samples (S₁₁₋₂₀) were obtained from another test field that was contaminated with
188 oxytetracycline for at least one year. Each processed soil extract (100 μ L) (i.e., the EDTA
189 soil extract, see Supplemental Text S3 for the pretreatment process of soils) was analyzed
190 by the methods described previously using the paper strip biosensor. The concentration
191 was then determined by colorimetrics based on the matrix matched calibration curve
192 described above.

193

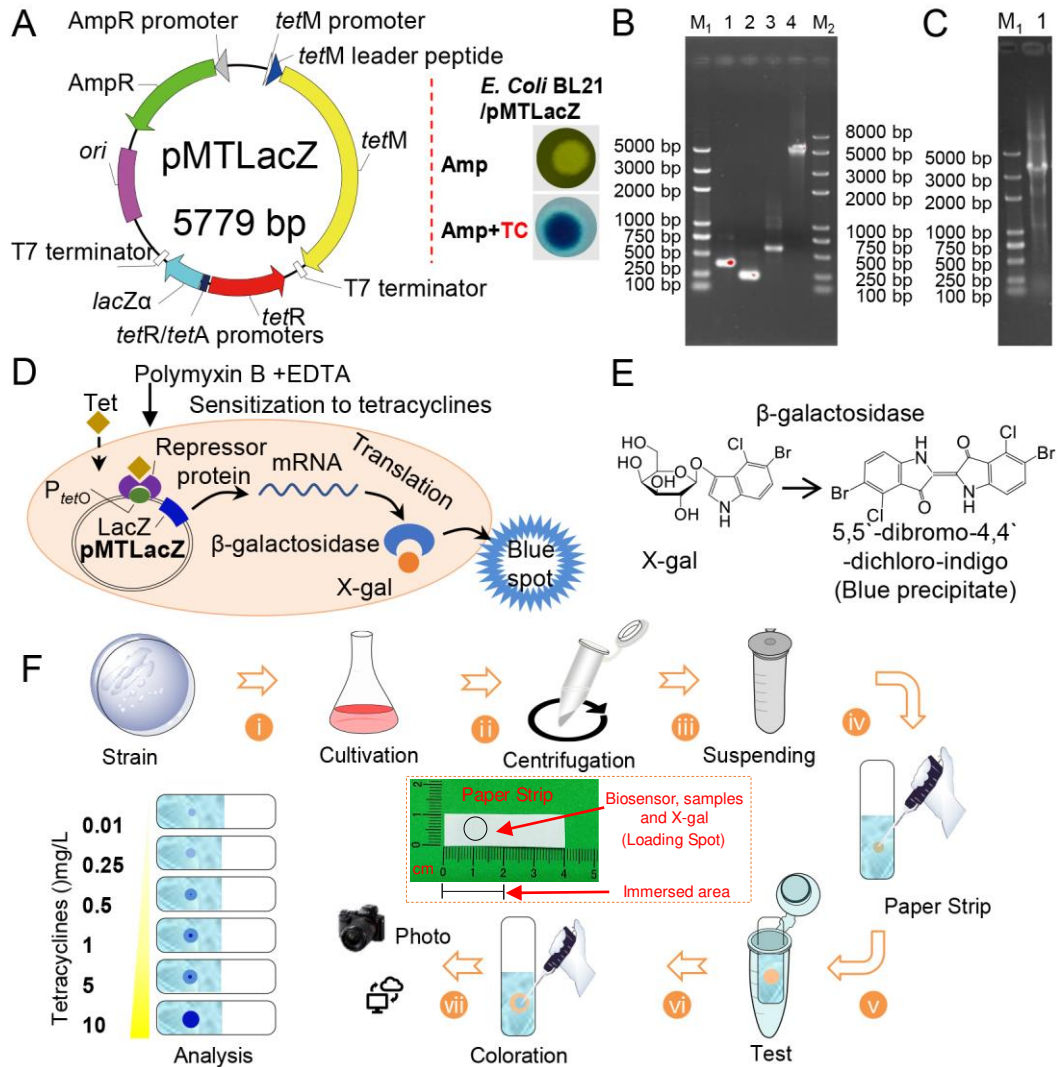
194 **3. Results and discussion**

195 *3.1 Biosensor construction and paper strip production*

196 The schematic of pMTLacZ plasmids used in this study is shown in Fig. 1A. The *lacZ*

197 gene and T7 gene were fused to obtain *lacZ*-T7 gene (Fig. 1B). To construct this biosensor
198 for the sensing of tetracyclines, the pMT gene (Fig. 1B), *lacZ*-T7 gene (Fig. 1B) and T7
199 gene (Fig. 1B) were used to construct the biosensor plasmids by DNA homologous
200 recombination. Additionally, the recombinant plasmid was further verified by PCR with
201 specific primers, and the amplicon lengths were consistent with those expected (pMTLacZ
202 3590 bp) (Fig. 1C). As shown in Fig. 1a, *E. coli* BL21 strain containing a pMTLacZ
203 plasmid produced the blue colony with X-gal when exposed to tetracycline. The
204 expression of the reporter gene is under tight transcriptional control of the tetracycline
205 repressor (*tetR*) on the plasmid (pMTLacZ, Fig. 1A and Fig. 1D). Binding of tetracycline
206 to *tetR* abolishes the binding of this gene to two operator sites (*tetO*) and thus allows
207 expression of the *lacZ* gene (Fig. 1D) and translation of β -galactosidase (Orth et al. 2000).
208 X-gal can be degraded by β -galactosidase, which then produces the blue signal
209 (5,5'-dibromo-4,4'-dichloro-indigo, Fig. 1E). The induction results of tetracyclines
