A glutaredoxin domain fused to the radical-generating subunit of ribonucleotide reductase (RNR) functions as an efficient RNR reductant

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Class I ribonucleotide reductase (RNR) consists of a catalytic subunit (NrdA) and a radical-generating subunit (NrdB) that together catalyze reduction of ribonucleotides to their corresponding deoxyribonucleotides. NrdB from the firmicute Flack-lamia ignava is a unique fusion protein with N-terminal additions of a glutaredoxin (Grx) domain followed by an ATP-binding domain, the ATP cone. Grx, usually encoded separately from the RNR operon, is a known RNR reductant. We show that the fused Grx domain functions as an efficient reductant of the F. ignava class I RNR via the common dithiol mechanism and, interestingly, also via a monothiol mechanism, although less efficiently. To our knowledge, a Grx that uses both of these two reaction mechanisms has not previously been observed with a native subunit of ribonucleotide reductase (RNR) functions as an on/off switch promoting ribonucleotide reduction in the presence of ATP and inhibiting RNR activity in the presence of dATP. We found that dATP bound to the ATP cone of F. ignava NrdB promotes formation of tetramers that cannot form active complexes with NrdA. The ATP cone bound two dATP molecules but only one ATP molecule. F. ignava NrdB contains the recently identified radical-generating cofactor MnIII/MnIV. We show that NrdA from F. ignava can form a catalytically competent RNR with the MnIII/MnIV-containing NrdB from the flavobacterium Leewenhoekiella blandensis. In conclusion, F. ignava NrdB is fused with a Grx functioning as an RNR reductant and an ATP cone serving as an on/off switch.

Ribonucleotide reductase (RNR)2 is an essential enzyme that catalyzes the synthesis of the DNA building blocks (dNTPs) by reduction of the four ribonucleotides. RNR plays a key role in DNA synthesis and DNA repair and consequently attracts biomedical interest as a potential target for antibacterial substances and for anticancer therapies. Currently, the RNR enzyme family comprises three different RNR classes and several subclasses. The three classes have a common reaction mechanism that builds on radical chemistry but differ in the way they initiate the radical mechanism (1–5). The class I RNRs consist of a larger catalytic component (NrdA) and a smaller radical-generating metal-containing component (NrdB) in which the dinuclear metal site differs between subclasses. Currently, class I RNRs have been subclassified based on radical cofactor type (subclasses 1a, Ib, Ic, Id, and Ie) or evolutionary history (subclasses NrdA/B followed by a small letter plus subclass NrdE/F) (1, 6). Metal content does not always follow phylogeny because two unrelated Mn2 subclasses exist, where one subclass contains a tyrosyl radical in the vicinity of a MnII/MnIII center (Ib, NrdE/F), and another recently identified subclass (Id, NrdA/B) contains a mixed valent MnII/MnIV metal center that harbors the unpaired electron (7–9). In eukaryotic RNRs and several evolutionarily unrelated bacterial class I subclasses, the NrdB component contains a stable tyrosyl radical in the vicinity of a diferric metal center (Ia). In another bacterial subclass (Ic), a mutational change in the radical-carrying tyrosine to phenylalanine is accompanied by a mixed valent MnII/MnIV/FeIII metal center (10, 11). Recently, a metal independent subclass (Ie) with an intrinsically modified dopa radical cofactor was discovered (12).

All class I RNRs contain a C-terminal redox-active cysteine pair in NrdA that functions as a reductant of a cysteine pair in the active site that is oxidized during catalysis. Physiological regeneration of active NrdA is performed by members of the redoxin family, with NADPH as ultimate electron source

2 The abbreviations used are: RNR, ribonucleotide reductase; a-site, allosteric overall activity site in the ATP cone; GEMMA, gas-phase electrophoretic macromolecule analysis; Grx, glutaredoxin; HED, 2-hydroxyethyl disulfide; ITC, isothermal titration calorimetry; NrdBΔGrx, NrdB protein lacking 69 N-terminal residues corresponding to the glutaredoxin domain; NrdBΔ169, NrdB protein lacking the glutaredoxin domain and the ATP-cone domain; SEC, size-exclusion chromatography; s-site, allosteric specificity site in NrdA.
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Figure 1. The F. ignava nrdAB operon and some other class I RNR operons with grx fusions. Transcriptional and translational directions are from left to right. The F. tularensis operon occur in all Francisella spp., and the displayed vibrio-phae operon is found in most vibrio phages (Table S1).

(13–15). Three types of redoxin have been found to reduce the C-terminal cysteines in class I RNRs: (i) thioredoxin that receives the electrons from NADPH via thioredoxin reductase, (ii) glutaredoxin (Grx) that receives the electrons from NADPH via GSH reductase and GSH, and (iii) NrdH-redoxin that also receives the electrons from NADPH via thioredoxin reductase even though NrdH is more similar to Grx than to thioredoxin. Whereas the nrdA and nrdB genes are mostly encoded close to each other in bacteria, the trx and grx genes are usually found elsewhere in the genome. Only the nrdH gene is predominantly encoded in the vicinity of the corresponding RNR genes, which, for historical reasons, in this particular subclass are called nrdE (encoding the catalytic subunit) and nrdF (encoding the radical-generating subunit).

We discovered an intriguing fusion of a grx gene to the nrdB gene in the bacterium Facklamia ignava, resulting in an ORF encoding a fusion protein. The F. ignava NrdB fusion protein consists of an N-terminal Grx domain followed by an ATP-cone domain and then the radical-generating subunit. An N-terminal grx fusion to the nrdB gene in Francisella tularensis was noticed by us previously (16). The redoxin domain in both of these fusions are most similar to the grxC domain family (COG0695). Whereas the γ-proteobacterium F. tularensis is a well-studied human pathogen causing tularemia (17), the Firmicutes genus Facklamia was first described in 1997 and has since been identified in samples from a wide range of animals and as a human pathogen (18–20).

