

Bacterial host and reporter gene optimization for genetically encoded whole cell biosensors

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Abstract Whole-cell biosensors based on reporter genes allow detection of toxic metals in water with high selectivity and sensitivity under laboratory conditions; nevertheless, their transfer to a commercial inline water analyzer requires specific adaptation and optimization to field conditions as well as economical considerations. We focused here on both the influence of the bacterial host and the choice of the reporter gene by following the responses of global toxicity biosensors based on constitutive bacterial promoters as well as arsenite biosensors based on the arsenite-inducible P_{ars} promoter. We observed important variations of the bioluminescence emission levels in five different *Escherichia coli* strains harboring two different *lux*-based biosensors, suggesting that the best host strain has to be empirically selected for each new biosensor under construction. We also investigated the bioluminescence reporter gene system transferred into *Deinococcus deserti*, an environmental, desiccation- and radiation-tolerant bacterium

that would reduce the manufacturing costs of bacterial biosensors for commercial water analyzers and open the field of biodetection in radioactive environments. We thus successfully obtained a cell survival biosensor and a metal biosensor able to detect a concentration as low as 100 nM of arsenite in *D. deserti*. We demonstrated that the arsenite biosensor resisted desiccation and remained functional after 7 days stored in air-dried *D. deserti* cells. We also report here the use of a new near-infrared (NIR) fluorescent reporter candidate, a bacteriophytochrome from the magnetotactic bacterium *Magnetospirillum magneticum* AMB-1, which showed a NIR fluorescent signal that remained optimal despite increasing sample turbidity, while in similar conditions, a drastic loss of the *lux*-based biosensors signal was observed.

Keywords Arsenite biosensor · Bioluminescence · Near-infrared fluorescent reporter · *Deinococcus deserti* · Desiccation · Bacteriophytochrome

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Introduction

An efficient surveillance of water networks requires automatic monitoring systems able to quickly raise the alarm in the event of an accidental or intentional pollution, reducing the risk of downstream contaminations with appropriate measures implemented as soon as possible. In this context, whole-cell biosensors offer sensitive and rapid alert systems for metal detection that can be integrated into inline water monitoring devices. These biological sensors are not intended to fully replace conventional analytical methods, but they offer the possibility to rapidly alarm the end-user in case of global or specific toxicities affecting the water network and take proper actions before conventional analyses are made for confirmation. Whole-cell biosensors are based on engineered bacteria genetically

transformed with reporter genes whose expression is driven by an inducible promoter; the selected promoter in its native genetic context is involved in the intracellular mechanism of metal resistance (Magrisso et al. 2008; Hynninen and Virta 2010). Typically, the interaction between the metal and a specific repressor protein, otherwise bound to the promoter region on the DNA sequence, results in the release of the repressor from the promoting region, allowing transcription of the reporter gene and detection of the presence or activity of the gene product. Thanks to this genetic modification, these bacterial biosensors are able to emit a semi-quantitative signal in the presence of targeted metals (Xu et al. 2013). Signal transduction by light emission (e.g., bioluminescence or fluorescence) is very popular due to its sensitivity and the variety of reporter genes available (Xiong et al. 2012). Among them, bacterial luciferase (encoded in the *lux* operon) is particularly well suited for the development of autonomous bacterial biosensors as the genes required for the production of the substrate for luciferase can be genetically co-introduced (Prévéral et al. 2016) (this special issue). Extensive searches for different promoters regulated by various metallic targets have been well documented in the literature (Sun et al. 2015); efforts to improve the dose-responses and the specificity by genetic modifications of the regulators have also been described (Peng et al. 2010), and an example is presented in this special issue for a nickel bioluminescent biosensor (Cayron et al. 2015).

Transferring light-emitting bacterial biosensors from the laboratory to on-field water analysis systems requires several optimizations, not all relying on biological research but also including technical or economical considerations. For a given host bacterial species, the genetic background is liable to affect the functioning of the biosensor at various stages, such as intracellular metal concentration, transcription efficiency of the reporter gene as well as the reporter protein expression, folding, and maturation; the integration of these various biological processes affect the biosensor global response in terms of sensitivity, specificity, or time response (Yagur-Kroll and Belkin 2014). In the case of bacterial luciferase, the genetic background may also affect the intracellular production of the luciferase substrate (an aldehyde) as it can also be genetically encoded. Thanks to countless studies and genetic tools available, *Escherichia coli* laboratory strains can conveniently host biosensors based on the reporter gene system but the selection of the proper host strain is critical. Whereas a light-emitting reporter gene system such as the bacterial luciferase can function perfectly well in laboratory conditions on idealized water samples, measurements on real samples taken from ground waters, drainage, industrial effluents, or sewage can be tampered by absorbing and fluorescent materials or particles in suspension. In such cases, it could be wise to use near-infrared (NIR)

light as a detectable signal: in this spectral region, light is less absorbed by organic molecules, more penetrating and less prone to diffusion. NIR-fluorescent reporters derived from bacteriophytochromes—biological light switches—have been efficiently engineered by Shu et al. to the purpose of *in vivo*, non-invasive light imaging where NIR light can be detected through the living tissues (Shu et al. 2009). Such reporters could be useful when the complexity of the environmental matrix prevents an efficient collection of light emitted by whole-cell biosensors usually operating in the visible light spectrum.

The results presented here were obtained during a 4-year translational project called COMBITOX funded by the French Agence Nationale pour la Recherche (ANR). This collaborative project, described in the Editorial of this special issue (Ansaldi et al. 2015), aimed at designing and building a prototype of semi-autonomous inline water analyzer including biosensors to detect pathogenic bacteria with bacteriophages, toxic metals with whole-cell biosensors and environmental toxins with antibodies. Regarding the whole-cell biosensors module, we emphasized in another paper in this special issue on the necessity to develop suitable conservation systems for their integration as bioreagents into a technical device aimed at water analysis (Prévéral et al. 2016); the main challenge here consists in maintaining acceptable performances for the bioreagent in the apparatus compared to the laboratory conditions. We showed that such biosensors can be conditioned by lyophilization for the intended usage, but this technique represents an additional cost in the production process. Unfortunately, a more simple and cost-effective solution such as desiccation proved to be unreliable in our case, probably due to the difficulty for *E. coli* to recover from the stress induced by desiccation. In addition to their intrinsic properties (specificity, sensitivity, time response, reliability), bacterial biosensors for inline water analysis must remain economically competitive with other techniques and this consideration prompted us to probe other microorganisms more amenable to simple conservation techniques such as desiccation.

In this context, we investigate here the response of our biosensors in different strains of *E. coli* in an attempt to identify an optimal strain to host luciferase reporters. We also report here the use of a new NIR fluorescent reporter candidate, a bacteriophytochrome from the magnetotactic bacterium *Magnetospirillum magneticum* AMB-1 that could circumvent the adverse effects of the complexity of environmental water samples on light detection. Lastly, with the objective of simplifying the conservation of the biosensors, we also investigate the transfer of the bioluminescence reporter gene system in *Deinococcus deserti*, a bacterium isolated from the desert and more amenable than are laboratory *E. coli* strains to desiccation.

Material and methods

Strains, plasmids, and primers

Escherichia coli strains used in this study are W3110 (Hayashi et al. 2006), TD2158 (Vinay et al. 2015), and Life Technologies-branded Dh10beta, TOP10 and BL21. *D. deserti* RD19 strain, a streptomycin-resistant mutant of wild-type strain VCD115 (de Groot et al. 2005; Dulermo et al. 2009), was used for desiccation experiments. In this article, when we refer to the L-arabinose-inducible P_{BAD} , arsenite-inducible P_{ars} and mercury-inducible P_{mer} promoters, the corresponding DNA sequences systematically include respectively the *araC*, *arsR*, and *merR* genes encoding the regulatory proteins. The plasmids pRpoD-lux, pMer-lux, and pArs-lux cited in this article have been described in this special issue (Prévéral et al. 2016). The vectors and the PCR primers used here are described in Tables 1 and 2, respectively. The host/vector combinations are listed in Table 3.

Culture conditions for *E. coli* and *D. deserti*

When required, antibiotics were added in the culture media. For *E. coli*, kanamycin, ampicillin, and tetracycline were used at 50, 50, and 10 µg/ml, respectively. For *D. deserti*, kanamycin was used at 10 µg/ml. *E. coli* strains were aerobically cultivated in Lysogeny Broth (LB) medium at 37 °C in

agitated Erlenmeyer flasks (150 rpm). When required, 1 % L-arabinose was added to induce the expression of the genes driven by the P_{BAD} promoter. *D. deserti* strains were aerobically cultivated in ten-fold diluted Tryptic Soy Broth (TSB) at 30 °C in agitated Erlenmeyer flasks (150 rpm).