210 revealed that exposure to an increased concentration of antibiotic can increase the enzyme
211 activity response of the biosensor with a dose-related effect (Fig. S3, Detailed illustration
212 in SI).

213 Based on this character of the constructed bacterium, the paper strip biosensor was
214 designed as follows (Fig. 1F). The biosensor cells were cultured in the LB medium,
215 centrifuged and resuspended in 1/4 LB with 10% lactose and PMB (Fig. 1F, step i-iii). The
216 suspension was spotted on Whatman filter paper strips (1 × 4 cm, Fig. 1F, step iv), then
217 dried by vacuum freeze-drying. The paper strip biosensor was immersed in LB broth with
218 samples at 37 °C for 1.5 h (Fig. 1F, step v). Once removed, the color development reagent
219 (X-gal) was added to the biosensor cells (Fig. 1F, step vi) before imaging and post-image
220 processing (Fig. 1F, step vii).



221
 222 **Fig. 1** The construction of the recombinant plasmids. Schematic of the recombinant plasmid
 223 (pMTLacZ) and the fluorescence response of the bacteria (*E. coli* BL21/pMTLacZ) induced by 20
 224 mg/L tetracycline (TC) (A), Agarose gel electrophoresis of fused gene fragment amplified by PCR
 225 (M₁-DNA marker DL5000, 1-*lacZ*, 2-T7, 3-*lacZ*-T7, 4-pMT, M₂-*Trans2K*[®] Plus II DNA Marker) (B),
 226 Agarose gel electrophoresis of cloning gene verified by PCR (M-DNA marker DL5000, 1-pMTLacZ
 227 partial fragment) (C). Genetic organization and mechanism of tetracycline-regulated TC-resistance
 228 determinant (D). The coupled enzyme reactions were catalyzed by β-galactosidase producing the blue
 229 signal (E). Tetracycline semi-quantification processing by whole-cell paper strip biosensor (F).
 230 Biosensor cultivation (i), preparation of paper strip biosensor (ii-iv), Semi-quantitative analysis of
 231 samples using biosensor (v-vii).
 232