RNR has been described as a textbook example of allosteric regulation in enzymes and employs two different allosteric mechanisms to regulate the synthesis of dNTPs (21, 22). One common mechanism regulates the balance between the four dNTPs in a sophisticated feedback control at the specificity site (s-site). Additional allosteric regulation is provided by the overall activity site (a-site), which works as a general on/off switch and constitutes a separate domain called the ATP cone. In short, the enzyme is active when ATP is bound and when dATP is bound, the enzyme is turned off. We have recently shown that the ATP cone can be horizontally transferred between different RNRs and even to different subunits of the holoenzyme (8, 23). In an overwhelming number of cases, the ATP cone is an N-terminal domain of the catalytic subunit of RNR (23). F. ignava RNR instead carries an ATP cone in its NrdB protein, between the N-terminal Grx domain and the radical-generating domain. We have recently reported a similar N-terminal ATP-cone fusion to NrdB in Leuwenhoekiella blandensis (8). Both of these fusion proteins belong to the NrdBi subclass, which harbors a few additional ATP-cone::NrdB fusions.

In this study we have used the F. ignava RNR to study two major questions: does the fused Grx domain function as a reductant for the holoenzyme, and does the fused ATP cone function as a general on/off switch? To investigate these questions, we used a series of biochemical assays to show that the Grx domain is indeed an efficient reductant of F. ignava RNR and that the fused ATP-cone domain is a functional allosteric domain.

Results

Glutaredoxin fusions to RNR components

The 496-residue F. ignava (Firmicutes) NrdB fusion protein consists of an N-terminal Grx domain (residues 4–61, with the characteristic cysteine pair at residues 12 and 15) followed by an ATP-cone domain (residues 84–169) and thereafter the NrdB proper. The nrdA gene is located 46 nucleotides downstream of the nrdB gene, and the two genes conceivably form an operon (Fig. 1). The F. ignava NrdB is a member of the NrdBi phylogenetic subclass (http://rnrdb.pfammap.org), like all other NrdBs in which we have detected N-terminal ATP cones (8).

Spurred by the discovery of the Grx fusion to F. ignava NrdBi, we performed a search of the RefSeq database for combinations of RNR proteins and Grx domains. Grx fusions were found in all RNR components (NrdA, NrdB, NrdD, and NrdJ), in some cases together with an ATP cone (Fig. 1 and Table S1). Grx fused to NrdB were detected in all Francisella spp. and Allofrancisella guangzhounensis (both γ-proteobacteria; subclass NrdBk) and in 24 viruses (NrdBe, NrdBg, and NrdBk) (Fig. 1 and Table S1). In addition, a grx::nrdF fusion was found in Streptococcus pneumoniae, a grx::nrdD fusion in Lachnospiraceae bacterium TWA4, a grx::nrdJ fusion in Labrenzia aggregata, and grx::nrdA fusions in two viruses (Table S1).

Because many Firmicutes lack GSH and instead produce another low molecular weight reductant called bacillithiol (24),

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we also searched the *F. ignava* genome for the presence of GSH biosynthesis and bacillithiol biosynthesis genes (*gshA*, *gshB*, *gshF*, *bshA*, *bshB1*, *bshB2*, and *bshC*). *F. ignava* and all other *Facklamia* spp. except one, encode the bifunctional *gshF* gene that is primarily found in Firmicutes (25). The *Facklamia* *gshF* has extensive similarity primarily to the *gshA* gene (Table S2). The GSH reductase gene *gor* was only found in *Facklamia sourekii*. The closest ortholog in *F. ignava* is a mercuric(II) reductase and a dihydrolipoyl dehydrogenase, both with ~50% similarity to *F. sourekii* *gor*. There were no genes corresponding to the bacillithiol biosynthesis genes in any *Facklamia* spp., apart from a glycosyl transferase gene with some similarity to *bshA*. Our results show that *F. ignava* and other *Facklamia* spp. have the capacity to synthesize GSH.

**Redox activity of the NrdB-fused glutaredoxin**

Using a series of cysteine-to-serine mutant proteins, we delineated the reaction mechanism of the fused Grx domain. Grx proteins usually reduce RNRs via a dithiol mechanism, but e.g. a human Grx has been reported to work via a glutathionylation mechanism (26–28). To test the capacity of the Grx domain in *F. ignava* NrdB to perform a dithiol reduction, we constructed two mutant proteins with a serine instead of cysteine in one or the other of the two redox-active residues in the Grx domain (C12S and C15S) and the corresponding double mutant (C12S/C15S).

In a first set of experiments the mutants were compared with the WT protein in a redox cycle with the artificial substrate 2-hydroxyethyl disulfide (HED). As evident from fig. 2A, the WT and C15S mutant proteins reduced the HED substrate, whereas the C12S mutant and the double mutant did not. The *Km* for HED was 0.6 ± 0.09 mM for the WT protein and 1.3 ± 0.24 mM for the C15S protein, and the *Vmax* was ~2-fold higher for the WT compared with C15S at saturating HED (Fig. 2B). In a GSH titration experiment with constant HED, the *Km* for GSH was 3 ± 0.9 mM for the C15S mutant protein, and the rate was 44 μM/min (Fig. 2C), corresponding to a redox *kcat* of 7.3 s⁻¹. Activity in the absence of one cysteine demonstrates that the Grx domain in *F. ignava* NrdB can work via a monothiol mechanism utilizing Cys-12 as the redox-active cysteine in presence of HED. The behavior of the WT protein in the GSH titration experiment (Fig. 2C) cannot be explained by a pure dithiol reaction mechanism. One possible explanation is that a monothiol mechanism may interfere at higher GSH concentrations.

