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A cofactor consumption screen identifies promising NfsB family nitroreductases for dinitrotoluene remediation --Manuscript Draft--

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Abstract:	Objectives		
	To survey a library of over-expressed nitroreductases to identify those most active with 2,4- and 2,6-dinitrotoluene substrates, as promising candidates for phytoremediation of soils and groundwater contaminated with poly-nitro toluene pollutants. Results To indirectly monitor dinitrotoluene reduction we implemented a nitroblue tetrazolium dye screen to compare relative rates of NADPH consumption for 58 nitroreductase candidates, over-expressed in a nitroreductase-deleted strain of Escherichia coli. Although the screen only provides activity data at a single substrate concentration, by		

	altering the substrate concentration and duration of incubation we showed we could first distinguish between more-active and less-active enzymes and then discriminate between the relative rates of reduction exhibited by the most active nitroreductases in the collection. We observed that members of the NfsA and NfsB nitroreductase families were the most active with 2,4-dinitrotoluene, but that only members of the NfsB family were proficient with 2,6-dinitrotoluene. Two NfsB family members, YfkO from Bacillus subtilis and NfsB from Vibrio vulnificus, appeared especially effective with these substrates. Purification of both enzymes as His6-tagged recombinant proteins enabled in vitro determination of Michaelis-Menten kinetic parameters with each dinitrotoluene substrate. Conclusions V. vulnificus NfsB is a particularly promising candidate for bioremediation applications, being ca. 5-fold more catalytically efficient with 2,4-dinitrotoluene and over 26-fold more active with 2,6-dinitrotoluene than the benchmark E. coli nitroreductases NfsA and NfsB.
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A cofactor consumption screen identifies promising NfsB family nitroreductases for dinitrotoluene remediation

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Abstract

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- 2 Objectives: To survey a library of over-expressed nitroreductases to identify those most
- active with 2,4- and 2,6-dinitrotoluene substrates, as promising candidates for
- 4 phytoremediation of soils and groundwater contaminated with poly-nitro toluene
- 5 pollutants.
- 6 Results: To indirectly monitor dinitrotoluene reduction we implemented a nitroblue
- 7 tetrazolium dye screen to compare relative rates of NADPH consumption for 58
- 8 nitroreductase candidates, over-expressed in a nitroreductase-deleted strain of Escherichia
- 9 coli. Although the screen only provides activity data at a single substrate concentration, by
- altering the substrate concentration and duration of incubation we showed we could first
- distinguish between more-active and less-active enzymes and then discriminate between
- the relative rates of reduction exhibited by the most active nitroreductases in the collection.
- 13 We observed that members of the NfsA and NfsB nitroreductase families were the most
- active with 2,4-dinitrotoluene, but that only members of the NfsB family were proficient
- with 2,6-dinitrotoluene. Two NfsB family members, YfkO from *Bacillus subtilis* and NfsB from
- 16 Vibrio vulnificus, appeared especially effective with these substrates. Purification of both
- 17 enzymes as His₆-tagged recombinant proteins enabled in vitro determination of Michaelis-
- 18 Menten kinetic parameters with each dinitrotoluene substrate.
- 19 Conclusions: V. vulnificus NfsB is a particularly promising candidate for bioremediation
- applications, being ca. 5-fold more catalytically efficient with 2,4-dinitrotoluene and over
- 21 26-fold more active with 2,6-dinitrotoluene than the benchmark *E. coli* nitroreductases NfsA
- and NfsB.
- 23 Keywords: Bioremediation; Dinitrotoluene; NADPH depletion assay; NfsA; NfsB;
- 24 Nitroreductase; YfkO.