Lyophilization of *E. coli* cells

E. coli cells were grown aerobically in LB medium at 37 °C under agitation (150 rpm) until early exponential phase ($OD_{600nm}=0.2-0.3$). The cultures were then centrifuged and cells resuspended in LB + 12 % sucrose to an OD_{600nm} equal to 0.5. Aliquots of 500 µl in 2 ml Eppendorf tubes of these bacterial cultures were frozen at −80 °C and then lyophilized for 24 h in a Christ Alpha 1–2 LD Freeze Dryer. The lyophilized aliquots were stored under nitrogen atmosphere at 4 °C until use.

Bioluminescence, NIR fluorescence, and absorbance measurements

Single-point absorbance measurements at 600 nm (OD_{600nm}) were done with a Cary® 50 UV-visible spectrophotometer from Agilent Technologies. Single-point bioluminescence emission measurements described in Fig. 1a were performed on a GloMax® 20/20 luminometer from Promega. For this, *E. coli* cultures were inoculated at $OD_{600nm}=0.01-0.03$ and

Table 1 Plasmids used in this study

Plasmid name	Features	Reference
pPΦ3295	Transcriptional fusion between L-arabinose-inducible P_{BAD} promoter and <i>Rhodopseudomonas palustris</i> apo-bacteriophytochrome gene + <i>hmuO</i> (heme oxygenase), [Amp ^R]	Giraud and Verméglio (2008)
pMagBphP	Transcriptional fusion between L-arabinose-inducible P_{BAD} promoter and <i>magBphP</i> (apo-bacteriophytochrome) + <i>hmuO</i> (heme oxygenase), [Amp ^R]	This study
pArs-lux	Transcriptional fusion between arsenite-inducible P_{ars} promoter and <i>luxCDABE</i> (bacterial luciferase), [Amp ^R]	Prévéral et al. (2016)
pRpoD-lux	Transcriptional fusion between constitutive P_{rpoD} promoter and <i>luxCDABE</i> , [Kan ^R]	Prévéral et al. (2016)
pAAM	Transcriptional fusion between L-arabinose-inducible P_{BAD} promoter and <i>magBphP</i> , [Tet ^R]	This study
pBMH	Transcriptional fusion between mercury-inducible P_{mer} promoter and <i>hmuO</i> , [Amp ^R]	This study
pBAH	Transcriptional fusion between arsenite-inducible P_{ars} promoter and <i>hmuO</i> , [Amp ^R]	This study
pALT	Insertion in <i>D. deserti amyE</i> of the promoter-less <i>luxCDABE</i> + translational fusion $P_{tuf}::kan$; replicates in <i>E. coli</i> [Amp ^R] but not in <i>D. deserti</i> [Kan ^R after integration in chromosome]	This study
pAGLT	In pALT, transcriptional fusion between the constitutive P_{gapdh} promoter and <i>luxCDABE</i> , [in <i>D. deserti</i> Kan ^R , in <i>E. coli</i> Amp ^R]	This study
pAALT	In pALT, transcriptional fusion between the arsenite-inducible P_{ars} promoter and <i>luxCDABE</i> , [in <i>D. deserti</i> Kan ^R , in <i>E. coli</i> Amp ^R]	This study
pBSK-lux-Km	Promoter-less <i>luxCDABE</i> + translational fusion between constitutive P_{tuf} promoter and kanamycin resistance gene, [in <i>E. coli</i> Amp ^R]	This study
pBSK-lux	<i>luxCDABE</i> , [Amp ^R]	This study
pSF0kan	Source of $P_{tuf}::kan$ for <i>D. deserti</i> : translational fusion between constitutive P_{tuf} promoter of <i>D. deserti</i> and kanamycin resistance gene, cloned as BamHI–XbaI fragment in pUC19, [in <i>E. coli</i> Amp ^R]	Laboratory stock
pUC18-AmyEDD	<i>amyE</i> (amylase), [Amp ^R]	This study

Table 2 Primers used in this study

Primer	Sequence	Description
1	GAAGATCTATGACAACGCTGTCCATGTTG	BglII-magBphP forward
2	CGGAATTCTTATAACCATTGGAGCAATGTG	EcoRI-magBphP reverse
3	ACAACGCCCAGATGAGTCTG	amyE forward
4	ACATCGAGCTCATCTTCGCC	amyE reverse
5	ATATTCTAGACCGATACCGGCACGGTAG	amyE-XbaI forward
6	ATATGGGCCCCGTCGAACCATTGCCCCA	amyE-ApaI reverse
7	CGAATGGTTCGACGGGGCCCTTGAGAAGCGCTTGGTG	InFusion gapdh forward
8	TTTTTTTAGTCATACGTACATGCTGTGCCTCCAGTCTTG	InFusion gapdh reverse
9	CGAATGGTTCGACGGGGCCCTAAACTGCCAGGAATTGGGG	InFusion ars forward
10	TTTTTTTAGTCATACGTATAGCAATCTAATTTTTACCGCAG	InFusion ars reverse
11	TGCCATACGGAATTCTAATGTGCCTGTCAAATGGAC	InFusion ara forward
12	TGCTCATCCGGAATTCCTTATAACCATTGGAGCAATGTGT	InFusion ara reverse
13	CGGCCAGTGAGCGCGCCCATGGTAGGGATTTTACCG	InFusion mer forward
14	ATTACGCCAAGCGCGCAATCTTCTCATCCGCCAA	InFusion mer reverse
15	GTATCGATAAGCTTGATATCGAATTCGATTCTAGAAGGAGGTTG	InFusion hmuO forward
16	CAACCTCCTTCTAGAATCGAATTCGATATCAAGCTTATCGATAC	InFusion hmuO reverse

cultivated until the exponential phase ($OD_{600nm}=0.3$ for pRpoD-lux cell-survival biosensor and $OD_{600nm}=0.2$ for pArs-lux arsenite biosensor). We measured the bioluminescence emission after 45 min of incubation at 30 °C in 2-ml Eppendorf tubes containing 500 μ l of *E. coli* culture, contaminated with 2.7 μ M arsenite for pArs-lux biosensor. In general, with the multi-mode micro-plate reader Infinite® 200 Pro from TECAN, the kinetics were measured at 30 °C under

regular shaking in sterilized Greiner 96-well black plate and clear bottom, with the apparatus set at 450 nm for bioluminescence, excitation at 685 nm, and emission at 755 nm for NIR fluorescence and 600 nm for absorbance. In these different experiments, for both *E. coli* and *D. deserti*, each well in the micro-plate contained 100 μ l of cell culture suspension and 100 μ l of a solution containing the additives when required (antibiotics, L-arabinose, sodium arsenite, or kaolin). As the

Table 3 Host/vector combinations used in this study

Strain	Vector	Promoter	Signal	Description	Figure
<i>E. coli</i> TD2158	pRpoD-lux	P _{rpoD}	Bioluminescence	Cell survival biosensor	1
<i>E. coli</i> W3110	pRpoD-lux	P _{rpoD}	Bioluminescence	Cell survival biosensor	1
<i>E. coli</i> DH10beta	pRpoD-lux	P _{rpoD}	Bioluminescence	Cell survival biosensor	1
<i>E. coli</i> TOP10	pRpoD-lux	P _{rpoD}	Bioluminescence	Cell survival biosensor	1
<i>E. coli</i> BL21	pRpoD-lux	P _{rpoD}	Bioluminescence	Cell survival biosensor	1
<i>E. coli</i> TD2158	pArs-lux	P _{ars}	Bioluminescence	Arsenite biosensor	1, 4
<i>E. coli</i> W3110	pArs-lux	P _{ars}	Bioluminescence	Arsenite biosensor	1
<i>E. coli</i> DH10beta	pArs-lux	P _{ars}	Bioluminescence	Arsenite biosensor	1
<i>E. coli</i> TOP10	pArs-lux	P _{ars}	Bioluminescence	Arsenite biosensor	1
<i>E. coli</i> BL21	pArs-lux	P _{ars}	Bioluminescence	Arsenite biosensor	1
<i>D. deserti</i> RD19	pALT	None	Bioluminescence		2
<i>D. deserti</i> RD19	pAGLT	P _{gapdh}	Bioluminescence	Cell survival biosensor	2
<i>D. deserti</i> RD19	pAALT	P _{ars}	Bioluminescence	Arsenite biosensor	3–5
<i>E. coli</i> DH10beta	pMagBphP	P _{BAD}	NIR fluorescence	Cell survival biosensor	7, 8
<i>E. coli</i> DH10beta	pBAH + pAAM	P _{ars} + P _{BAD}	NIR fluorescence	Arsenite biosensor	9