In a second set of experiments, we compared the ability of the WT and mutant Grx domains to function as reductants in RNR assays. High specific activity (*kcat* 1.4 ± 0.06 s⁻¹) with an apparent *Km* for GSH of 1.2 ± 0.2 mM was only obtained with the WT protein (Fig. 3). Of the mutant proteins, C12S and the C12S/C15S were deficient in ribonucleotide reduction with both 4 and 10 mM GSH, whereas their specific activity was on par with the WT enzyme when the Grx domain was bypassed using DTT as reductant (Fig. 3, inset). Interestingly, the C15S mutant protein promoted a low but significant GSH-dependent ribonucleotide reductase activity, as measured both as consumption of NADPH (Fig. 3A) and as formation of dCDP (Fig. 3C), but it was not possible to reach a *Vmax* for the RNR activity of the C15S protein even at 20 mM GSH (Fig. 3B and data not shown). The GSH concentration of *Facklamia* spp. is not known, but GSH concentrations in studied bacteria range between 0.1 and 10 mM, with Firmicutes generally on the high side (25, 29). Conceivably, the Grx fused to *F. ignava* NrdB is most efficiently promoting turnover of the *F. ignava* RNR via a dithiol mechanism, and at 10 mM GSH concentration, the C15S mutant protein can promote ~4-fold less efficient ribonucleotide reduction via a monothiol mechanism involving Cys-12.

**Substrate specificity regulation of *F. ignava* RNR via the s-site**

Using a four-substrate activity assay in the presence of saturating concentrations of the substrate specificity site (s-site) effectors ATP, dTTP, or dGTP, we found that *F. ignava* RNR has a similar specificity regulation pattern to most characterized RNRs (3). ATP stimulated the reduction of CDP and UDP, whereas dTTP stimulated the reduction of GDP, and dGTP stimulated the reduction of ADP and GDP (Fig. 4). There was also a low activity of predominantly CDP reduction in the absence of allosteric effectors. Using mixtures of allosteric effectors, we observed that dTTP-induced GDP reduction increased in the presence of ATP (Fig. 5A), as is commonly seen in RNRs (3).

**Overall activity of *F. ignava* RNR is regulated via the NrdB-linked ATP-cone**

We performed a series of activity assays with CDP as substrate to elucidate the potential roles of ATP and dATP in activating and inhibiting the enzyme. The presence of ATP acti-

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**Figure 2. HED reduction capacity of *F. ignava* NrdB.** Glutaredoxin activity was measured as NADPH consumption in presence of 0.1 μM protein. A, WT and mutant proteins in the presence of 0.75 mM HED and 4 mM GSH. Assays were performed in triplicate with standard deviations shown. B, HED titration of WT and C15S proteins in the presence of 4 mM GSH. C, GSH titration of WT and C15S proteins in presence of 0.75 mM HED.
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...either ATP or dATP (Fig. 5, NrdA was saturated with dTTP, and GDP was used as sub-
ATP or dATP at the ATP cone of NrdB, the specificity site of 
dATP-dependent inhibition was 1.3
ATP-dependent activation was 47
complex,
formed in triplicate with standard deviations shown.
RNR activity measured as NADPH consumption in presence of 0.5
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enzyme activity at 3
activating the enzyme at low concentrations and inhibiting 
results.
To elucidate the mechanism of allosteric overall activity reg-
governed by the NrdB-linked ATP-cone, oligomer-
distribution experiments were performed by gas-phase electro-
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showed that the NrdB subunit (β) was in a dimer–tetramer 
equilibrium and that the tetramer formation was stimulated by 
dATP and suppressed by ATP (Fig. 6A). If the ATP cone is 
removed as in NrdBΔ169, the protein lost the ability to form 
tetramers, indicating that the process depends on the ATP cone 
(Fig. 6B). In the Grx deletion mutant, the ability to form tetram-
ers was decreased but not lost completely (Fig. 6B). The NrdA 
subunit (α) was in a monomer–dimer equilibrium favoring dimers, especially in the presence of dATP where the mono-
ners were below the detection limit (Fig. 6C). When both pro-
teins were mixed together with dATP, an additional peak 
corresponding to an αβ4 complex and to a minor extent also an αβ4 complex (Fig. 6D). In the absence of alloste-
eric effectors or in the presence of ATP, αβ2 complexes were 
formed instead. The subunit compositions of the 234-, 344-, 
and 470-kDa peaks were determined by comparing the results 
with each subunit alone. NrdB tetramer formation was very 
efficient in the absence of effectors or in the presence of ATP, 
indicating that the 234-kDa peak only to a minor extent can be 
explained by NrdB tetramers and mostly contains αβ2 complexes, 
resulting from the interaction of NrdA and NrdB, the 
major two species formed in the absence of effectors. In the 
presence of dATP, the two major species NrdA dimers and 
NrdB tetramers interacted to form the αβ4 complex and to 
some extent also an αβ4 complex if an additional NrdA dimer 
binds.