Introduction

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2,4-dinitrotoluene and 2,6- dinitrotoluene, widely used as propellants or as precursors in the manufacture of 2,4,6-trinitrotoluene (TNT), are toxic and persistent contaminants of soil and groundwater (Dontsova et al, 2009). The strongly electron-withdrawing nitro groups on poly-nitro toluenes delocalise the electronic charge distribution in the aromatic rings, rendering them resistant to oxidative degradation by other enzymes (Roldan et al, 2008). Bacterial type I nitroreductases, which reduce nitro substituents via concerted two-electron transfer steps, yield derivatives that are far more amenable substrates for ring degrading enzymes (Williams et al, 2015; Roldan et al, 2008). Plants engineered for high-level expression of bacterial nitroreductases have shown enhanced transformation and tolerance of poly-nitro toluenes, offering prospects for phytoremediation of contaminated environments (Zhang et al, 2019; Van Dillewijn et al, 2008; Hannink et al, 2007). However, despite there being a great diversity of bacterial nitroreductases available in nature (Akiva et al, 2017), very few candidates have been surveyed to assess their relative potential for these applications. A dearth of characterised nitroreductase candidates is particularly evident in the context of dinitrotoluene remediation. Effective surveying of novel enzyme candidates requires an efficient activity screen. We previously described how Escherichia coli cells over-expressing nitroreductase enzymes that are active with dinitrotoluene substrates became more sensitive to these toxins, and that IC₅₀ data were somewhat predictive of relative levels of catalytic efficiency (k_{cat}/K_M) for the purified enzymes (Rich et al, 2018). However, as IC₅₀ calculations require measurements to be made across a broad range of substrate concentrations, this is not an approach

amenable to large-scale screening of candidate gene libraries. Here we describe our implementation of an alternative assay, based on cofactor consumption, to rapidly identify nitroreductase enzymes proficient in reduction of both 2,4- and 2,6-dinitrotoluene.

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Materials and Methods

Chemicals, plasmids, bacterial strains and nitroreductases 53 All reagents were purchased from Sigma-Aldrich (Saint Louis, MO). For protein purification, 54 55 the genes encoding YfkO from Bacillus subtilis and NfsB from Vibrio vulnificus were expressed from the His6-tag expression vector pET28a+ in E. coli strain BL21(DE3) (Novagen, 56 Merck Millipore; Billerica, MA) as previously described (Prosser et al, 2013). For library 57 screening and IC₅₀ assays, nitroreductase candidates were over-expressed from plasmid 58 pUCX in E. coli 7NT, a gene-deleted derivative of strain W3110 that lacks seven endogenous 59 60 oxidoreductase genes (nfsA, nfsB, azoR, nemA, yieF, ycaK and mdaB) to minimise 61 background nitroreductase activity and an efflux pump gene (tolC) to maximise substrate retention (Copp et al, 2014). The 58-membered nitroreductase gene library was constructed 62 as previously described (Mowday et al, 2016) and the full list of candidate genes (with 63 UniProt default identifiers) is summarised in Table 1. 64

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Nitroblue tetrazolium assay

<Insert Table 1 here>

Cells of *E. coli* 7NT pUCX:nitroreductase strains were used to inoculate wells of a sterile 96-well microplate containing 200 μ L lysogeny broth (LB) supplemented with ampicillin (100 μ g.mL⁻¹) and glucose (0.2% w/v). Cultures were incubated at 30 °C, 200 rpm for 16 hours,

after which 10 µL of each overnight culture was used to inoculate wells of a sterile 96-well microplate containing 190 μL induction medium (LB medium supplemented with ampicillin (100 µg.mL⁻¹), glucose (0.2% w/v) and IPTG (0.05 mM). Induced cultures were incubated at 30 °C, 200 rpm for 6 hours. Crude cell lysates were prepared by incubating 100 μL of each culture with BugBuster® Protein Extraction Reagent at a 1:1 (v/v) ratio at room temperature for 30 min. To monitor cell lysate NADPH consumption, duplicate 200 µL reactions were prepared consisting of 100 mM KPO₄ buffer pH 8.0, crude cell lysate (40 µL for full library screening assays or 5 μL for NfsB-family screening assays) and 150 μM substrate (2,4dinitrotoluene or 2,6-dinitrotoluene), and reactions were initiated by addition of 50 µL of 800 µM NADPH. Reactions were incubated at room temperature for 30 min for full library screening assays, or 10 min for NfsB-family screening assays. Reactions were halted by the addition of 50 µL nitroblue tetrazolium solution (nitroblue tetrazolium (2 mg.mL⁻¹), phenazine methosulfate (0.3 mg.mL⁻¹, prepared fresh prior to use) and 100 mM KPO₄ buffer (pH 8.0)). Levels of formazan production, corresponding to the amount of NADPH remaining in each well, were quantified 3-5 min post addition of the nitroblue tetrazolium solution by measuring absorbance at 590 nm using an EnSpireTM 2300 Multilabel Reader (PerkinElmer, Waltham, MA). In pilot tests, four E. coli 7NT pUCX:nitroreductase strains (expressing the genes encoding NfrA_Bli, NfsA_Vv, Ydgl_Bs and YcnD_Bs) exhibited a substantial decrease in NADPH levels in the absence of substrate. Reasoning that this high background NADPH oxidase activity would confound measurement of substrate consumption, these strains were excluded from the full library analysis.