For *Escherichia coli* strains, vectors are replicative plasmids; for *Deinococcus deserti*, vectors are inserted within the amylase gene by homologous recombination. The pRpoD-lux and pArs-lux vectors have been described elsewhere (Prévéral et al. 2016)

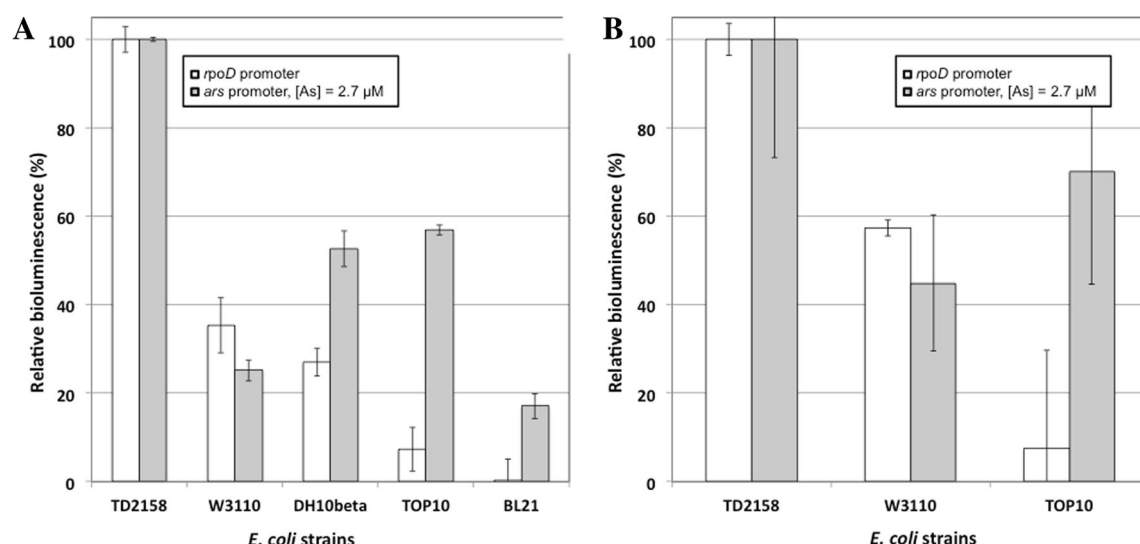


Fig. 1 Impact of the *Escherichia coli* host strain on the biosensor relative bioluminescence emission. **a** Measurements made on growing cells after 45 min incubation. 2.7 μ M arsenite was added to the pArs biosensor at $t = 0$ mn. **b** Measurements made on revived lyophilized cells after 60 min incubation. 2.7 μ M arsenite was added to the pArs biosensor at $t = 0$ mn. For a given biosensor, the signal is set at 100 % for the strain exhibiting

the maximum bioluminescence emission level. *rpoD* (white square) is the cell-survival biosensor and *ars* (gray square) is the arsenite biosensor. The *E. coli* strains used here are the wild-type strain TD2158 and the engineered strains W3110, Dh10beta, TOP10, and BL21. Measurements were done in triplicates (standard errors shown in panels **a** and **b**)

preparation details for the 100- μ l cell suspension depended on the strain and the type of experiment performed, they are summarized in Table 4.

Bioinformatics

For genomic analysis, the genome sequence of *M. magneticum* AMB-1 (Matsunaga et al. 2005) was imported into the microbial genome expert annotation system MaGe (Vallenet et al. 2006) provided by the Génoscope (<https://www.genoscope.cns.fr>). We identified a previously undetected gene, which we named *magBphP*, encoding a putative bacteriophytochrome. The gene sequence was deposited in Genbank (no. BK009263) as a third party annotation.

Bacteriophytochrome expression and purification

Giraud et al. designed the PPΦ3295 plasmid for heterologous expression in *E. coli* of functional bacteriophytochromes from *Rhodospseudomonas palustris* (Giraud and Verméglio 2008). Briefly, the apo-bacteriophytochrome gene is cloned between the BglII and EcoRI restrictions sites, resulting in addition of a 6-histidine tag at the N-terminus of the protein; it is followed by the *hmuO* gene from *Bradyrhizobium* sp. ORS 278 that encodes the heme oxygenase required to oxidize a heme into biliverdin. The expression of both genes is driven by the L-arabinose inducible promoter *araBAD* (P_{BAD}) (Guzman et al. 1995). We amplified the *magBphP* gene by high-fidelity PCR from *M. magneticum* AMB-1 genomic DNA (primer nos. 1 and 2); this PCR fragment was then digested with BglII and EcoRI restriction enzymes and cloned into the PPΦ3295 after

Table 4 Cell suspension preparation for *E. coli* and *D. deserti* prior to absorbance, bioluminescence, and NIR-fluorescence kinetics with the multi-mode micro-plate reader Infinite® 200 Pro from TECAN

Strain	Figure	Cell preparation for 96-wells micro-plates
<i>Escherichia coli</i>	1B	Lyophilized aliquot resuspended in 500 μ l of water and incubated 20 min at 30 °C
<i>Deinococcus deserti</i>	2, 3	Inoculated from overnight culture.
<i>Deinococcus deserti</i>	5A	Overnight culture inoculated at $OD_{600nm} = 0.01$, harvested in exponential phase at $OD_{600nm} \approx 0.1$ and resuspended at $OD_{600nm} = 1.2$
<i>Deinococcus deserti</i>	5B, 5C	Air-dried cells resuspended in TSB/10 at $OD_{600nm} = 1.2$
<i>Escherichia coli</i>	7–9	Inoculation at $OD_{600nm} = 0.01–0.03$, harvested in early exponential phase at $OD_{600nm} \approx 0.25$ and resuspended at $OD_{600nm} = 1.2$

Overnight cultures are always prepared prior to the experiments in the prescribed media with the required additives when needed (antibiotics, L-arabinose), serving as inoculum

digestion by the same restriction enzymes and excision of the *R. palustris* bacteriophytochrome gene, yielding the pMagBphP plasmid. The latter was then introduced into *E. coli* Dh10beta. For protein expression, two 3-L Erlenmeyer flasks containing 500 ml of Terrific Broth medium (Life technologies) were inoculated (1/100) with an overnight pre-culture of *E. coli* transformed with pMagBphP plasmid. Cells were grown at 37 °C, 150 rpm agitation, until OD_{600nm} 0.6 was reached. Protein expression was then induced by the addition of 0.1 % (w/v) of L-arabinose (Sigma-Aldrich), bacterial growth taking place now at 25 °C overnight. Cells of emerald green color were harvested (7500g, 10 min, 4 °C), washed twice in buffer A (20 mM phosphate buffer, 500 mM NaCl, pH 8), and finally resuspended in 30 ml of the same buffer. DNase I and protease inhibitor cocktail (Sigma-Aldrich) were added prior to cell disruption by French Press (three times, 1000 psi, 4 °C). Unbroken cells and debris were removed by centrifugation (7500g, 10 min, 4 °C). Ultracentrifugation of the supernatant (45,000g, 90 min, 4 °C) yielded 24 ml of a dark emerald green soluble fraction. After filtration on a 0.22-μm disposable filtration unit, this supernatant was loaded (1 ml min⁻¹) onto a 5-ml HisTrap Nickel affinity column (Amersham) equilibrated with buffer A. The column was washed with 50 ml of buffer A, then 50 ml of buffer A + 50 mM imidazole. The bound protein was eluted by 25 ml of buffer A + 125 mM imidazole and concentrated to a final volume of 2.5 ml on a Centricon NW30 unit (Millipore). The entire protein sample was then loaded onto a desalting PD-10 column (Life Technologies) equilibrated with Buffer B (20 mM HEPES, pH 8) then eluted with 3.5 ml of the same buffer.