To complement the GEMMA analyses of oligomer forma-
tion, we performed analytical size-exclusion chromatography (Fig. 7) using higher protein concentrations and physiologically 
reasonable concentrations of effectors (3 mM ATP and 0.1 mM 
dATP) (30, 31). The SEC experiments confirmed the GEMMA 
results. The NrdA protein was predominantly a dimer, and the 
dimeric form was further enhanced by binding of dATP and 
ATP to the s-site (Fig. 7A). The NrdB subunit doubled in mass 
in the presence of dATP compared with when ATP was present, 
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dimer and tetramer had larger masses than expected, indicating 
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effectors, the NrdB protein seemed to be a dimer that gradually 
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NrdA was saturated with dTTP, and GDP was used as sub-
strate, giving a starting specific activity (normalized to 100% in 
Fig. 5) even in the absence of added ATP. In WT NrdB Ks for 
ATP-dependent activation was 47 ± 12 μM (Fig. 5A), and Ks for 
dATP-dependent inhibition was 1.3 ± 0.23 μM (Fig. 5B). The 
activity of the deletion mutant that lacks both the Grx domain 
and the ATP cone (NrdBΔ169) was not affected by addition of 
either ATP or dATP (Fig. 5, A and B). The initial kcat of the 
NrdBΔ169 in the presence of dTTP-loaded NrdA was 2.2 s-1,
i.e. almost three times higher than that of full-length NrdB (0.8 
s-1). However, ATP addition increased the activity of WT 
NrdB to 1.8 s-1 (Fig. 5A), i.e. on par with the NrdBΔ169 mutant 
and other RNR enzymes. Titration with dADP inhibited the 
WT enzyme activity (Fig. S2), although less strongly than dATP did.

dATP binding to NrdB induces formation of higher oligomeric 
complexes

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both with ATP and without effector, and a larger species with dATP. There was a gradual movement to a larger species when the protein concentration was increased up to a mass indicating an F. ignava NrdB complex.

Binding of nucleotides to F. ignava NrdB was investigated using isothermal titration calorimetry (ITC). Binding curves for dATP and ATP to NrdB at 20 °C were consistent with a single set of binding sites (Fig. 8). In dATP titrations, the fitted apparent $N$ value was significantly above 1 ($n = 1.4 \pm 0.1$), suggesting that the protein binds two dATP molecules per ATP cone provided our preparation contains 70% active protein. Fit of ATP titrations, performed with the same protein preparation and at the same day resulted in $n = 0.55 \pm 0.02$, suggesting binding of only one ATP per ATP cone. $K_d$ for the different nucleotides (Fig. 8E) indicated a 20-fold lower affinity for ATP compared with dATP. Thermodynamic parameters (Fig. 8E) indicated that the interactions are predominantly enthalpy-driven, with negative $\Delta H$ values of $-80$ and $-60$ kJ/mol for dATP and ATP, respectively. As observed earlier for L. blandensis NrdB (8) dATP also binds to the ATP cone of F. ignava NrdB with a $K_d$ of 5.8 $\mu$M at 25 °C, i.e. considerably weaker than the $K_d$ for dATP (Fig. S3).

We performed an additional set of ITC experiments at 10 °C, which resulted in lower $K_d$ values (0.4 $\mu$M for dATP and 6.8 $\mu$M for ATP) but otherwise similar conclusions. Fitted stoichiometries were $1.5 \pm 0.1$ for dATP and $0.57 \pm 0.01$ for ATP in agreement with the 20 °C results and underscoring our interpretation that the F. ignava NrdB protein binds two molecules of dATP and one molecule of ATP.

**Type of radical cofactor in the F. ignava NrdB protein**

To elucidate the nature of the radical cofactor in the F. ignava NrdB protein, we employed EPR spectroscopy. X-band EPR spectra recorded on samples of NrdB expressed in the presence of excess Mn$^{2+}$ and purified via affinity chromatography revealed an intense multiline signal with a signal width of 125–130 mT (Fig. 9). The signal varied in a uniform fashion in the interval 5–15 K and can thus be attributed to a single paramagnetic species (Fig. 9, compare 5-, 10-, and 15-K spectra). Increasing the temperature further resulted in a complete disappearance of the signal at 30 K, with no new signal appearing. The shape, width, and temperature dependence of the signal is in good agreement with an anti-ferromagnetically coupled Mn$^{III}$/Mn$^{IV}$ complex, where the complex line shape is a result of an $S = 1/2$ system where the unpaired electron is interacting with two $I = 5/2$ manganese centers. In a biological context, similar Mn$^{III}$/Mn$^{IV}$ species have been observed in the case of superoxidized manganese catalase and as a short-lived intermediate during the assembly of the Mn$^{III}_{2}$–Y cofactor in NrdF (32, 33). The presence of such an intense multiline signal in our purified samples suggests that this high-valent species is stable at least in the time scale of hours.

![Figure 4](attachment:image1.png) **Figure 4.** Substrate specificity of F. ignava class I RNR. Enzyme assays were performed in mixtures with 0.5 mM of each of the four substrates (ADP, CDP, GDP, and UDP) and a saturating concentration of one effector nucleotide at a time. Assays were performed in triplicate with standard deviations shown.

![Figure 5](attachment:image2.png) **Figure 5.** Inhibition and activation of WT and mutant enzyme activity by dATP and ATP. A and B, ATP titration (A) and dATP titration (B) of enzyme loaded with 2 mM dTTP and GDP as substrate. Specific activities of NrdB proteins were measured with a 10-fold excess of NrdA. WT NrdB (●) had a starting activity of 830 ± 120 nmol/min/mg in the absence of added ATP and reached a $V_{satur}$ of 1850 ± 200 nmol/min/mg ($k_{cat} = 1.8$ s$^{-1}$) in the presence of ATP, whereas NrdBΔ169 (○) had a specific activity of 3400 ± 500 nmol/min/mg ($k_{cat} = 2.2$ s$^{-1}$) in the absence of ATP that was not affected by addition of ATP or dATP.
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RNR activity in mixtures of F. ignava and L. blandensis NrdA and NrdB proteins