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IC₅₀ assays

Cells of E. coli 7NT pUCX:nitroreductase strains were used to inoculate wells of sterile 96well microplates containing 200 μL LB medium supplemented with ampicillin (100 μg.mL⁻¹) and glucose (0.2% w/v). Cultures were incubated at 30 °C, 200 rpm for 16 hours, after which they were diluted 20-fold into 15 mL centrifuge tubes containing 2 mL induction medium (LB medium supplemented with ampicillin (100 μg.mL⁻¹), glucose (0.2% w/v) and IPTG (0.05 mM). Induced cultures were incubated at 30 °C, 200 rpm for 2.5 hours. Aliquots (30 μL) of each culture were added to wells of a sterile 384-well microplate containing 30 μL induction medium ± two-fold the desired final dinitrotoluene concentration. In total, each strain was challenged in duplicate across 7 to 15 conditions containing 1.5-fold increasing titrations of 2,4-dinitrotoluene or 2,6-dinitrotoluene together with a medium-only control. The absorbance of each well at 600 nm was measured using an EnSpire[™] 2300 Multilabel Reader (PerkinElmer, Waltham, MA). Cultures were incubated at 30 °C, 200 rpm for a further 4 hours, after which absorbance readings at 600 nm were recorded once more. The increase in absorbances at 600 nm of corresponding challenged and unchallenged wells for each strain were compared and used to calculate percentage growth following challenge. IC₅₀ values (the concentration of 2,4-dinitrotoluene or 2,6-dinitrotoluene estimated to reduce growth by 50% relative to the unchallenged control) were calculated using a doseresponse inhibition four-parameter variable slope equation in GraphPad Prism 7.0 (GraphPad Software Inc., La Jolla, CA).

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His₆-tagged protein purification and Michaelis-Menten kinetics

Recombinant His₆-tagged nitroreductases were purified by nickel-affinity chromatography using Ni/NTA resin (Novagen, Merck, Darmstadt, Germany), followed by incubation with a molar excess of FMN cofactor and buffer exchange as previously described (Prosser et al,

2010). Apparent steady state enzyme kinetics were determined via monitoring decrease in absorbance at 340 nm (NADPH extinction co-efficient, 6,220 M⁻¹ cm⁻¹) at a fixed concentration of NADPH and varying concentrations of 2,4-dinitrotoluene or 2,6-dinitrotoluene. Reactions were carried out in 60 μL volumes in UVettes™ (Eppendorf, Hamburg, Germany), using a 1 cm path length, and the wavelength was measured using a Helios γ UV-Vis spectrophotometer (Thermo Fisher Scientific, Waltham, MA). Reactions consisted of 10 mM Tris-Cl pH 7.0 buffer, 250 μM NADPH, 2% DMSO and substrate concentrations ranging from 0 - 400 μM 2,4-dinitrotoluene and 0 -1600 μM 2,6-dinitrotoluene. Reactions were initiated by addition of an appropriate dilution of enzyme and the linear decrease in absorbance monitored for the first 20-40 s post enzyme addition. Non-linear regression analyses and Michaelis-Menten curve fitting was performed using GraphPad Prism v 7.0 (GraphPad Software, San Diego, California, USA).

Results and Discussion

To identify enzymes capable of activating diverse nitroaromatic prodrugs we previously generated a 58-membered library of nitroreductase candidates in *Escherichia coli* reporter strains that express either a β -galactosidase or GFP gene under control of a DNA damage responsive (SOS) promoter (Mowday et al, 2016; Copp et al, 2017). We tested whether dinitrotoluene reduction could likewise be monitored via SOS assays, reasoning that reduction of TNT is known to yield genotoxic metabolites (Bolt et al, 2006). However, we found that both 2,4-dinitrotoluene and 2,6 dinitrotoluene evoked high levels of background signal in nitroreductase-expressing reporter strains that precluded accurate screening (not shown). We therefore sought to implement an alternative screening procedure based on

consumption of the NAD(P)H cofactor that acts as an electron source for bacterial Type I nitroreductases. For this, we adapted a colorimetric nitroblue tetrazolium assay, previously described by Mayer and Arnold (2002) for directed evolution of 6-phosphogluconate dehydrogenase enzymes.