Construction of bioluminescent biosensors in *D. deserti*

The genetic modification of *D. deserti* was obtained by chromosomal insertion of constructed cassettes. For the insertion location on the bacterial chromosome, we selected the amylase-encoding *amyE* gene (gene ID *Deide_08810*), which is not essential for growth of *D. deserti*. We designed three bioluminescent strains where the *luxCDABE* operon encoding the bacterial luciferase from *Photorhabdus luminescens* (Winson et al. 1998) is controlled by a *D. deserti* constitutive promoter P_{gapdh} (from gene *Deide_11420* encoding glyceraldehyde-3-phosphate dehydrogenase) (plasmid pAGLT), by an *E. coli* arsenite-inducible promoter P_{ars} (plasmid pAALT) or is left promoter-less (plasmid pALT). The pUC18 plasmid, not replicative in *D. deserti*, was used as a backbone (Dulermo et al. 2009). We first amplified by high-fidelity PCR the *amyE* sequence from *D. deserti* genomic DNA (primer nos. 3 and 4) and cloned it into pUC18 (Norlander et al. 1983) between HincII and SacI restriction sites, yielding the pUC18-amyEDD plasmid. We obtained the *luxCDABE* operon by enzymatic restriction (SnaBI and

BamHI) from the plasmid pArs-lux and cloned it into a pBluescript II SK(+) from Agilent Technologies (pBSKII) at the corresponding sites, yielding the pBSK-lux plasmid. The *Ptuf::kan* fragment, comprising the kanamycin resistance gene under the control of the *D. deserti* P_{tuf} promoter (Dulermo et al. 2009), was obtained by enzymatic restriction (BamHI and XbaI) of pSF0kan and cloned at the corresponding sites in pBSK, yielding the pBSK-lux-*Ptuf::kan* plasmid. We then amplified the entire pUC18-AmyEDD with a primer pair (primer nos. 5 and 6) introducing ApaI and XbaI restriction sites in the middle of the *amyE* sequence. After enzymatic restriction by ApaI and XbaI, this PCR product was ligated to the *luxCDABE* + *Ptuf::kan* cassette obtained by enzymatic restriction (ApaI and XbaI) of pBSK-lux-*Ptuf::kan*, yielding the pALT plasmid (promoter-less expression of *lux*). Insertions of P_{gapdh} and P_{ars} promoters were realized by the In-Fusion® HD cloning technique (Clontech) in pALT digested with ApaI and SnaBI restriction enzymes, yielding pAGLT and pAALT plasmids, respectively. P_{gapdh} and P_{ars} promoters were obtained by PCR from *D. deserti* genomic DNA (primer nos. 7 and 8) and pArs-lux plasmid (primer nos. 9 and 10), respectively. The insertion of the various *lux* cassettes in *D. deserti*'s genome was obtained via double homologous recombination in *D. deserti* genome at the *amyE* locus after transformation of the cells with pALT, pAGLT, or pAALT. Transformation of chemically competent cells was performed as described (Dulermo et al. 2009). The double homologous recombination at the correct locus and insertion of the cassette of interest were systematically confirmed by diagnostic PCR.

Cell survival after desiccation of *E. coli* and *D. deserti* containing the arsenite P_{ars} biosensor

We transferred the corresponding arsenite biosensors into *E. coli* TD2158 and *D. deserti* RD19. For desiccation, exponential (OD_{600nm} = 0.1 for *D. deserti* and 0.6 for *E. coli*) phase cells were concentrated to an OD_{600nm} = 20 and placed on sterile glass slides (100 μl cells per slide), dried and stored at room temperature in a sealed desiccator over anhydrous CaSO₄. The CaSO₄ desiccant is impregnated with CoCl₂. This latter chemical compound is a visual moisture indicator: anhydrous CoCl₂ is blue while pink in the presence of water. After storage at room temperature for 1 and 7 days, cells were resuspended in TSB/10 (*D. deserti*) or LB (*E. coli*), serially diluted, and plated. After growth, colonies were counted to determine survival compared to non-desiccated cells.

Construction of a near-infrared fluorescent biosensor in *E. coli*

The NIR fluorescent biosensor relies on two plasmids co-introduced into *E. coli*. The pAAM plasmid comprises the *M. magneticum* AMB-1 bacteriophytochrome structural gene

magBphP whose expression is driven by the L-arabinose-inducible P_{BAD} promoter. The pBAH plasmid includes the heme oxygenase gene *hmuO* from *Bradyrhizobium sp.* ORS 278 under the control of the arsenite-inducible P_{ars} promoter from *E. coli*. Plasmid pAAM was constructed by cloning the fragment containing the regulatory gene *araC*, the P_{BAD} promoter and *magBphP*, obtained from pMagBphP by high-fidelity PCR (primer nos. 11 and 12), into pACYC184 plasmid (Rose 1988) at the EcoRI restriction site by In-Fusion® HD cloning technique (Clontech). An intermediary plasmid pBMH was constructed by In-Fusion® HD cloning technique. The enzymatic restriction of plasmid pBSKII(+) by BssHII produced Fragment no. 1; the sequence containing the mercury-inducible P_{mer} promoter was obtained by high-fidelity PCR (primer nos. 13 and 14) from pMer-lux plasmid, yielding Fragment no. 2; and the *hmuO* gene sequence was obtained by high-fidelity PCR (primer nos. 15 and 16) from pMagBphP, yielding Fragment no. 3. The ordered assembly of Fragment nos. 1, 2, and 3 was realized according to the manufacturer protocol. The pBMH plasmid was then digested with NcoI and ApaI restriction enzymes to remove the P_{mer} promoter, yielding Fragment no. 4; the sequence containing the arsenite-inducible P_{ars} promoter was obtained by high-fidelity PCR from pArs-lux and ligated to Fragment no. 4 with the In-Fusion® HD cloning technique, yielding plasmid pBAH. After verification by DNA sequencing, plasmids pAAM and pBAH were co-introduced into *E. coli* Dh10beta chemically competent cells.

Nephelometry

A kaolin stock solution at 4000 Nephelometric turbidity units (NTU) was prepared by dissolving 400 mg of kaolin into 100 ml of water. Serial dilutions were made in water to obtain a range of 0–500 NTU. For bioluminescence and NIR fluorescence assays, we used the cell-survival pRpoD-lux biosensor in *E. coli* TD2158 and pMagBphP in *E. coli* Dh10beta, respectively. Absorbance, bioluminescence, and NIR fluorescence kinetics were then recorded as described above.

Results

Influence of the host strain for a bioluminescent bacterial biosensor

Are there optimal E. coli strains to harbor bacterial biosensors?

Each academic laboratory involved in project COMBITOX used its own selection of *E. coli* strains to host its biosensors and it became fairly evident that the genetic background had an impact on the bioluminescence emission. In order to

validate general rules relevant for the construction and optimization of new whole-cell detectors within the COMBITOX framework, we measured the bioluminescence emission in 5 different *E. coli* strains (TD2158, W3110, DH10beta, TOP10 and BL21) harboring 2 different *lux*-based biosensors. In the cell-survival pRpoD-lux biosensor, the *lux* operon is transcriptionally fused to the constitutive P_{rpoD} promoter, whereas pArs-lux is a metal biosensor where the *lux* operon is transcriptionally fused to the P_{ars} (arsenite-inducible) promoter. Both pRpoD-lux and pArs-lux biosensors have been fully characterized and are published in this special issue (Prévéral et al. 2016).

In Fig. 1a, we summarized the relative bioluminescence levels that were measured with growing *E. coli* cells. Depending on the *E. coli* host strain, we observed important variations of the bioluminescence emission levels for each biosensor that was tested 45 min after the beginning of the experiment. At this time, the bioluminescence emission is maximum for all strains tested and the emission kinetics are similar, with no lag phase recorded (data not shown). Variations are the most important for the cell-survival pRpoD-lux biosensor, with the maximal signal obtained in TD2158 (100 %), only 7 % remaining in TOP10 and almost nothing measured in BL21. For the arsenite biosensor, there is more than a five-fold decrease of light emission when transferred from TD2158 to BL21. To summarize, the wild-type *E. coli* TD2158 strain would be a reasonable compromise in living cells with optimum bioluminescence emission measured for both the cell survival and the arsenite biosensor; nevertheless, we observed in another context on fresh cells for a cadmium biosensor designed on the same scaffold as P_{rpoD} and P_{ars} that TD2158 comes in a second position after TOP10 (unpublished data). Lyophilized cells when revived and eventually challenged with arsenite do follow the same trend as the non-lyophilized cultures (Fig. 1b), although with a 15-min delay when compared to the fresh cells (Fig. 1a). Indeed, the emission kinetics are similar for all strains but this time with an initial lag phase resulting in a maximum bioluminescence emission observed 60 min after the beginning of the experiment (data not shown). These kinetics indicate that the global effect of the lyophilization procedure on *E. coli* is similar regardless the strain or the biosensor. All together, these observations suggest that there is not a universal, optimal host strain within the same species, each bioluminescent biosensor requiring a separate optimization.