The NrdB core of the firmicute F. ignava from residue 170 and onwards has extensive similarity (61% sequence identity; Fig. S4) to the core of the NrdB protein from the flavobacterium L. blandensis. They both harbor a mixed valent MnIII/MnIV center with capacity to initiate the radical-based enzyme reaction (this study and Ref. 8). Both the F. ignava NrdB and the L. blandensis NrdB also harbor an ATP-cone domain that functions as an on/off switch for the activity of its RNR holoenzyme by forming tetrameric NrdB structures in presence of dATP to which the NrdA protein is prevented from binding in a productive fashion (above and in Ref. 8). However, the ATP cones of F. ignava and L. blandensis NrdB proteins are more different (28% sequence identity; Fig. S4) and align extensively only over their C-terminal sequences, which in the L. blandensis structure has been shown to interact primarily with one of the two bound dATP molecules (8). The similarity between the two corresponding NrdA proteins is extensive (61% sequence identities; Fig. S5). Based on these similarities, we designed a set of experiments to test whether RNR enzyme activity can be achieved in heterologous mixtures of F. ignava and L. blandensis NrdA and NrdB proteins and whether the unique Grx domain would disturb a heterologous interaction. Heterologous mixtures of class I RNR subunits have primarily been tested for distantly related enzymes, e.g. class I RNR subunits from Escherichia coli and bacteriophage T4 with negative results (34). On the other hand, several thioredoxins are known to cross-react with heterologous RNRS, whereas Grxs usually do not (35).

Fig. 10 shows that the heterologous F. ignava NrdA/L. blandensis NrdB holoenzyme was active and regulated by ATP and dATP via the ATP cone linked to L. blandensis NrdB, whereas the heterologous L. blandensis NrdA/F. ignava NrdB holoenzyme was inactive. The same was true for heterologous mixtures with F. ignava NrdBΔGrx, as well as for F. ignava NrdBΔ169 (Fig. 10A). Kᵢ for ATP was ∼300 μM, and Kᵢ for dATP was ∼70 μM for the ATP cone of L. blandensis NrdB in the heterologous mixture (Fig. 10B), i.e. more than 3 times higher than the Kᵢ(dATP) of 96 μM and the Kᵢ(dATP) of 20 μM for the L. blandensis holoenzyme (8). The Vₐ₉ obtained in the heterologous holoenzyme was 250 nmol/min/mg, corresponding to a k₉ of ∼0.2 s⁻¹, approximately a fourth of the activity of the L. blandensis holoenzyme (Fig. 10A).

Discussion

We have shown that the multidomain radical-generating component of the F. ignava class I RNR contains a gene fusion of an N-terminal Grx that is fully functional as a reductant of the RNR holoenzyme and an ATP-cone that serves as a general on/off switch of the enzyme. We also identified fusions of Grx-domains with NrdB proteins in Francisella spp., A. guangzhouensis, and several viruses (Fig. 1 and Table S1), but none of the other cases were in the NrdBi subclass that the F. ignava protein belongs to. This strongly suggests that the F. ignava grx:nrdBi fusion was a separate evolutionary event, not related to the grx:nrdB fusions discovered in other organisms and viruses. On the contrary, the presence in F. ignava of a fusion between an ATP-cone domain and NrdBi appears to be the result of horizontal gene transfer because the majority of ATP cones fused with nrdBi genes occurs in flavobacteria (8).
Glutaredoxin and ATP-cone fusions to ribonucleotide reductase

Figure 7. Size-exclusion chromatography analysis of F. ignava RNR components in the presence of nucleotides. A, 5 and 10 μM of the NrdA subunit was analyzed in the presence of 3 mM ATP or 100 μM dATP or without effectector. B, corresponding analysis of the NrdB subunit. In this case the experiment without effectector was performed at 1.25, 2.5, 5, and 10 μM protein. The position of the peaks indicate a larger size than expected, which is typical for elongated proteins, and the interpretation above the peaks is based on a comparison with the GEMMA results. C, analysis of the combination of both subunits. Each subunit was used at 10 and 20 μM concentration except in the experiment without effectector, where only 10 μM was used.

thus appears most parsimonious to suggest that the ATP cone::NrdBi fusion gene was first transferred to F. ignava and that the gene was subsequently fused with the grx gene in the F. ignava genome.

Grx was first described as a physiological reductant for RNR in E. coli (26) and has since also been observed to be involved in sulfate assimilation, detoxification, and development and proliferation, primarily in eukaryotic cells (27, 36). Similarly to other redoxins, the active site of dithiol Grxs consists of a cysteine pair separated by two residues (predominantly -CPYC-) (27). The corresponding sequence in F. ignava Grx::NrdBi is -CPWC- (Fig. S4). Grxs differ from other redoxins in that they form mixed disulfides with GSH and also promote glutathionylation/deglutathionylation reactions, which may lead to reduction of protein disulfides (36). E. coli Grx has been shown to use the dithiol mechanism in its reduction of E. coli RNR, whereas a human Grx was interpreted to reduce mammalian RNR via a glutathionylation mechanism (28). However, recent theoretical studies, as well as thorough experimental studies on Grx-dependent reduction of protein disulfides with heterologous components from eukaryotic and bacterial origins, show that the monothiol–dithiol mechanisms occur in parallel and that GSH concentration and dominance of specific steps in the mechanism determine the preferred path taken (37–39). In this study we show that the Grx::NrdB fusion protein of F. ignava can reduce its class I RNR holoenzyme via a dithiol mechanism and that the C15S mutant in the Grx active site can reduce RNR less efficiently via a monothiol mechanism (Fig. 11). To our knowledge this is the first demonstration of parallel dithiol–monothiol reduction mechanisms in a native system between Grx and its oxidized substrate from the same species.