In the presence of NADPH yellow nitroblue tetrazolium is reduced to a purple formazan dye that can be quantified at 590 nm. This allowed us to measure the relative levels of dinitrotoluene reduction in lysates derived from *E. coli* strains over-expressing nitroreductase candidates. To minimise background nitroreductase activity, screening of the full 58-membered nitroreductase candidate library was performed in a strain of *E. coli* lacking endogenous *nfsA*, *nfsB*, *azoR*, *nemA*, *yieF*, *ycaK* and *mdaB* genes (strain 7NT; Copp et al, 2014). Lysates were incubated with NADPH and either 2,4-dinitrotoluene or 2,6-dinitrotoluene, then reactions were stopped with nitroblue tetrazolium chloride and the amount of formazan in challenged lysates compared to unchallenged duplicates.

The full library screen confirmed that, of the enzymes tested, NfsA and NfsB family nitroreductases consistently consumed the most NADPH in the presence of dinitrotoluene substrates (Figure 1A). Notably, although members of both the NfsA and NfsB families were represented among the top 25 most active enzymes with 2,4-dinitrotoluene as substrate, the NfsB family enzymes markedly outperformed those from the NfsA family with 2,6-dinitrotoluene as substrate (Figure 1A). To distinguish the relative activities of the NfsB family enzymes it was necessary to reduce the amount of culture lysate from 40 μ L for the full library screening assay to 5 μ L and the incubation time from 30 min to 10 min (Figure 1B).

<Insert Figure 1 here>

We next sought to test whether results from the nitroblue tetrazolium assay were generally congruent with IC₅₀ data by performing detailed IC₅₀ assays for all strains expressing nfsB family genes (Table 2) and correlating the measured IC₅₀ values (μ M) with NADPH consumption (percent Δ A₅₉₀) (Figure 2). The inverse correlation between IC₅₀ and NADPH consumption for 2,4-dinitrotoluene was moderate rather than strong (r^2 = 0.35); this is likely a consequence of the nitroblue tetrazolium assay providing a 'snapshot' of activity at a single substrate concentration, whereas IC₅₀ data derive from a more comprehensive midpoint of activity across a titration range. However, an additional advantage of the rapid nitroblue tetrazolium assay over performing more extensive IC₅₀ measurements was evident in its ability to generate data for 2,6-dinitrotoluene, whereas IC₅₀ values could not be calculated for the majority of nfsB-expressing strains owing to insufficient toxicity at the achievable solubility limit (Table 2).

<Insert Table 2 here> <Insert Figure 2 here>

Based on NADPH consumption data, the NfsB family enzyme *Bacillus subtilis* YfkO (YfkO_Bs) appeared most active with 2,4-dinitrotoluene, whereas *Vibrio vulnificus* NfsB (NfsB_Vv) appeared most active with 2,6-dinitrotoluene (Figure 1B). We purified each enzyme as a His₆-tagged recombinant protein by nickel affinity chromatography and measured their Michaelis-Menten kinetic parameters with either 2,4-dinitrotoluene or 2,6-dinitrotoluene as the reductive substrate (Table 3). In terms of catalytic efficiency, the k_{cat}/K_M values for NfsB_Vv and YfkO_Bs with 2,4-dinitrotoluene were 4- to 7-fold higher than those previously reported for either of the canonical Type I nitroreductases NfsA_Ec and NfsB_Ec (Rich et al, 2018). Their superiority with 2,6-dinitrotoluene was even more profound, with the measured k_{cat}/K_M values for NfsB_Vv and YfkO_Bs being 20- to 37-fold higher than those previously reported for NfsA_Ec and NfsB_Ec (Rich et al, 2018). NfsB_Vv was particularly

effective with each substrate, and appears a promising candidate for expression in transgenic dinitrotoluene remediation systems.