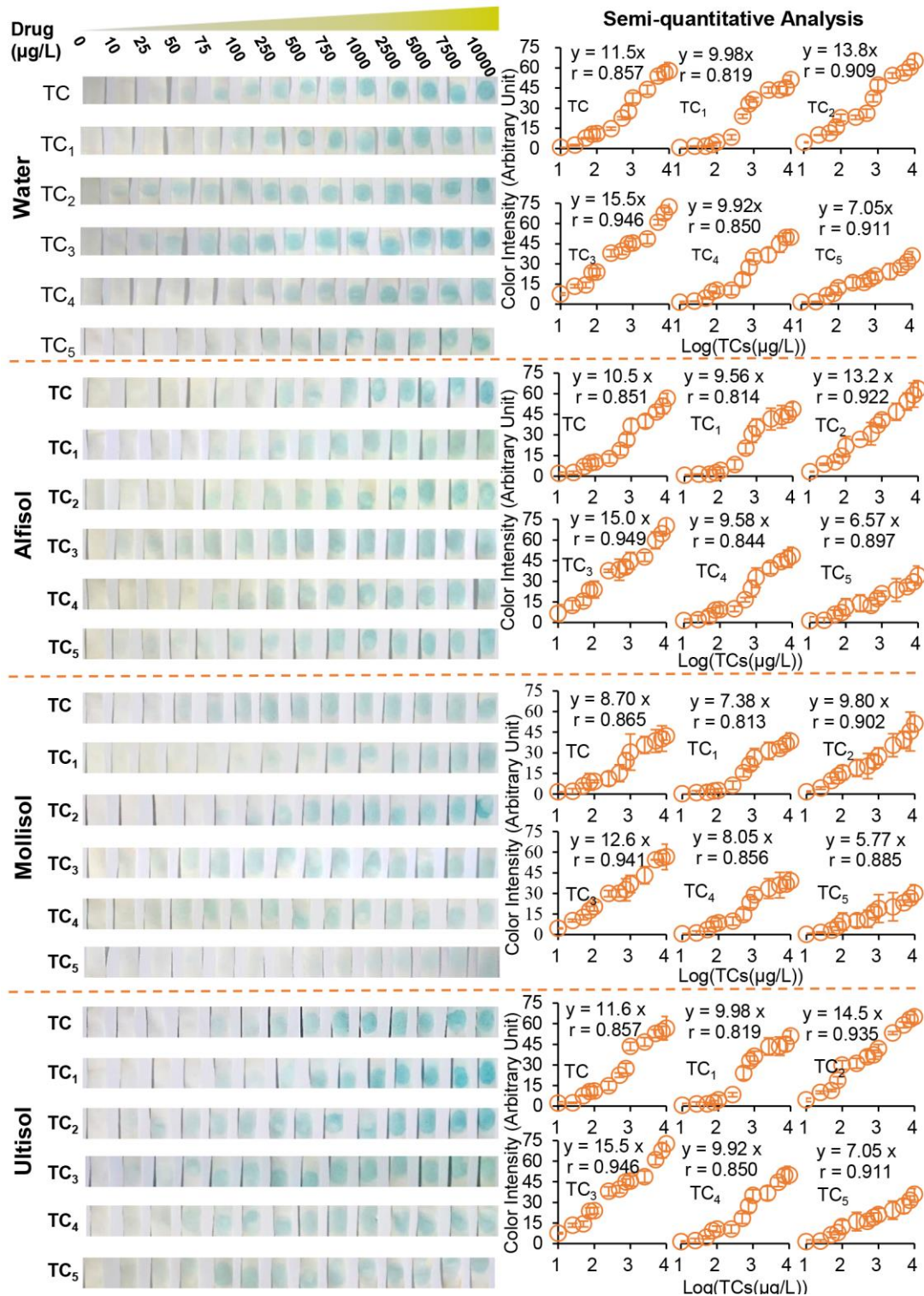
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 240 **Fig. 2** Matrix matched calibration of paper strip biosensors for water and soil extracts and the
 241 correlations between spiked tetracycline concentrations in and color intensity measured using the
 242 software ImageJ upon digital image acquisition. TC, TC₁, TC₂, TC₃, TC₄ and TC₅ denote tetracycline,
 243 oxytetracycline, chlorotetracycline, deoxytetracycline, minocycline, and methacycline, respectively.