Over and above the fused Grx domain, the remarkable F. ignava NrdB protein exhibits two other unusual characters: a fused ATP cone and a mixed valent Mn\textsuperscript{III}/Mn\textsuperscript{IV} metal site. Both of these features were recently described in L. blandensis NrdB and in several NrdBi proteins in Flavobacteriales (8). The ATP cone in F. ignava NrdB binds two dATP molecules like the cone in L. blandensis NrdB, but their amino acid sequences across the ATP cones are surprisingly dissimilar in the N-terminal half (Fig. S4). This may relate to our finding that the cone in F. ignava binds only one ATP molecule, whereas the cone in L. blandensis binds two ATP molecules. The ATP-loaded active F. ignava holoenzyme is αβ, whereas the dATP-inhibited complexes are β\textsubscript{2} for NrdB and α\textsubscript{2}β\textsubscript{4} plus α\textsubscript{2}β\textsubscript{4} for the holoenzyme. All of these complexes were also observed in the L. blandensis RNR (8).

The mixed valent Mn\textsuperscript{III}/Mn\textsuperscript{IV} metal site in F. ignava NrdB has a distinct EPR signal in the temperature range of 5–15 K, with no other manganese-related EPR signals at 30 K and no trace of a tyrosyl radical. A similar high valent manganese dimer was recently found to be present in NrdB from L. blandensis (8). Later, Boal and co-workers (9) also reported a similar multiline signal in Flavobacterium johnsoniae class I RNR. However, in both of these latter cases, the multiline feature represented only a fraction of the total metal content. Conversely, in our F. ignava NrdB samples presented here, the Mn\textsuperscript{III}/Mn\textsuperscript{IV} signal is clearly the dominant metal species. These observations underscore the catalytic relevance of the Mn\textsuperscript{III}/Mn\textsuperscript{IV} signal and support the notion that the NrdBi proteins represent a new subclass of class I RNRs, denoted subclass Id (6, 8, 9).

The similarities between F. ignava and L. blandensis NrdB proteins is further manifested by the enzyme activity observed in a heterologous mixture of F. ignava NrdA and L. blandensis NrdB, which is almost a third of that in the L. blandensis...
holoenzyme. Conversely, heterologous mixtures of *L. blandensis* NrdA and *F. ignava* NrdB lack activity even in the absence of the Grx domain or for the NrdB/H9004169 protein that lacks both the Grx domain and the ATP cone. The *F. ignava* NrdB core may have undergone significant structural changes to accept the fusion of the Grx domain, which may also pertain to the divergent N-terminal of the ATP-cone sequence. Future studies will be directed to clarify this point.

All in all, we have shown that the unique NrdB protein in *F. ignava* carries an N-terminal Grx domain with capacity to act as a physiological reductant of its corresponding holoenzyme via a dithiol mechanism and less efficiently via a monothiol mechanism in the C15S mutant variant. The ATP-cone domain, which is fused between the Grx domain and the NrdB core, functions as an allosteric on/off switch, promoting an enzymatically active [H9251]2[H9252]2 complex in presence of ATP and enzymatically inactive [H9251]4[H9252]4 and [H9251]4[H9252]4 complexes in the presence of dATP. The radical cofactor in *F. ignava* NrdB is a mixed valent dinuclear MnIII/MnIV site, which forms in the absence of an NrdI activase and lacks a tyrosyl radical.

*F. ignava* NrdB is an enthralling illustration of how RNR subclasses continuously evolve via gain and loss of accessory domains and RNR-related proteins.

### Experimental procedures

#### Bioinformatics

The RefSeq database (40) was downloaded March 16, 2018, and searched with the Pfam (41) profiles for Grx (PF00462) and the ATP cone (PF03477) plus profiles developed in-house for RNR proteins (http://rnrdb.pfitmap.org)3 using the HMMER...
software version 3.1b2 (42). For RNR proteins, only hits covering at least 90% of the length of the profile were kept. For the Grx and ATP-cone profiles, only hits with a higher bitscore than the Pfam-specified gathering scores (21.50 in both cases) were kept.

Cloning

DNA fragments encoding NrdA (<i>F. ignava</i> WP_006702002) and NrdB (<i>L. blandensis</i> EKB53615/WP_006702003) were amplified by PCR from <i>F. ignava</i> CCUG 37419 genomic DNA, obtained from the Culture Collection at the University of Gothenburg, using specific primers: NrdA, FiR1_For 5'-tcctcCATATGACCGCA-CAATTAAAGAATC-3' and FiR1_Rev 5'-cagaGGATCCTT-AAGCTTAAGCTCAAGCTAAGC-3'; NrdB: FiR2_For 5'-tcctcCATATGACTCAAGTACAAGTTTATAG-3' and FiR2_REV 5'-cagaGGATCCTTAGAATAGGTCGTCGGC-3'.

The PCR products were purified, cleaved with NdeI and BamHI restriction enzymes, and inserted into a pET-28a(+) expression vector (Novagen, Madison, WI). The obtained constructs pET-nrdA, pET-nrdB, pET-nrdB<sub>Grx</sub>, pET-nrdB<sub>169</sub>, pET-nrdB<sub>C12S</sub>, pET-nrdB<sub>C15S</sub>, and pET-nrdB<sub>C12SC15S</sub> containing nucleotide mismatches T34A, G44C, and T34A/G44C, respectively, were ordered from GenScript.

Protein expression

Overnight cultures of <i>E. coli</i> BL21(DE3)/pET28a(+) bearing pET-nrdA, pET-nrdB, pET-nrdB<sub>Grx</sub>, pET-nrdB<sub>Δ169</sub>, pET-nrdB<sub>C12S</sub>, pET-nrdB<sub>C15S</sub>, or pET-nrdB<sub>C12SC15S</sub> were diluted to an absorbance at 600 nm of 0.1 in LB (Luria-Bertani) liquid medium, containing kanamycin (50 μg/ml) and shaken vigorously at 37 °C. At an absorbance of A<sub>600</sub> 0.8 isopropyl-β-D-thiogalactopyranoside (Sigma) was added to a final concentration of 0.5 mM; the cultures expressing NrdB were further supplemented with MnSO4 (final concentration, 0.5 mM) during the induction. The cells were grown overnight at 30 °C and harvested by centrifugation.