<Insert Table 3 here>

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Conclusions

We have previously used activation of the E. coli SOS response to report on genotoxicity as a measure of nitroreductase-mediated activation of prodrug substrates. Here, to monitor activity with dinitrotoluene substrates that did not yield a clear SOS signal upon reduction, we implemented a more generally applicable screen for nitroreductase activity based on consumption of exogenously added NADPH that provided a dominant electron source for the reaction. In addition to providing much higher throughput than IC₅₀ assays, we showed that our screen could provide relative activity data with substrates such as 2,6dinitrotoluene that are insufficiently toxic at achievable concentrations for IC50 values to be calculated. The linear range of the assay was insufficient to discriminate between the more active NfsB family nitroreductases while still detecting the less active NfsA and AzoR family enzymes. However, by reducing both the substrate concentration and duration of incubation we were able to discriminate between the different NfsB family nitroreductases and identified V. vulnificus NfsB and B. subtilis YfkO as two of the most active dinitrotoluene reducing enzymes in our collection. Purification of these nitroreductases as His6-tagged proteins enabled steady state kinetic assays to be performed, demonstrating V. vulnificus NfsB to be over 5-fold more active with 2,4-dinitrotoluene and over 26-fold more active with 2,6-dinitrotoluene than either of the canonical nitroreductases *E. coli* NfsA and NfsB.

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References

217 218 Akiva E, Copp JN, Tokuriki N, Babbitt PC (2017) Evolutionary and molecular foundations of 219 multiple contemporary functions of the nitroreductase superfamily. Proc Natl Acad Sci U S A 114:E9549-E9558. 220 Bolt HM, Degen GH, Dorn SB, Plottner S, Harth V (2006) Genotoxicity and potential 221 222 carcinogenicity of 2,4,6-trinitrotoluene: structural and toxicological considerations. Rev Environ Heatl 21:217–228. 223 224 Copp JN, Williams EM, Rich MH, Patterson AV, Smaill JB, Ackerley DF (2014) Toward a high-225 throughput screening platform for directed evolution of enzymes that activate genotoxic 226 prodrugs. Protein Eng Des Sel 27:399-403. 227 Copp JN et al. (2017) Engineering a multifunctional nitroreductase for improved activation of prodrugs and PET probes for cancer gene therapy. Cell Chem Biol 24:391-403 228 229 Dontsova KM, Pennington JC, Hayes C, Simunek J, Williford CW (2009) Dissolution and 230 transport of 2,4-DNT and 2,6-DNT from M1 propellant in soil. Chemosphere 77:597–603. 231 Hannink NK, Subramanian M, Rosser SJ, Basran A, Murray JA, Shanks JV, Bruce NC (2007) Enhanced transformation of TNT by tobacco plants expressing a bacterial nitroreductase. Int 232 J Phytoremediation 9:385-401. 233 234 Mayer KM, Arnold FH (2002) A colorimetric assay to quantify dehydrogenase activity in 235 crude cell lysates. J Biomol Screen 7:135-140.

Mowday AM et al. (2016) Rational design of an AKR1C3-resistant analog of PR-104 for 236 237 enzyme-prodrug therapy. Biochem Pharmacol 116:176-187. Prosser GA et al. (2010) Discovery and evaluation of Escherichia coli nitroreductases that 238 activate the anti-cancer prodrug CB1954. Biochem Pharmacol 79:678-687. 239 240 Prosser GA et al. (2013) Creation and screening of a multi-family bacterial oxidoreductase library to discover novel nitroreductases that efficiently activate the bioreductive prodrugs 241 CB1954 and PR-104A. Biochem Pharmacol 85:1091-1103. 242 243 Roldan MD, Perez-Reinado E, Castillo F, Moreno-Vivian C (2008) Reduction of polynitroaromatic compounds: the bacterial nitroreductases. FEMS Microbiol Rev 32:474-244 500. 245 246 Rich MH, Sharrock AV, Hall KR, Ackerley DF, MacKichan JK (2018) Evaluation of NfsA-like nitroreductases from Neisseria meningitidis and Bartonella henselae for enzyme-prodrug 247 248 therapy, targeted cellular ablation, and dinitrotoluene bioremediation. Biotechnol Lett 249 40:359-367. 250 Van Dillewijn P, Couselo JL, Corredoira E, Delgado A, Wittich RM, Ballester A, Ramos JL 251 (2008) Bioremediation of 2,4,6-trinitrotoluene by bacterial nitroreductase expressing transgenic aspen. Environ Sci Technol 42:7405–7410. 252 253 Williams EM et al. (2015) Nitroreductase gene-directed enzyme prodrug therapy: insights and advances toward clinical utility. Biochem J 471:131-153. 254 255 Zhang L, Rylott EL, Bruce NC, Strand SE (2019) Genetic modification of western wheatgrass (Pascopyrum smithii) for the phytoremediation of RDX and TNT. Planta 249:1007-1015. 256