244 For semi-quantification of tetracycline concentrations in the samples, a 14-point

245 calibration was prepared for each of the six tetracycline compounds (tetracycline,
246 oxytetracycline, chlorotetracycline, deoxytetracycline, minocycline, and methacycline)
247 spiked in aquafarm water at concentrations ranging from 0–10,000 µg/L. An increase in
248 blue color intensity was observed with increasing concentrations of tetracyclines (Fig. 2).
249 The visual results of the paper strip biosensor were able to indicate low tetracycline
250 concentrations down to 10 µg/L upon color development for 90 min. A good linear
251 relationship between the color intensity and log value of tetracycline concentration in
252 water was observed (Fig. 2, $r > 0.850$, $P < 0.01$). The concentration of each tetracycline in
253 water as a function of the color intensity on the paper strip can be expressed as equation 1
254 and 2 (Table S5, Detailed illustration in SI).

$$255 \quad \log TC = \alpha I \quad (1)$$

256 or

$$257 \quad TC = 10^{\alpha I} \quad (2)$$

258 where TC is the concentration of tetracycline, I is the color intensity detected by the
259 paper strip biosensors and, and α is the slope of standard curve (Values given in Fig. 2).
260 Using the color intensity reading of the biosensors paper strip and Eq. (2), TC can be then
261 calculated by I and α (Fig. 2) obtained from their corresponding standard calibration curve.
262 After 90 min incubation, good linearity in the range of 75–10000 µg/L were found by
263 paper strip biosensors for six tetracyclines (Table 1). The detection limits of the biosensor
264 method in water were between 5.23 and 17.1 µg/L for all types of soils and tetracyclines
265 as determined by S/N ratio. The paper strip biosensor produced a slightly lower detection
266 limit for chlorotetracycline (5.44 µg/L) and deoxytetracycline (5.23 µg/L) than for other

267 tetracyclines in water. In comparison, Zhu et al. (2010) obtained a detection limit of
 268 0.20–0.28 µg/L for tetracyclines in groundwater by HPLC. Although the primary aim of
 269 this study was to develop a paper strip biosensor for the onsite, high-throughput screening
 270 of tetracyclines in water, it is conceivable that the detection limits achieved by HPLC
 271 could also be achieved by the biosensor if concentration of the water samples was
 272 conducted (Zhu et al. 2001).

273 **Table 1** Detection parameters of six tetracyclines measured by biosensors paper strip in water and soil
 274 extracts (Alfisol, Mollisol, and Ultisol)

TCs	Water		Alfisol		Mollisol		Ultisol	
	DL (µg/L)	LR (µg/L)	DL (µg/L)	LR (µg/L)	DL (µg/L)	LR (µg/L)	DL (µg/L)	LR (µg/L)
TC	5.86	25-10000	5.88	25-5000	20.4	50-7500	5.32	50-10000
TC ₁	5.85	75-10000	5.95	75-10000	30.1	75-10000	5.76	75-7500
TC ₂	5.44	25-10000	5.78	25-7500	19.1	10-10000	5.68	25-10000
TC ₃	5.23	10-10000	5.67	25-10000	17.0	10-10000	5.21	25-10000
TC ₄	17.1	75-10000	23.4	25-10000	30.2	25-7500	15.4	50-7500
TC ₅	9.05	75-10000	35.3	10-10000	28.4	25-7500	12.6	10-7500

275 TCs, TC, TC₁, TC₂, TC₃, TC₄ and TC₅ are tetracyclines, tetracycline, oxytetracycline, chlorotetracycline,
 276 deoxytetracycline, minocycline, and methacycline, respectively. DL and LR are detection limit and Linear range,
 277 respectively.