Is an environmental desiccation-resistant strain (D. deserti) more adapted to harbor biosensors?

We further investigated the influence of the cellular chassis by constructing a cell survival and an arsenite biosensor in *D. deserti*, a bacterium isolated from a desert environment (de Groot et al. 2005) exhibiting a high resistance to DNA-

damaging stresses such as UV and gamma irradiation as well as desiccation. This bacterial strain is thus likely to be more resistant than are laboratory *E. coli* strains to on-field measurements on environmental samples. Keeping in mind the economical constraints on the production of biosensors as reagents for water analyzer, *D. deserti* would also be a good candidate to host biosensors with desiccation as a simple, cost-effective conservation process.

We intend here to demonstrate the feasibility of a bioluminescent arsenite sensor hosted in *D. deserti* derived from the pArs-lux biosensor previously hosted in *E. coli*. To start with, this proof-of-concept requires evaluating the ability of *D. deserti* to express a functional, bioluminescent bacterial luciferase. To address this, we inserted the transcriptional fusion between the *D. deserti gapdh* promoter (P_{gapdh}) and the *luxCDABE* operon into the amylase gene (*amyE*), which is not essential for aerobic growth of *D. deserti* (Fig. 2). The *gapdh* gene is among the most highly expressed genes in *D. deserti* on the basis of RNAseq data (de Groot et al. 2014). The bioluminescence emission kinetics shown in Fig. 2 demonstrate the ability of *D. deserti* to produce both the bacterial luciferase and its substrate under the supervision of the P_{gapdh} promoter, although the time course of bioluminescence emission is slow. Little background bioluminescence was observed in the absence of any promoter fused to the *lux* operon.

We then replaced P_{gapdh} by the arsenite-inducible P_{ars} promoter in the chromosomal insertion and measured the

bioluminescence emission after arsenite addition (Fig. 3). This genetic modification led to a functional biosensor that emits light in a dose-response manner to arsenite, although the P_{ars} promoter originates from a distantly related bacterium. As observed with pArs-lux in *E. coli* (Prévéral et al. 2016), the baseline bioluminescence is significantly high without arsenite added, most probably due to an incomplete binding of the ArsR repressor to the *ars*-promoting region. Nevertheless, a concentration as low as 100 nM arsenite results in a significant increase of bioluminescence emission with respect to this baseline level (Fig. 3a). Saturation is reached between 5 and 10 μM , corresponding to a range of arsenite concentrations where *D. deserti* growth begins to be affected (not shown). The time response of this arsenite biosensor in *D. deserti* is slow as shown in Fig. 3b: it takes between 10 and 15 h to be sensitive enough to detect small variations of arsenite concentrations. This is 10 to 20 times slower than in *E. coli* where reliable measurements can be obtained between 30 and 60 min after contamination. As a conclusion, the time response in *D. deserti* is slow whatever the promoter (P_{gapdh} or P_{ars}) and this may be related to the slower cell cycle of *D. deserti* and a lower expression.

We completed the proof-of-concept of a *D. deserti*-based arsenite biosensor by investigating the bioluminescence recovery and response to arsenite with air-dried *D. deserti* cells. Using a very simple, non-optimized desiccation procedure applied to the arsenite biosensor hosted both in *E. coli* and *D. deserti*, we showed that the cell survival rate was more than five orders of magnitude higher for *D. deserti* than for *E. coli* after 7 days of desiccation (Fig. 4). One has to note that the 4 % cell survival rate observed for the genetically modified *D. deserti* strain after 7 days is significantly lower than for the WT strain (50 %) measured 40 days after desiccation in another laboratory (de Groot et al. 2009). We believe that the genetic modification and the use of antibiotics likely impacts the cell fitness, hence its resistance to desiccation, and justifies a separate study for optimization of the desiccation protocol. Moreover, we used in this study the simplest procedure to air-dry the cells and different parameters can be adjusted to improve the cell survival rate such as temperature, evaporation kinetics, protectants, or vacuum drying. Figure 5 depicts the bioluminescence emission kinetics recorded after addition of increasing concentrations of arsenite to fresh *D. deserti* cells (Fig. 5a) and to cells revived 1 (Fig. 5b) and 7 days (Fig. 5c) after desiccation. In all cases, the arsenite biosensor responds in a similar fashion to arsenite concentrations ranging from 0.1 to 5 μM (Fig. 5d). For *E. coli*, little or no bioluminescence could be reliably observed (data not shown). We thus observe that desiccation preserves the sensitivity and the dynamic range of the bioluminescence emission in desiccated *D. deserti* cells but also impacts the time response of the biosensor. Indeed, we measured similar bioluminescence

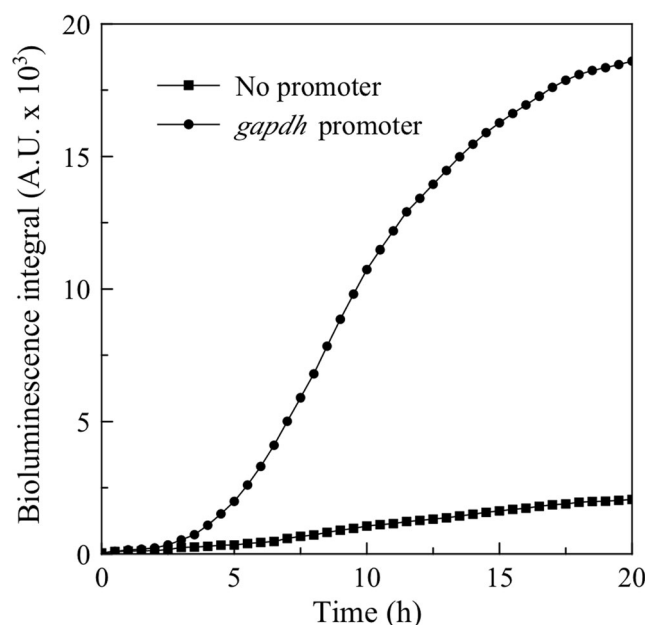


Fig. 2 Bioluminescence emission kinetics in *Deinococcus deserti*. Chromosomal insertion within the *amyE* gene of the *lux* operon without promoter (black square) or under the control of P_{gapdh} promoter (black circle)

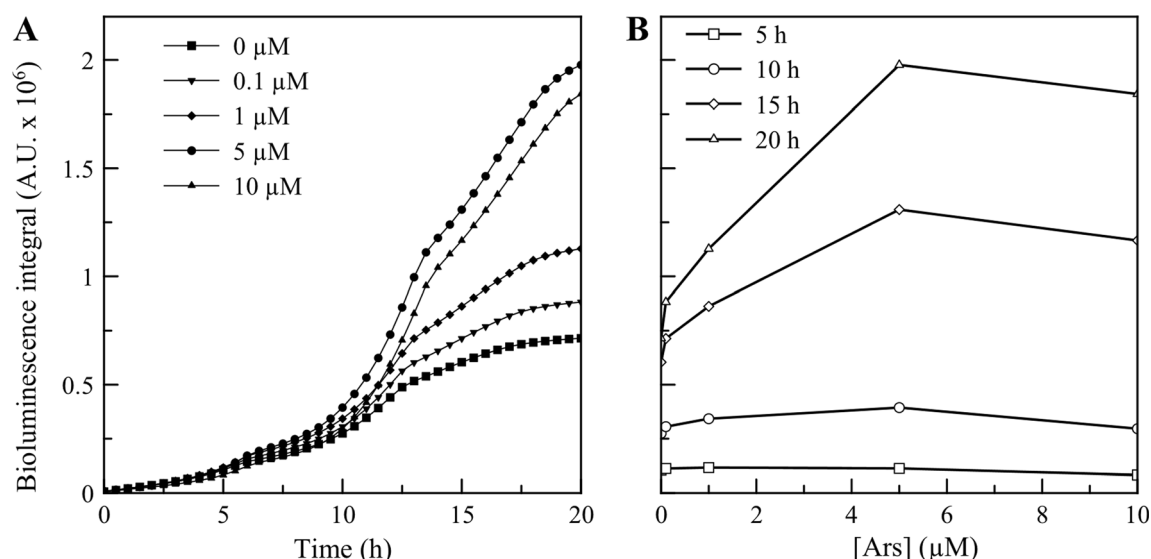


Fig. 3 Arsenite-induced bioluminescence emission in recombinant *Deinococcus deserti*. **a** Integral of the bioluminescence emission over time for increasing concentrations of arsenite. **b** Bioluminescence dose

response at different sampling times (5, 10, 15, and 20 h) after addition of arsenite (0, 0.1, 1, 5, and 10 μM). Initial $\text{OD}_{600\text{nm}}$ is equal to 0.1 and arsenite is added at $t = 0$ h

emission levels with 5- μM arsenite added for the fresh and desiccated cells but with a 5 and 7 h delay for the 1- and 7-day desiccated cells, respectively. As a conclusion, the proof-of-concept of a bioluminescent arsenite biosensor with *D. deserti* is sound, although liable to optimization (codon usage, insertion locus, desiccation procedure), and a desiccation-resistant strain such as *D. deserti* remains a promising candidate to host bioluminescent biosensors.