Protein purification

The cell pellet was resuspended in lysis buffer: 50 mM Tris-HCl, pH 7.6, containing 300 mM NaCl, 10% glycerol, 2 mM DTT, 10 mM imidazole, 1 mM phenylmethylsulfonyl fluoride. The cells were disrupted by high pressure homogenization, and the lysate was centrifuged at 18,000 × g for 45 min at 4 °C. The
recombinant His-tagged protein using ÄKTA prime system (GE Healthcare): the supernatant was loaded on a HiTrap FF nickel-Sepharose column (GE Healthcare), equilibrated with lysis buffer (without phenylmethylsulfonyl fluoride), washed thoroughly with buffer, and eluted with buffer containing 500 mM imidazole.

NrdB_C125, NrdB_C155, NrdB_C12SC155, and the WT NrdB were used for measuring the redox activity of the NrdB fused Grx were then desalted on a Sephadex G-25 PD10 column (GE Healthcare) equilibrated with buffer containing 50 mM Tris-HCl, pH 7.6, 300 mM NaCl, 10% glycerol, and 1 mM DTT; frozen in liquid nitrogen; and stored at −80 °C until used.

For NrdA, NrdB, NrdBΔGrx, and NrdBΔ169, further purification was accomplished by FPLC on a 125-ml column packed with HiLoad 16/60 Superdex 200-pg column (GE Healthcare) using ÄKTA prime system, equilibrated with buffer containing 50 mM Tris-HCl, pH 7.6, 300 mM NaCl, 10% glycerol, and 2 mM DTT. Eluted protein was frozen until used.

NrdA was further applied to hydrophobic interaction chromatography using the HiLoad 16/60 phenyl-Sepharose column (GE Healthcare) in 50 mM Tris-HCl, pH 7.6, 2 mM DTT, 0.75 mM (NH₄)₂SO₄; washed extensively (15 column volumes) with the same buffer; and eluted with buffer without ammonium sulfate. The protein was resuspended in excess of buffer containing 50 mM Tris-HCl, pH 7.6, 300 mM NaCl, 10% glycerol, 2 mM DTT; concentrated; and frozen until used. The hydrophobic interaction chromatography removed residual nucleotide contamination from NrdA. L. blandensis NrdA and NrdB were expressed and purified as previously described (8).

Protein concentrations were determined by measuring the UV absorbance at 280 nm based on protein theoretical extinction coefficients 99,700 M⁻¹ cm⁻¹ for NrdA, 72,770 M⁻¹ cm⁻¹ for NrdB (and cysteine to serine mutants), 54,320 M⁻¹ cm⁻¹ for NrdBΔGrx, and 51,340 M⁻¹ cm⁻¹ for NrdBΔ169. Protein purity was evaluated by SDS–PAGE (12%) stained with Coomassie Brilliant Blue. Proteins were concentrated using Amicon Ultra-15 centrifugal filter units (Millipore), frozen in liquid nitrogen, and stored at −80 °C until used. For EPR measurements, NrdBΔ169 was purified using affinity chromatography as described above but transferred to EPR tubes and flash-frozen in liquid nitrogen in EPR tubes immediately upon elution.

**RNR activity measurements**

RNR activity assays were performed at room temperature in 50 mM Tris-HCl, pH 8.0, in volumes of 50 μl. Reaction conditions giving maximal activity were determined experimentally. In a standard reaction the constituents were 10 mM DTT, 40 or 20 mM Mg(CH₃CO₂)₂ (when NrdA of F. ignava or L. blandensis was used, respectively), 10 mM KCl, 0.8 mM CDP, and various concentrations of allosteric effectors ATP or dATP. Mixtures of 0.1–1 mM of NrdB, 0.07 mM NrdBΔ169, 0.5 mM NrdB_C12S, NrdB_C155, or NrdB_C12SC155 and a 10-fold excess of NrdA were used. In specific experiments some components were varied as indicated in the text.

In experiments aimed to determine the redoxin activity of the NrdB-fused Grx, DTT was omitted. Instead, 4 or 10 mM reduced GSH, 11 μg ml⁻¹ GSH reductase (from yeast; Sigma) and 1 mM NADPH were added to the reaction mixtures. CDP (0.8 mM) was used as substrate, and ATP (3 mM) was used as effector. Protein concentration of 0.5 μM for WT NrdB, NrdB_C12S, NrdB_C155, or NrdB_C12SC155 were used in combination with 5 μM NrdA.

When dTTP (2 mM) was used as an s-site effector, 0.8 mM GDP was used as substrate. In the four-substrate assays, the substrates CDP, ADP, GDP, and UDP were simultaneously present in the mixture at concentrations of 0.5 mM each with 2 mM of one of the effectors (ATP, dTTP, or dGTP). The substrate mixture was added last to start the reactions.

Enzyme reactions were incubated for 2–30 min at room temperature and then stopped by the addition of methanol. Substrate conversion was analyzed by HPLC using a Waters Symmetry C18 column (150 × 4.6 mm, 3.5-μm pore size) equilibrated with buffer A. Samples of 25–100 μl were injected and eluted at 0.4 ml/min at 10 °C with a linear gradient of 0–30% buffer B over 40 min for the separation of CDP and dCDP or 0–100% buffer B over 45 min for the separation of GDP and dGDP (buffer A: 10 mM potassium phosphate buffer, pH 7.0, supplemented with 10 mM tributylammonium hydroxide; buffer B: 30% methanol in 50 mM potassium phosphate buffer, pH 7.0, supplemented with 10 mM tributylammonium hydroxide). Compound identification was achieved by comparison with injected standards. Relative quantification was obtained by peak height measurements in the chromatogram (UV absorbance at 271 or 254 nm) in relation to standards. Specific activities are given as nmol product formed per min and mg of protein.