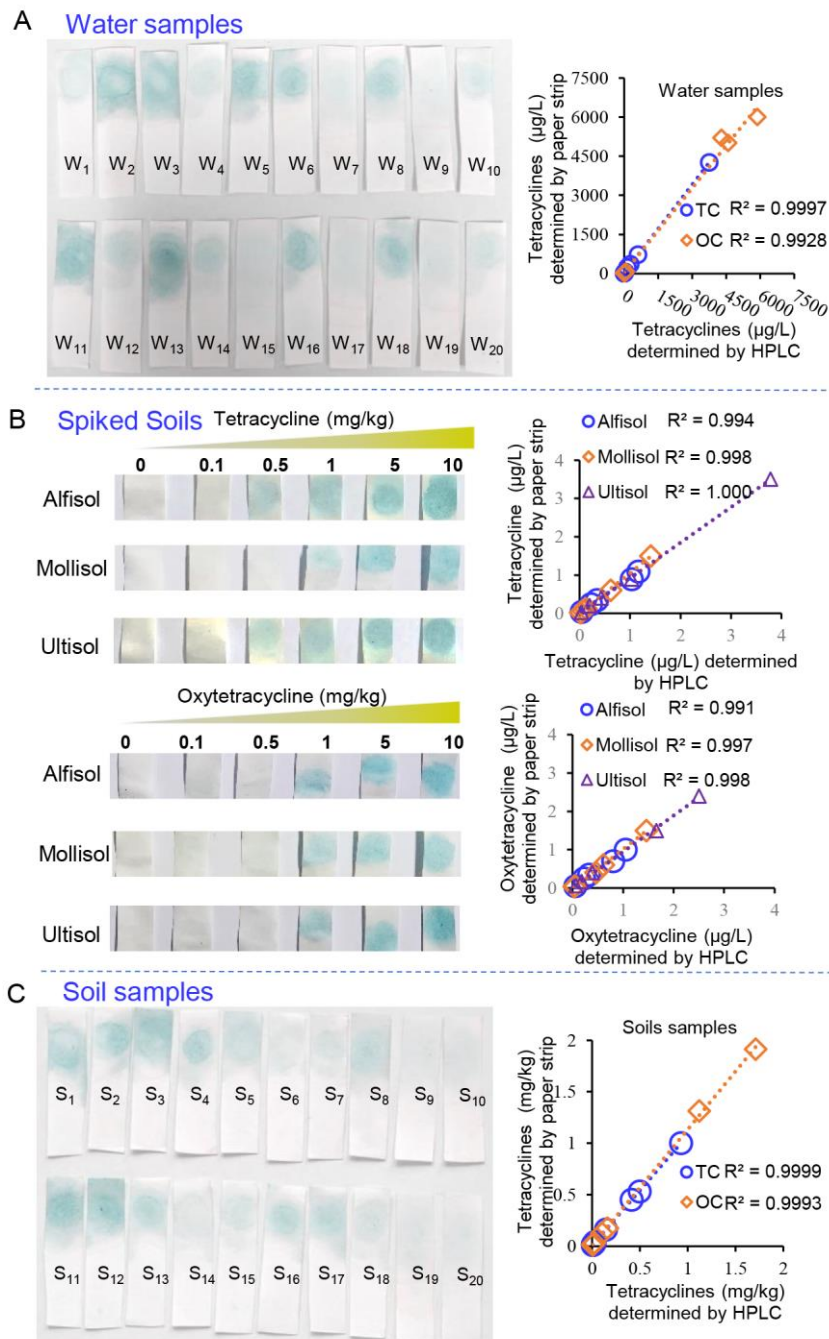
278 The results above demonstrated the capacity of paper strip biosensors for
 279 semi-quantitative detection of tetracyclines in water samples. Extractable fractions of
 280 tetracyclines in soils are often used to assess their bioavailability to biota and their
 281 mobility through soil profiles (Ikehata et al. 2006; Li et al. 2016; Liu et al. 2014). The
 282 EDTA extractable fraction of tetracyclines can be used to evaluate their corresponding
 283 environmental and health risks in soils (Bergan et al. 1973; Cipullo et al. 2018; Hansen
 284 and Sørensen 2001). Therefore, further validation of the dose-response curves of six
 285 tetracyclines from soil extracts using the paper strip biosensor was carried out using a
 286 matrix matched approach of spiked soil extracts from three different soils. The sensitivity
 287 of tetracyclines in soil extracts, as measured by the color intensity of the paper strip