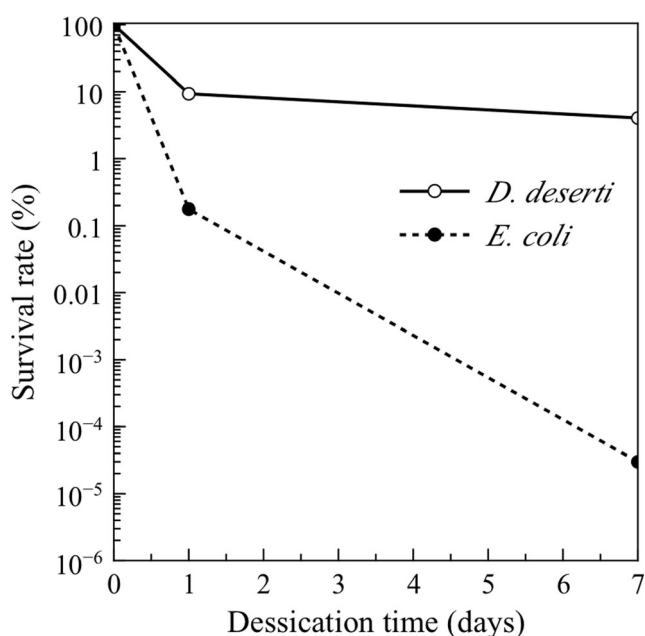


Fig. 4 *Deinococcus deserti* resistance to desiccation. Comparison of bacterial survival after desiccation for the arsenite biosensors hosted in *D. deserti* RD19 (white circle) and *E. coli* TD2158 (black circle). The 100 % values correspond to 8.5×10^6 and 3.3×10^7 colony forming units for *D. deserti* and *E. coli*, respectively

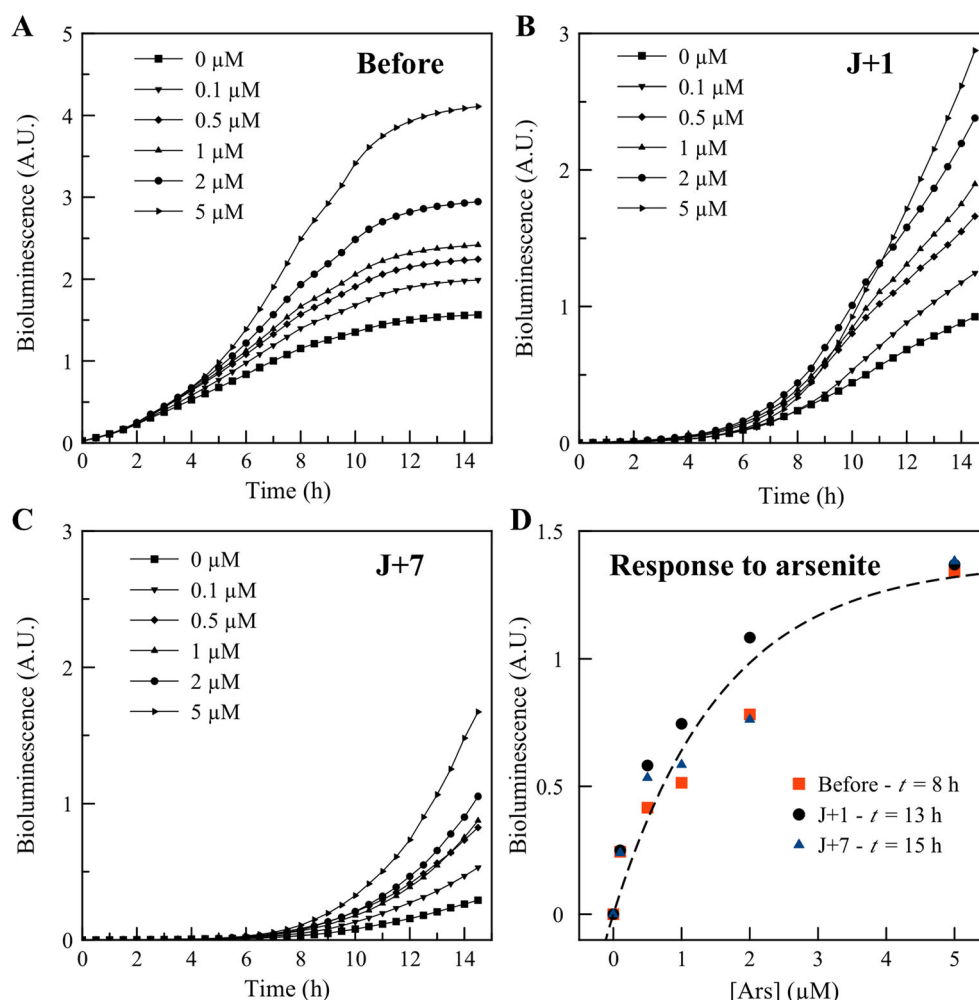
Bacteriophytochromes as near-infrared fluorescent reporters for light biosensors

As we previously evidenced, the whole-cell biosensor response is affected by the choice of the bacterial host and it is also strongly influenced by the selection of the reporter gene. This is notably true in the context of using biosensors under real life conditions, where environmental samples can be much more complex than in laboratory experiments; for instance turbidity and absorbing materials can drastically affect the emission and collection of bioluminescence emitted in *lux*-based whole-cell biosensors. In an attempt to propose an alternative to luciferase in such conditions, we turned to the characterization and the use of a near-infrared fluorescent reporter, the bacteriophytochrome MagBpHP we identified while re-annotating the genome of the magnetotactic bacterium *Magnetospirillum magneticum* AMB-1. At variance with most bacteriophytochromes identified to date, MagBpHP is naturally restricted to the chromophore-binding domain, with no transducing domain fused at the C-terminus as customary (Giraud and Verméglio 2008; Sharrock 2008). For the record, the engineered bacteriophytochrome from *Deinococcus radiodurans* used as a reporter for in vivo imaging in rodents is truncated to the same size and functional domain as MagBpHP (Shu et al. 2009).

In vitro characterization of a near-infrared fluorescent bacteriophytochrome

The recombinant histidine-tagged MagBpHP protein was purified on Ni-NTA column (Fig. 6a), yielding an emerald green-colored protein typical of biliverdin-binding NIR light

Fig. 5 Bioluminescence emission kinetics of *D. deserti* pArs biosensor upon addition of arsenite. Integration over time of the ratio (Bioluminescence emission/OD_{600nm}) for **a** fresh cells, **b** cells revived after 1 day of desiccation, and **c** cells revived after 7 days of desiccation. **d** Biosensor's response to arsenite for fresh (red square), 1-day desiccated (black circle), and 7-day desiccated (blue triangle) cells. In each case, we selected the sampling time where equivalent bioluminescence increases were observed with 5- μ M arsenite added with respect to the baseline, i.e. no addition of arsenite. The corresponding times are 8, 13, and 15 h for the fresh, 1-day desiccated, and 7-day desiccated cells, respectively



sensors (Fig. 6b). The purified bacteriophytochrome is fluorescent in the NIR spectrum, with a peak excitation at 715 nm and a maximum emission at 725 nm (Fig. 6c). These spectroscopic features suggest that MagBphP is well suited for NIR detection and a good candidate for bacterial biosensors based on the reporter gene system.

