Reconstructed ancient nitrogenases suggest Mo-specific ancestry

Amanda K. Garcia¹, Hanon McShea²,a, Bryan Kolaczkowski³, Betül Kaçar¹,4

¹Department of Molecular and Cellular Biology, University of Arizona, Tucson, Arizona
²Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, Massachusetts
³Department of Microbiology and Cell Science, University of Florida, Gainesville, Florida
⁴Department of Astronomy and Steward Observatory, University of Arizona, Tucson, Arizona
aPresent address: Department of Earth System Science, Stanford University, Stanford, California

Corresponding author information:

Betul Kacar
1007 E. Lowell St., Tucson, AZ 85721
betul@arizona.edu

ABSTRACT

The nitrogenase metalloenzyme family, essential for supplying fixed nitrogen to the biosphere, is one of life’s key biogeochemical innovations. The three isozymes of nitrogenase differ in their metal dependence, each binding either a FeMo-, FeV-, or FeFe-cofactor for the reduction of nitrogen. The history of nitrogenase metal dependence has been of particular interest due to the possible implication that ancient marine metal availabilities have significantly constrained nitrogenase evolution over geologic time. Here, we combine phylogenetics and ancestral sequence reconstruction — a method by which inferred, historical protein sequence information can be linked to functional molecular properties — to reconstruct the metal dependence of ancient nitrogenases. Inferred ancestral nitrogenase sequences at the deepest nodes of the phylogeny
suggest that ancient nitrogenases were Mo-dependent. We find that active-site sequence identity can reliably distinguish extant Mo-nitrogenases from V- and Fe-nitrogenases, as opposed to modeled active-site structural features that cannot be used to reliably classify nitrogenases of unknown metal dependence. Taxa represented by early-branching nitrogenase lineages lack one or more biosynthetic \( nifE \) and \( nifN \) genes that are necessary for assembly of the FeMo-cofactor, suggesting that early Mo-dependent nitrogenases may have utilized an alternate pathway for Mo-usage predating the FeMo-cofactor. Our results underscore the profound impacts that protein-level innovations likely had on shaping global biogeochemical cycles throughout Precambrian, in contrast to organism-level innovations which characterize Phanerozoic eon.

**KEYWORDS**

ancestral sequence reconstruction, cofactor pocket, metalloenzyme, nitrogenase, nitrogen fixation
1 INTRODUCTION

All known life requires nitrogen for the synthesis of essential biomolecules, including nucleotides and amino acids. Though the atmosphere contains nearly 80% N\(_2\) by volume, most organisms are not able to assimilate N\(_2\) due to the enormous energetic cost of breaking the N≡N bond (MacKay & Fryzuk, 2004). Select bacteria and archaea called diazotrophs accomplish biological nitrogen fixation by nitrogenase metalloenzymes (E.C. 1.18.6.1), which catalyze the reduction of N\(_2\) to bioavailable NH\(_3\). Nitrogenases are an ancient family of enzymes; oldest isotopic biosignatures interpreted as evidence of nitrogenase activity date to ~3.2 billion years (Ga) (Stueken, Buick, Guy, & Koehler, 2015). Because nitrogen has been suggested to be an important limiting nutrient on geologic timescales (Falkowski, 1997), nitrogenases have likely played a key role in the expansion of the biosphere since the Archean.