From a direct plot of activity versus concentration of effector, the $K_i$ values for binding of effectors to the s-site and the a-site were calculated in SigmaPlot using the following equation.

$$v = V_{\text{max}} \times \frac{(\text{dNTP})}{(K_i + \text{dNTP})}$$

$K_i$ for noncompetitive dATP inhibition at NrdB was calculated in SigmaPlot using the following equation.

$$v = V_{\text{max}} \times (1 + (\text{dNTP})/K_i)$$

**Photometric activity assays**

Photometric assays for NrdB-fused Grx based on the artificial electron acceptor HED were performed as described in earlier studies (43, 44). The standard Grx assay contained 50 mM Tris, pH 8.0, 0.1 mg/ml BSA, 11 μg ml⁻¹ GSH reductase (from Saccharomyces cerevisiae), 4 mM GSH, 0.75 mM HED, and 0.4 mM NADPH. The above ingredients were mixed and incubated for 3 min, after which the reaction was started by the addition of 0.1 μM WT or mutant Grx (NrdB fused). The reference cuvette contained all ingredients, except Grx. $A_{340}$ was recorded for 3 min at room temperature using a Lambda 35 UV-visible spectrophotometer (PerkinElmer Life Science). Linear decrease in $A_{340}$ was used to calculate moles of NADPH consumed using its extinction coefficient of 6220 M⁻¹ cm⁻¹.

The combined redoxin/RNR assays contained 0.5 μM NrdB, 5 μM NrdA, the indicated amount of GSH, 11 μg ml⁻¹ GSH reductase, 0.25 mM NADPH, 10 mM Mg(CH₃CO₂)₂, and 3 mM ATP. The reaction was started by the addition of 0.8 mM CDP. The reaction was monitored by the change of $A_{340}$ using a Cary
60 UV-visible spectrophotometer (Agilent Technologies). In the calculation of specific activity, 1 mol of consumed NADPH equals formation of 1 mol of dCDP.

**GEMMA analysis**

In GEMMA, biomolecules are electrosprayed into gas phase and neutralized to singly charged particles, and the gas-phase electrophoretic mobility is measured with a differential mobility analyzer. The mobility of an analyzed particle is proportional to its diameter, which therefore allows for quantitative analysis of the different particle sizes contained in a sample (45). The GEMMA instrumental setup and general procedures were as described previously (46). NrdA, NrdB, NrdBΔGrx, and NrdBΔ169 proteins were equilibrated by Sephadex G-25 chromatography into a buffer containing 100 mM NH₄CH₃CO₂, pH 7.8, and 2 mM DTT. Prior to GEMMA analysis, the protein samples were diluted to a concentration of 0.05 mg/ml in a buffer containing 20 mM NH₄CH₃CO₂, pH 7.8, 1 mM DTT, 0.005% (v/v) Tween 20, nucleotides (when indicated), and Mg(CH₃CO₂)₂ (equimolar to the total nucleotide concentration), incubated for 5 min at room temperature, centrifuged, and applied to the GEMMA instrument. The runs were conducted at low flow rate, resulting in 1.4 – 2 p.s.i. pressure. The GEMMA system contained the following components: 3480 electrospray aerosol generator, 3080 electrostatic classifier, 3085 differential mobility analyzer, and 3025A ultrafine condensation particle counter (TSI Corp., Shoreview, MN).

**Analytical SEC**

The SEC experiments were performed at room temperature with a Superdex™ 200 10/300 column (GE Healthcare) equilibrated with a mobile phase containing 50 mM KCl, 10 mM MgCl₂, 0.1 mM DTT, and 50 mM Tris-HCl, pH 7.6. When nucleotide-dependent protein oligomerization was studied, 3 mM ATP or 0.1 mM dATP was also included in the mobile phase. The injection loop volume was 100 μl, and the flow rate was 0.5 ml/min. The UV trace was recorded with a Jasco UV-2075 Plus detector (Jasco Inc., Easton, MD) at 290 nm to limit the absorbance from the nucleotides. The proteins were incubated in mobile phase for 5 min prior to injection onto the column.

**Isothermal titration calorimetry measurements**

ITC experiments were carried out on a MicroCal ITC 200 system (Malvern Instruments Ltd.) in a buffer containing 50 mM Tris, pH 7.65, 300 mM NaCl, 10% glycerol, 2 mM tris(2-carboxyethyl)phosphine, and 10 mM MgCl₂. Measurements were done at 20 and 10 °C. The initial injection volume was 0.5 μl over a duration of 1 s. All subsequent injection volumes were 2–2.5 μl over 4–5 s with a spacing of 150–180 s between the injections. Data for the initial injection were not considered. For dATP binding analysis, the concentration of NrdB in the cell was 40 μM, and dATP in the syringe was 600 μM. For titration of ATP into NrdB, cell and syringe concentrations were 103 μM NrdB and 1.2 mM ATP. The data were analyzed using the built-in one set of sites model of the MicroCal PEAQ-ITC analysis software (Malvern Panalytical). Standard deviations in thermodynamic parameters, N and Kᵋ, were estimated from the fits of three different titrations.

**EPR spectroscopy**

Measurements were performed on a Bruker ELEXYS E500 spectrometer using an ER049X SuperX microwave bridge in a Bruker SHQ0601 cavity equipped with an Oxford Instruments continuous flow cryostat and using an ITC 503 temperature controller (Oxford Instruments). The Xepr software package (Bruker) was used for data acquisition and processing of spectra.

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**References**


A glutaredoxin domain fused to the radical-generating subunit of ribonucleotide reductase (RNR) functions as an efficient RNR reductant

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