Nephelometric measurements

In the following study, we compared the evolution of the light signal with turbidity in two light-emitting *E. coli* strains, a bioluminescent cell survival biosensor (*lux* operon expression driven by the constitutive P_{tpoD} promoter) and a near-infrared fluorescent cell survival biosensor (transcriptional fusion between the L-arabinose-inducible P_{BAD} promoter controlling the expression of both the apo-bacteriophytochrome *magBphP* and heme oxygenase *hmuO* genes). The turbidity of the bacterial LB growth medium was modified with the addition of a kaolin suspension to achieve a range of 0 to 500 nephelometric turbidity units (NTUs). Kaolin was preferred to formazine because we observed bacteriostatic effects of

formazine on the growth of *E. coli* (data not shown). Cellular growth (Fig. 7a) and the fluorescence and bioluminescence kinetics (Fig. 7b) were recorded on a micro-plate reader. During the experiment, the bacteria were inoculated in the kaolin-modified LB medium (with 1 % L-arabinose when required) and incubated at 30 °C under shaking. In these conditions, the microorganisms developed as shown in Fig. 7a by the increase of the OD_{600nm} without any significant effects of kaolin on the growth. We thus normalized the fluorescence and bioluminescence data with the OD_{600nm} values to correct the variations due to biomass increase (Fig. 7b). We then plotted the fluorescence or luminescence intensities against the turbidity at different times (Fig. 8), in the 60–120-min time range for the NIR fluorescence signal and 810–1020-min time range for the bioluminescence signal. In the latter case, the bioluminescence emission was significantly delayed. At each time point, data were normalized by the intensity measured at 0 NTU. For the bioluminescence signal, attenuation occurs very quickly when turbidity is increased (50 % loss at 75 NTU), while the signal remains optimal for NIR fluorescence in a large range of turbidity values (20 % loss at 500 NTU). We thus demonstrate

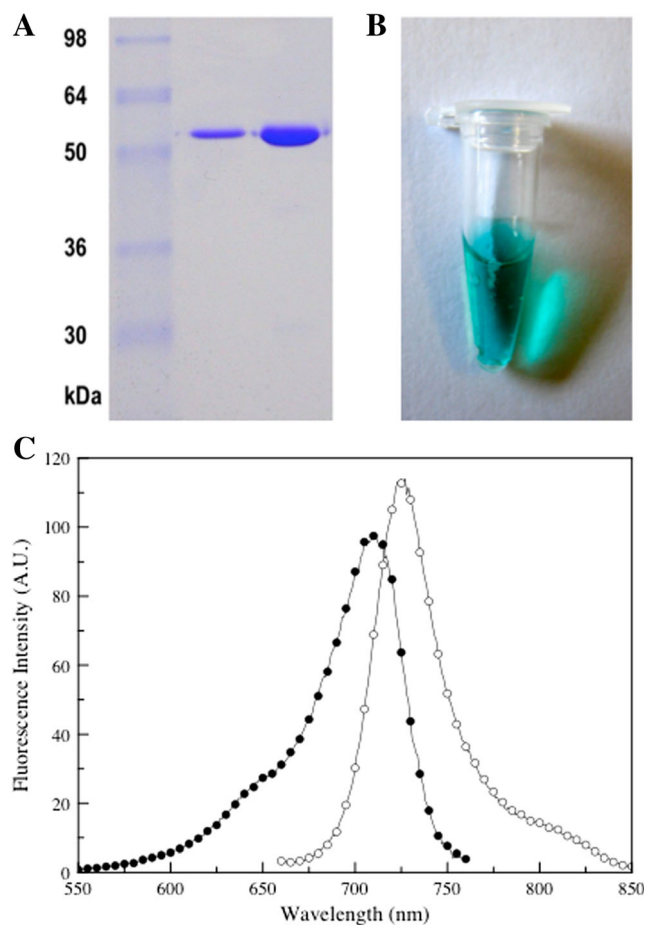


Fig. 6 MagBphP, a NIR fluorescent bacteriophytochrome from *Magnetospirillum magneticum* AMB-1. **a** SDS-PAGE analysis of purified MagBphP after affinity chromatography (1 and 5 μ g of protein deposits, from left to right). MW markers are indicated on the left. **b** Photography of the purified protein displaying a typical emerald-green color due to the biliverdin moiety. **c** MagBphP excitation spectrum with fluorescence emission measured at 770 nm (black circle) and emission spectrum with fluorescence excitation set at 650 nm (white circle)

here that the use of a bacteriophytochrome as a fluorescent reporter in the NIR spectrum represents a significant improvement to maintain the light signal fairly constant in a wide range of turbidity values.

Bacteriophytochrome-based arsenite biosensor

As evidenced above, NIR fluorescence bacterial biosensors reduce the adverse effects of turbidity in water samples. To go further in the proof-of-concept, we used as a blueprint the bioluminescent arsenite biosensor pArs-lux, replacing the bacterial luciferase by the NIR fluorescent bacteriophytochrome from *M. magneticum* AMB-1. We engineered a two-plasmid system where the arsenite-inducible P_{ars} promoter drives the expression of the heme oxygenase gene *hmuO* required for biliverdin production, while expression of the apo-bacteriophytochrome structural gene *magBphP* is controlled on a separate plasmid by the L-arabinose-inducible P_{BAD} promoter. We hypothesized that we can achieve a more sensitive measurement with the NIR fluorescent biosensor by separating the expression of the two genes. The rationale behind this assumption is that the production of a single molecule of HmuO is sufficient to produce several biliverdin molecules ready to be incorporated into a pool of constitutively expressed apo-bacteriophytochrome MagBphP. This approach was successfully led by Yagur-Kroll and Belkin on a *lux*-based biosensor with the luciferase structural genes *luxAB* separated from the substrate production-related genes *luxCDE* (Yagur-Kroll and Belkin 2010). Figure 9a summarizes the integration over time of the NIR fluorescence emission kinetics for different arsenite concentrations added to the culture together with 1 % L-arabinose. When L-arabinose is omitted (i.e., no apo-bacteriophytochrome gene expression), we record no fluorescence kinetics (data not shown). When arsenite

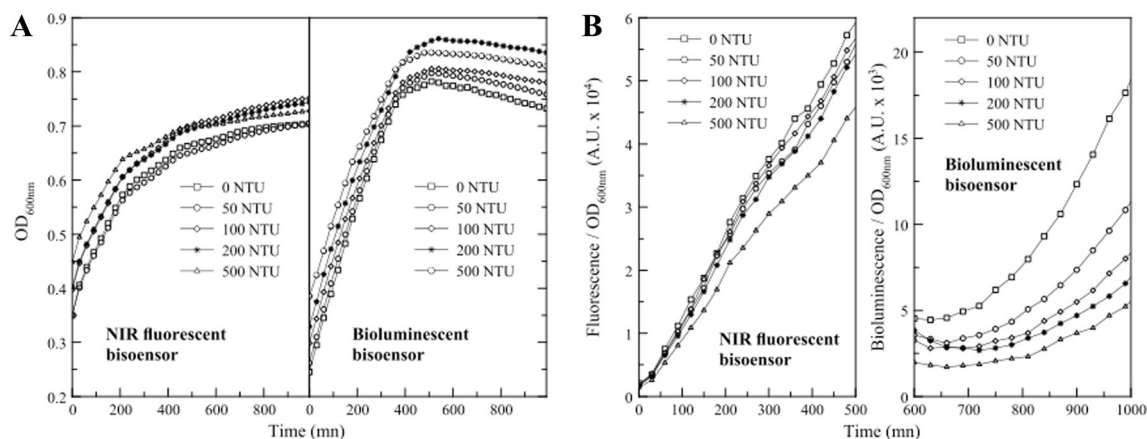


Fig. 7 Light emission and bacterial growth in turbid media. **a** OD kinetics measured at 600 nm for the NIR fluorescent (left panel) and the bioluminescent (right panel) biosensors at various turbidity values. **b** NIR fluorescence (left panel) and bioluminescence (right panel) emission

kinetics for various turbidity values, normalized by the OD_{600nm} . Fluorescence emission was measured at 755 nm with excitation set at 685 nm. Turbidity was varied in the 0–500 NTU range

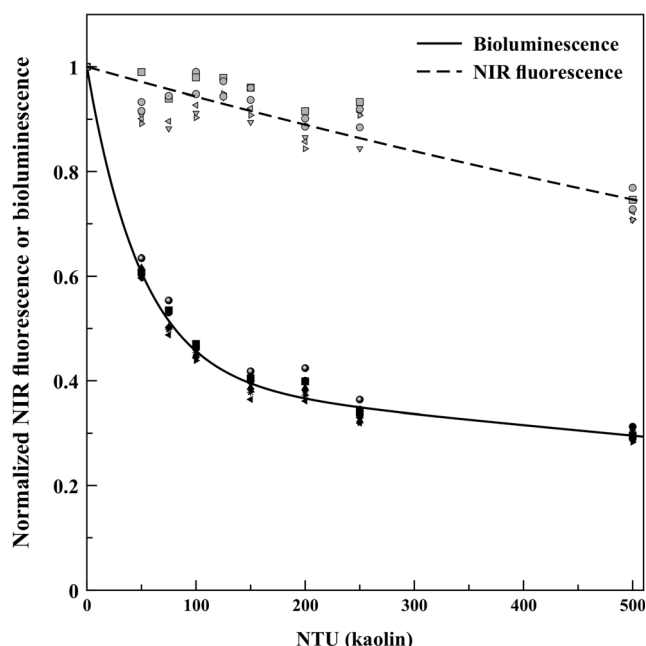


Fig. 8 Light signal attenuation in turbid media. Normalized bioluminescence intensities (*rpoD::luxCDABE*) and normalized NIR fluorescence intensities (*ara::magbph::hmuO*) are indicated with *black* and *gray symbols*, respectively