Table 1. Nitroreductase candidate library

Bacterial strain nitroreductase sourced from	Nitroreductase gene	UniProt identifier
Bacillus amyloliquefaciens CAMR 0454	ywrO	Q8VSR5
Bacillus coagulans ATCC 7050	nfsA	G2TLJ0
Bacillus licheniformis ATCC14580	nfrA	Q65DM9
	ycnD	Q65ND8
	yfkO	Q65MG6
Bacillus subtilis NZ isolate	nfrA	P39605
	ycnD	P94424
	ydgI	P96707
	yfkO	O34475
	ywrO	P80871
Bacillus thuringiensis serovar konkukian str. 97-27	nfsA	Q6HKT9
Citrobacter koseri ATCC 27156	nfsA	A8AIR6
	nfsB	A8AJP2
Cronobacter (previously Enterobacter) sakazakii	nfsA	A7MF39
ATCC 29544	nfsB	A7MK61
Erwinia carotovora subsp. atroseptica SCRI1043	nfsA	Q6D3R4
Escherichia coli W3110	azoR	P41407
	mdaB	POAEY5
	nemA	P77258
	nfsA	P17117
	nfsB	P38489
	wrbA	A1A9Q9
	усаК	P43340
	ycdl	P75894
	ydjA	P0ACY1
	yieF	P0AGE6
Klebsiella pneumoniae ATCC 13883	nemA	A6T9Z9
	nfsA	A6TT6U7
	nfsB	A6T5Y2
	ycdI	A6T798
Lactobacillus sakei subsp. sakei 23K	nfsA	Q38UW5
Listeria innocua Clip11262	nfsA	Q92D83
	ywrO	Q92AM4
Listeria welshimeri ATCC 35897	nfsA	A0AH51
Mycobacterium smegmatis mc²155	nfsA	A0QWF2
Neisseria meningitidis strain H44/76	nfsB	E6N0E6
Nostoc punctiforme PCC73102	nfsA	B2J8C5
Pseudomonas aeruginosa PAO1	усаК	Q9I4B3
	ydgI	Q9HTZ9
	yieF	O31038
Pseudomonas putida KT2440	azoR	Q88EC8
	nfsA	Q88K03

	nfsB	Q88K59
Pseudomonas savastanoi pv. phaseolicola	mdaB	Q48NL7
	wrbA	E7PSG4
Salmonella typhi ATCC 19430	azoR	P63462
	nemA	Q8Z6P3
	nfsA	Q8Z848
	nfsB	Q8Z8M5
Vibrio fischeri ATCC 7744	frasel	P46072
	nfsA	B5EVB7
Vibrio harveyi ATCC 14126	frp	Q56691
	nfsB	A6AP86
Vibrio harveyi KCTC 2720	frp, herein referred to as frp2	AAM73721 (Genbank)
Vibrio vulnificus ATCC 27562	azoR	Q8DA68
	nfsA	Q8D4B5
	nfsB	Q7MCD2
	nemA	Q8D549

Table 2. Dinitrotoluene IC_{50} values for *E. coli* 7NT pUCX: *nitroreductase* strains

	IC ₅₀ (μM)		
Nitroreductase	2,6-dinitrotoluene	2,4-dinitrotoluene	
YfkO_Bs	670 ± 60	330 ± 60	
NfsB_Nme	680 ± 70	310 ± 60	
NfsB_Vv	800 ± 70	510 ± 110	
NfsB_Cs	940 ± 130	530 ± 60	
YfkO_Bli	>1000	470 ± 40	
NfsB_Pp	>1000	700 ± 140	
NfsB_Kp	>1000	740 ± 20	
Frasel_Vf	>1000	750 ± 50	
NfsB_Vh	>1000	770 ± 60	
NfsB_St	>1000	790 ± 20	
NfsB_Ck	>1000	800 ± 60	
NfsB_Ec	>1000	840 ± 60	

Table 3. Michaelis-Menten kinetic parameters for reduction of 2,4-dinitrotoluene and 2,6-dinitrotoluene by purified His₆-tagged nitroreductases