288 biosensor response decreased from Ultisol and Alfisol to Mollisol (Fig. 2). The visual
289 detection limit of the paper strip in soil extracts was between 25 and 100 $\mu\text{g/L}$ for all of
290 the soils and tetracycline compounds. Measurement of the color intensities using
291 appropriate software were shown to enhance sensitivity (Fig. 2). Image analysis indicated
292 that the logarithmic concentration of each tetracycline in the soil extracts as a function of
293 the color intensity followed also followed equation 1 ($r > 0.813$, $P < 0.01$) (Fig. 2). A good
294 linearity in the range of 75–7500 $\mu\text{g/L}$ was determined for the paper strip biosensors for
295 all six of the tetracyclines in the soil extracts (Table 1). The detection limits of
296 tetracyclines in the EDTA-extracts of Inceptisol, Mollisol, and Ultisol were 5.67–35.3,
297 17.0–30.2, and 5.21–15.4 $\mu\text{g/L}$ for this biosensor method (Table 1). Similar to the water,
298 chlorotetracycline and deoxytetracycline showed the lowest paper strip biosensor
299 detection limit in all three soil extracts whereas minocycline and methacycline were
300 highest at 5.21 $\mu\text{g/L}$ in all three soil extracts. Overall, a lower limit of detection was
301 observed in EDTA-extracts from Ultisol than either Inceptisol or Mollisol. The detection
302 limits of the EDTA-extractable tetracyclines in three soils was calculated as $DL = (DL_{SE}$
303 $\times 1 \text{ mL})/1 \text{ g}$ (DL_{SE} denote the detection limit of tetracyclines in soil EDTA-extracts). So,
304 the detection limit of the biosensor method for EDTA-extractable tetracyclines in three
305 soil follow the same principles (5.21–30.2 $\mu\text{g/kg}$, TableS7).

306

307

308



310

311 **Fig. 3** Tetracycline (TC) and oxytetracycline (OC) semi-quantitation in twenty water samples (A) and
312 twenty soil samples (B) and tetracyclines-spiked soils (C) by whole-cell paper strip biosensors. Results
313 represent the average of triplicate assays whereby prepared whole cell paper strip biosensors were
314 incubated in soil extracts at 37 °C and color intensity measured using the software ImageJ upon digital
315 image acquisition.

316 The utility of the paper strip biosensor was verified with the analysis of real water

317 samples. For the samples collected from fishponds, the data (Fig. 3A) were almost
318 identical to those obtained from water samples in laboratory, which indicates the sample
319 matrices had a negligible effect on the sensitivity of the biosensors for tetracyclines. For
320 twenty water samples collected from local fishponds in Nanjing (China), the paper strip
321 biosensor indicated that tetracycline concentrations of W₁–W₁₀ were 45.8, 3756, 609, 6.32,
322 263, 144, 7.75, 15.4, 7.9 and 6.78 µg/L, respectively (Fig. 3A and Table S7). The
323 tetracycline concentrations of W₁–W₁₀ water samples measured by HPLC were 59.7, 4260,
324 718, 9.26, 350, 230, 12.2, 24.3, 11.2 and 10.4 µg/L, respectively, and were slightly higher
325 than the results generated by biosensor method. Standard deviations (between estimated
326 tetracycline concentration by biosensors paper strip and that detected by HPLC, SD) in the
327 analysis of the 10 samples were between 13.4–59.6%. Oxytetracycline concentrations of
328 W₁₁–W₂₀ water samples were 4288, 22.4, 5882, 10.7, 66.8, 4585, 13.7, 26.8, 10.8 and 15.2
329 µg/L, respectively (see Fig. 3A and Table S7). The oxytetracycline concentrations of
330 W₁₁–W₂₀ water samples measured by HPLC were 5201, 30.2, 6000, 14.2, 78.3, 5001, 15.2,
331 30.3, 11.3 and 17.0 µg/L, respectively, and again were higher than the detected values by
332 paper strip. The SD in the analysis of 10 samples contained oxytetracycline were
333 2.01–33.5% (Fig. 3A and Table 2).

334

335

336

337 **Table 2** Tetracycline (TC) and oxytetracycline (OC) semi-quantitative values in twenty water samples and
338 twenty soil samples by whole-cell paper strip biosensors