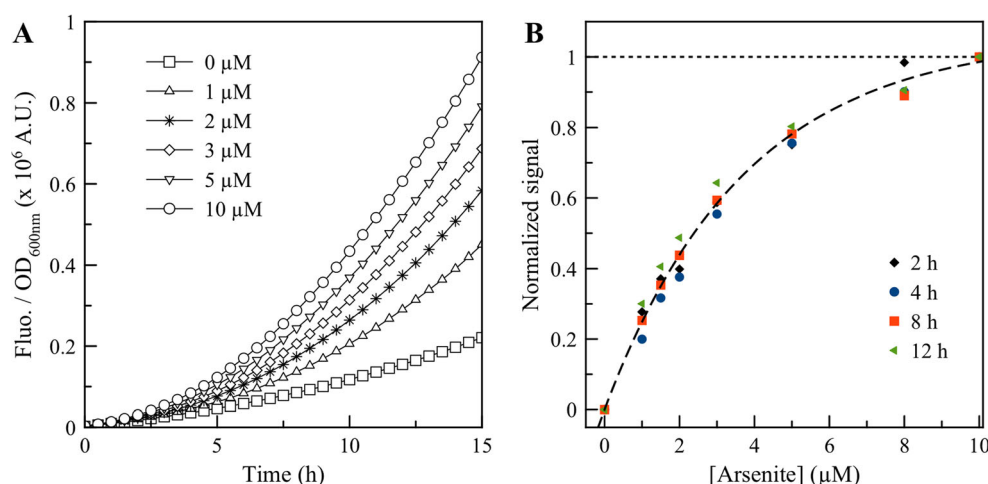
is added, we observe a dose-dependent increase of the NIR fluorescence signal (Fig. 9a). In the absence of arsenite (Fig. 9a, open squares), we still can measure a NIR fluorescence emission kinetics reminiscent of the bioluminescent signal observed with the *lux*-based arsenite biosensor; we believe this is probably due to the incomplete binding of the ArsR repressor to the P_{ars} promoter. The bacterial growth during the experiment is not significantly affected by the addition of either arsenite or L-arabinose (data not shown) and we chose to represent in Fig. 9a the integration over time of the (fluorescence/ OD_{600nm}) ratio to correct the fluorescence variations due to cellular biomass differences. We then used these

ratios to determine a calibration curve for the response of the biosensor to arsenite at different times after arsenite addition (Fig. 9b). In such conditions, we can estimate a detection limit of 0.5 μM for arsenite with the NIR-fluorescent biosensor, with a time response to arsenite that is reliable 2 h after contamination, i.e. about two times longer than with the bioluminescent pArs-*lux* biosensor hosted in *E. coli*. Owing to its performances in turbid media, the NIR fluorescent reporter gene system is an interesting alternative to other light-emitting biosensors. Further optimization is required to increase the performance of this arsenite biosensor in terms of sensitivity and time response, for instance by selecting a proper host strain and increasing the fluorescence quantum yield of the bacteriophytochrome by site-directed mutagenesis.

Conclusions

At the most basic level, the optimal functioning of bacterial biosensors based on the reporter gene system depends on the careful selection of an operator/regulator pair for efficient translation of the input signal (concentration of a molecule) into a detectable output such as bioluminescence or fluorescence. However, the specificity, the sensitivity, and the time-response of such biosensors do not exclusively rely on transcriptional regulation but are also strongly influenced by the host strain for the same bacterial species. Indeed, for two different *lux*-based biosensors, we observed in this study important variations of bioluminescence emission levels in five different *E. coli* strains used by the partners in the COMBITOX project. These observations were consistent in both fresh cultures and lyophilized cells. These discrepancies between *E. coli* strains may be linked either to variations of the promoter efficiency, the regulator stability or modification of the functional behaviors of the reporter proteins in the different yet closely related *E. coli* strains. Since our approach revealed

Fig. 9 A NIR fluorescent arsenite biosensor. **a** Integration over time of the fluorescence/ OD_{600nm} ratio kinetics after arsenite addition. **b** Standard curve for arsenite measured at different times. Data in Panel **a** are normalized by the value computed at $[As] = 10 \mu M$ after subtraction of the offset value at 0 μM arsenite. Fluorescence emission was measured at 755 nm with excitation set at 685 nm



that it is not possible to find an optimal and universal *E. coli* strain covering all the tested biosensors, biosensor design requires at early stages the empirical screening of host strains. The biosensor response can in a second step be improved by specific genome editing of the selected strain, such as deletion of export pumps or overexpression of specific metal uptake systems as illustrated elsewhere in this special issue (Cayron et al. 2015).

The use of environmental bacterial species as cellular chassis can also represent an interesting alternative to laboratory *E. coli* strains for optimal biodetection under field conditions with complex water samples. In such a context, we transferred our genetic constructs in *D. deserti*, an environmental strain highly resistant to desiccation; such ability can be exploited for the development of a simpler conservation process than lyophilization. We successfully obtained two functional biosensors hosted in *D. deserti*: (i) the constitutive P_{gapdh} promoter was efficiently used to induce *lux* operon expression, resulting in a cell-survival biosensor with a usage for global toxicity similar to the one developed in *E. coli* and described in this special issue (Prévéral et al. 2016); (ii) using P_{ars} promoter from *E. coli* in *D. deserti*, we were able to detect a concentration as low as 100 nM of arsenite in the medium. The time response of these biosensors is slow (around 10 h) and optimizations are required in the future. We showed that air-dried *D. deserti* cells remain responsive to arsenite even after 7 days of storage at room temperature in conditions where *E. coli* fails to recover. To the best of our knowledge, this study presents the first evidence of a strain from the *Deinococcus* genus able to produce both the bacterial luciferase and its substrate and to generate bioluminescence. Such genetic modifications result in whole-cell biosensors with potential resistance to DNA-damaging radiation, bestowing robustness to the whole-cell biosensor module of the COMBITOX-like water analyzer exposed to chemical, biological, radiological, or nuclear (CBRN) threats. Owing to its resistance to desiccation, we evidenced that *D. deserti* is a promising candidate to host metal biosensors that can be simply dried for storage while retaining its sensitivity toward the pollutant. The proof-of-concept brought by the prototype arsenite bioluminescent sensor described here paves the way to the development of robust biosensors with improved sensitivity, time response, and shelf life.

In a reporter-based biosensor, the selection of the transducing biological system represents an additional and critical optimization step. Depending on the origin of the water samples, the signal emitted by the reporter gene product can be detected and quantified with varying efficiency; for bioluminescent or fluorescent biosensors, light emission in the visible range of the spectrum is prone to re-absorption in colored water samples or diffusion in turbid media. Fluorescent organic molecules present in environmental samples are also liable to interact with the light emitted by the biosensors, resulting in

reduced sensitivity or specificity. The COMBITOX project focuses on the construction of a semi-autonomous inline water analyzer encompassing various biosensors; it is destined to monitor real water networks and the actual physicochemical properties of the water samples prompted us to improve the transducing element of our biosensors. Thanks to its fluorescence in the near infrared spectrum, we tested the bacteriophytochrome from *M. magneticum* AMB-1 (MagBphP) as an alternative to bacterial luciferase to perform biodetection in turbid samples. We compared the responses of different cell biosensors based on the continuous expression of either the *lux* operon (promoter P_{IpoD}) or *magBphP* and its companion heme oxygenase gene (L-arabinose-inducible P_{BAD} promoter) and evidenced a drastic loss of the luminescent signal when turbidity is increased, while for bacteriophytochrome-based biosensors, the NIR fluorescent signal remains optimal in a large range of turbidity values. We then designed an arsenite biosensor relying on NIR fluorescence emission and evidenced a detection limit of 0.2 μM of arsenite, with a very good reproducibility. Although the time response should be optimized in the future, our NIR-fluorescent reporter gene system represents an interesting alternative to other light-emitting biosensors, notably to transfer laboratory-optimized biosensors into on-field measurements.

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