Nitroreductase	Substrate	$K_M^1(\mu M)$	<i>k_{cat}</i> ¹ (s ⁻¹)	$k_{cat}/K_{M} (M^{-1} s^{-1})$
B. subtilis YfkO	2,4-dinitrotoluene	62 ± 7	140 ± 42	440,000 ± 140,000
	2,6-dinitrotoluene	510 ± 80	22 ± 2	43,000 ± 7,000
V. vulnificus NfsB	2,4-dinitrotoluene	320 ± 50	200 ± 20	640,000 ± 110,000
	2,6-dinitrotoluene	1600 ± 110	90 ± 4	57,000 ± 5,000

 $^{^{1}\}text{Apparent kinetic parameters}$ as measured at 250 μM NADPH

Figure Legends

Figure 1. NADPH consumption by E. coli 7NT pUCX:nitroreductase cell lysates in the presence of 2,4- or 2,6-dinitrotoluene compounds. A. Crude cell lysates (40 µl) of nitroreductase-expressing E. coli 7NT strains were incubated with 200 μM NADPH and 150 μM of either 2,4-dinitrotoluene (2,4-DNT) or 2,6-dinitrotoluene (2,6-DNT) for 30 min. Addition of nitroblue tetrazolium post-incubation yielded formazan dye in proportion to the remaining NADPH, which was quantified by measuring absorbance at 490 nm. Plotted values indicate the percentage of NADPH consumed by dinitrotoluene-challenged lysates during the reaction compared to unchallenged duplicate controls. Individual E. coli 7NT pUCX: nitroreductase strains are labelled according to the name of the nitroreductase (NTR) followed by a two- or three-letter abbreviation that defines the genus and species of the bacteria that the enzyme was derived from (a full list of all NTRs tested in this study and their accession codes is provided in the Materials and Methods). Data are the average of three independent assays ± SD. B. To delineate the activities of NfsB-family NTRs, the assay was repeated as in panel A, only using a smaller volume of crude cell lysate (5 μl) and shorter incubation time (10 min).

Figure 2. Scatter diagram and correlation analysis of NfsB-family nitroreductase activities with 2,4-dinitrotoluene in nitroblue tetrazolium and IC₅₀ assays. Correlation analysis showed that 2,4-dinitrotoluene (2,4-DNT) IC₅₀ values for *E. coli* 7NT pUCX:nitroreductase strains were negatively correlated (p < 0.05, Pearson's correlation test) with the percentage of NADPH consumed by the corresponding strain lysates in nitroblue tetrazolium assays (r^2 =0.35).

Figure 1

₽

100 **-**

80 90

NfsA-family

NfsB-family

2,6 DNT 2,4 DNT

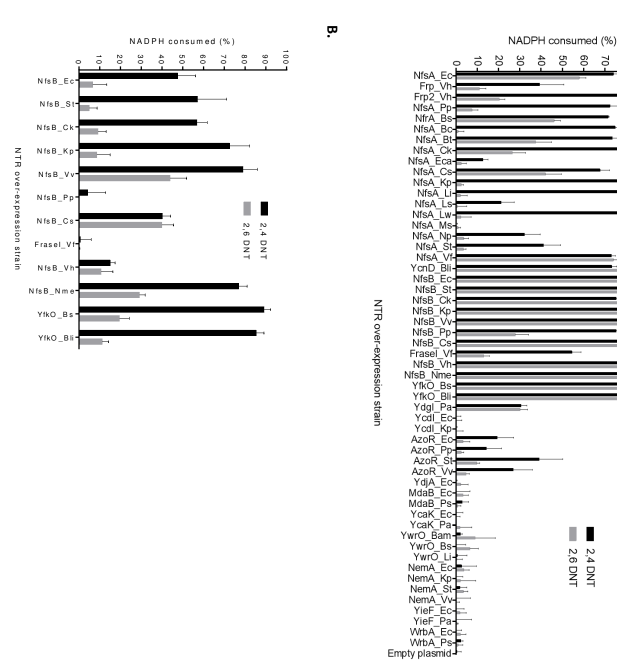
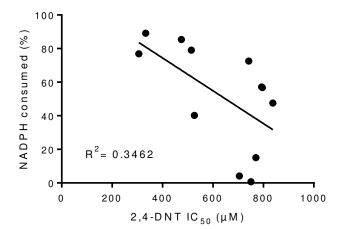


Figure 2



NIA

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