Sample	Cl	PC (mg/kg)	HC (mg/kg)	SD (%)
W ₁	19.1	45.8	59.7	30.2
W ₂	41.1	3756	4260	13.4
W ₃	32.0	609	718	17.8
W ₄	9.21	6.32	9.26	46.5
W ₅	27.8	263	350	32.9
W ₆	24.8	144	230	59.6
W ₇	10.2	7.75	12.2	57.8
W ₈	13.7	15.4	24.3	57.7
W ₉	10.3	7.90	11.2	42.4
W ₁₀	9.56	6.78	10.4	52.6
W ₁₁	36.3	4288	5201	21.3
W ₁₂	13.5	22.4	30.2	34.9
W ₁₃	37.6	5882	6000	2.01
W ₁₄	10.3	10.7	14.2	33.5
W ₁₅	18.2	66.8	78.3	17.2
W ₁₆	36.5	4585	5001	9.09
W ₁₇	11.3	13.7	15.2	11.4
W ₁₈	14.3	26.8	30.3	13.3
W ₁₉	10.3	10.6	11.3	5.71
W ₂₀	11.8	15.2	17.0	12.0
S ₁	28.3	0.498	0.532	6.86
S ₂	27.5	0.414	0.450	8.72
S ₃	31.2	0.928	0.100	7.78
S ₄	22.7	0.143	0.161	12.4
S ₅	15.3	0.029	0.041	42.5
S ₆	13.7	0.020	0.030	48.3
S ₇	10.3	0.009	0.016	72.8
S ₈	14.2	0.022	0.024	6.22
S ₉	9.5	0.008	0.011	41.3
S ₁₀	8.6	0.006	0.011	58.1
S ₁₁	30.9	1.71	1.91	11.7
S ₁₂	29.2	1.12	1.31	17.2
S ₁₃	13.2	0.024	0.030	25.4
S ₁₄	9.21	0.009	0.010	11.4
S ₁₅	9.79	0.011	0.013	25.3
S ₁₆	19.3	0.011	0.013	19.0
S ₁₇	21.1	0.163	0.172	5.90
S ₁₈	13.0	0.023	0.030	33.7
S ₁₉	7.26	0.006	0.008	43.4
S ₂₀	8.32	0.007	0.011	40.9

339 Color Intensity (CI, Arbitrary Unit), tetracycline concentration measured by paper strip biosensors (PC),
340 concentration determined by HPLC (HC), and standard deviation (SD) between estimated tetracycline
341 concentration, PC, and HPLC determined tetracycline concentration (HC) are presented.

342 When the values calculated by the biosensor were plotted against the HPLC derived
343 concentrations, a strong linear relationship was established. ($R^2 > 0.990$, Fig. 3A). This
344 shows that variation between results obtained from the two methods are not concentration
345 dependent and suggests the potential to correct for the underestimated concentrations
346 observed via biosensor analysis.

347 *3.4 Analysis of extractable tetracyclines fraction in contaminated soil*

348 The paper strip biosensor was also validated for the analysis of the extractable
349 tetracycline fraction in contaminated soils following the same extraction procedure from
350 soil as described above. The results of the analysis in tetracycline and
351 oxytetracycline-spiked soils revealed that the concentrations of extractable tetracycline
352 measured by paper strip biosensors were closer to the values of extractable tetracycline
353 concentration measured by HPLC (Fig. 3B). Two tetracyclines concentrations by
354 whole-cell paper strip biosensors showed a good linear relationship with the
355 concentrations of that of detection by HPLC ($R^2 > 0.991$). The paper strip was further
356 tested to detect tetracycline concentrations in soils sampled from a test field that were
357 contaminated with either tetracycline or oxytetracycline. Using the paper strips, the
358 extractable tetracycline concentration in S₁-S₁₀ soils ranged from 0.006–0.498 mg/kg, (Fig.
359 3C and Table 2), which were slightly lower than the concentrations measured by the
360 HPLC method (0.011–0.532 mg/kg). The extractable oxytetracycline concentrations in
361 S₁₁-S₂₀ soils measured by paper strip biosensors ranged from were 0.006–1.71 mg/kg
362 which were very similar to the concentrations measured by HPLC (0.008–1.91 mg/kg).
363 The SD values of the extractable tetracycline in soils measured by the paper strip

364 biosensor S₁-S₁₀ (6.86–72.8%) were greater than the SD values of the extractable
365 oxytetracycline in S₁₁-S₂₀ soils (5.90–43.4%). However, both SDs provide adequate
366 reproducibility for a semi-quantitative method. The same as for water, a strong linear
367 relationship existed between the concentrations of both compounds determined by paper
368 strip biosensors and HPLC ($R^2 > 0.999$, Fig. 3C and Table 2). Combined with no
369 observable matrix interferences, these results confirm that these paper strip biosensors
370 provide reliable semi-quantitative evaluation of tetracycline concentrations from soil
371 extracts.

372 3.5 Advantages

373 The paper strip biosensors include greater ability to accommodate the variation of
374 environmental conditions in water or soil, such as a wide range of ionic strengths, pH
375 (4–8), and temperature (Iglesias et al. 2009) that is hard to overcome the impact other
376 types of paper strip sensors such as those based on immunochromatographic lateral flow
377 and immobilization of antibodies. This biosensor is more robust and has an excellent
378 capacity to endure various environmental conditions, such as those expected in pond water
379 and agricultural soil where contamination with tetracycline antibiotics is commonly
380 associated with the utilization of feed additives in aquaculture and the use of livestock
381 manure as a soil fertilizer amendment.

382 In this study, the construction of *Escherichia coli*/pMTLacZ for whole-cell biosensor
383 was similar to that of *Escherichia coli*/pMTGFP and *Escherichia coli*/pMTmcherry from
384 our previous study (Ma et al. 2020). These cellular reporters can rapidly and accurately
385 detect tetracyclines in water or soil. However, cost effective and readily available
386 Whatman filter paper was utilized as the carrier for the biosensor scaffolding (Fig. 1),
387 which has substantially greater portability and is more economical than the 96-microwell
388 plate method described in previous studies (Wang et al. 2010; Zou et al. 2018). Biosensor

389 paper strips were produced by a simple culture process (3 hours at 37 °C with vigorous
390 shaking at 150 r/min) in cheap media (Lysogeny Broth), which dramatically reduces
391 analysis cost. This semi-quantitative method has adequate sensitivity and high selectivity,
392 only detecting tetracycline antibiotics, and requires minimal sample pretreatment. This
393 paper strip biosensor offers a lower analysis cost and little instrumental expertise
394 compared to conventional analytical methods (gas chromatography, liquid
395 chromatography, mass spectrometry) (Aga et al. 2016; Arefev et al. 1987). Meanwhile,
396 developing EDTA solvent extraction for the detection of tetracyclines in soils is an
397 inexpensive, easily accessible and environmentally-friendly method compared to many
398 common methods which use organic solvents needed to perform chromatographic
399 separations (Batt and Aga 2005; Hamscher et al. 2005). The combination of EDTA and
400 polymyxin B as the agent to sensitize this material resulted in significantly improve
401 method precision and accuracy (Belkin 2003; Parlanti et al. 2000).

402 At present, portable colorimeters and office scanners pixelated by GIMP software into
403 an RGB profile can evaluate the color intensity of the paper strip, but it's exorbitant for
404 onsite application and not nearly as portable as reagent kits. In our study, optical changes
405 of the sensor were acquired by a digital camera and analyzed by Image J software that
406 enhanced the portability of these paper strip kits while vastly reducing the expense of
407 antibiotic detection. Based on the image processing described in this study, the color
408 analysis software could be developed as a mobile phone APP to directly capture and
409 analyze images and output detection results which would increase user-friendliness while
410 further decreasing analysis time.

411 **4. Conclusion**

412 We have developed a self-contained portable sensing system based on genetically
413 engineered whole-cells physically adsorbed on filter-paper strips for on-site

414 semiquantitative visual monitoring of tetracyclines in environmental samples. Additionally,
415 we demonstrate their ability to provide quantitative measurements by means of digital
416 image analysis. This novel filter-paper strip biosensor method obtained a quantification
417 range of 75–10000 $\mu\text{g/L}$ with detection limits of 5.23–17.1 $\mu\text{g/L}$ for 6 tetracycline
418 antibiotics in water and a detection limit of 5.21–35.3 $\mu\text{g/kg}$ for the EDTA-extractable
419 tetracyclines from three soils. This study describes a fast and convenient method for the
420 detection of tetracyclines, which could be employed for first-level screening of a variety of
421 environmentally and clinically relevant samples. A filter-paper-based biosensor provides
422 easy transportation and storage, and does not require instrumentation or trained personnel;
423 therefore, it could be a component of a simple and inexpensive field kit.

424

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428

429 **Competing interests**

430 The authors declare no competing interests.

431

432 **Data availability**

433 The data that support the findings of this study are available from the corresponding
434 author upon reasonable request.

435

436 **Additional information**

437 Supplementary information is available for this paper at

438

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