The nitrogenase family consists of three homologous isozymes (Boyd, Hamilton, & Peters, 2011; Raymond, Siefert, Staples, & Blankenship, 2004) named for the differential metal content of the active-site cofactor: Mo-nitrogenase (Nif, encoded by \(nif\)), V-nitrogenase (Vnf, encoded by \(vnf\)), and Fe-nitrogenase (Anf, encoded by \(anf\)) (Bulen & LeComte, 1966; Eady, 1996; Joerger & Bishop, 1988; Mus, Alleman, Pence, Seefeldt, & Peters, 2018) (Figure 1). Of the three isozymes, Mo-nitrogenases are the most common and widely studied; V- and Fe-nitrogenases are comparatively rarer and only known in taxa that also possess Mo-nitrogenase (Boyd, Hamilton, et al., 2011; Dos Santos, Fang, Mason, Setubal, & Dixon, 2012). All three nitrogenase isozymes are structurally and functionally similar, each containing two protein components: the electron delivery component (NifH, VnfH, or AnfH) is a homodimer and the catalytic component is either an \(\alpha_2\beta_2\) heterotetramer (MoFe protein, NifDK) or an \(\alpha_2\beta_2\gamma_2\) heterohexamer (VFe protein, VnfDGK).
or FeFe protein, AnfDGK) (**Figure 1a**) (Bulen & LeComte, 1966; Hales, Case, Morningstar, Dzedan, & Mauterer, 1986; Schmid et al., 2002; D. Sippel & Einsle, 2017). During catalysis, the electron delivery component transiently associates with and delivers electrons to the catalytic component (Hageman & Burris, 1978). Electrons accumulate at the active site for N\textsubscript{2} binding and reduction (Hoffman, Lukoyanov, Yang, Dean, & Seefeldt, 2014), which houses a homocitrate-metallocluster cofactor unique to each isozyme: the FeMo-cofactor in Nif (**Figure 1b**), FeV-cofactor in Vnf (**Figure 1c**), and FeFe-cofactor in Anf (**Figure 1d**) (Eady, 1996; Harris, Lukoyanov et al., 2018; Krahn et al., 2002; Mus et al., 2018; D. Sippel & Einsle, 2017; Spatzal et al., 2011). Available spectral evidence suggest that these cofactors are structurally similar except for the substitution of a Mo, V, or additional Fe atom (Eady, 1996; Krahn et al., 2002; Daniel Sippel et al., 2018; Spatzal et al., 2011). Nevertheless, biochemical studies demonstrate variable catalytic properties among the three nitrogenase isozymes, including differential abilities to reduce alternative substrates (Harris, Lukoyanov et al., 2018; Harris, Yang, Dean, Seefeldt, & Hoffman, 2018; B. Hu et al., 2018; Y. Hu, Lee, & Ribbe, 2011; Zheng et al., 2018). These catalytic variations likely arise due to a combination of the aforementioned cofactor compositional differences as well as differences in the surrounding protein environment (Fixen et al., 2016; Harris, Yang, et al., 2018; Lee et al., 2018; Rebelein, Lee, Newcomb, Hu, & Ribbe, 2018; Zheng et al., 2018).
**Figure 1.** Structure and genetics of the three nitrogenase isozymes. (a) Structure of the *A. vinelandii* Mo-nitrogenase enzyme complex (NifHDK; PDB 1M34 (Schmid et al., 2002)) and V-nitrogenase VFe-protein component (VnfDGK; PDB 5N6Y (D. Sippel & Einsle, 2017)) (Fe-nitrogenase structure not previously published). The active-site FeMo-cofactor of Mo-nitrogenase and FeV-cofactor of V-nitrogenase are circled. (b-d) Catalytic genes and cofactor structures of *A. vinelandii* Mo-nitrogenase (b; PDB 3U7Q (Spatzal et al., 2011)), V-nitrogenase (c; PDB 5N6Y (D. Sippel & Einsle, 2017)), and Fe-nitrogenase (d; proposed structure (Harris, Lukoyanov, et al., 2018)). Cofactor atom coloring is as follows: C, tan; Fe, rust; Mo, cyan; N, blue; O, red; S, yellow.

Metal cofactor incorporation in nitrogenases is constrained at multiple levels. At the level of single enzyme functionality, nitrogenase biochemical and biophysical properties shape metal binding specificity. At a higher level, constraints arise from the partner proteins that constitute the biosynthetic mechanism for active-site cofactor assembly and insertion, best studied in Mo-nitrogenases (Curatti et al., 2007; Y. Hu & Ribbe, 2011; Rubio & Ludden, 2008). In the *Azotobacter vinelandii* (*A. vinelandii*) model system, FeMo-cofactor assembly requires several
associated proteins encoded within the *nif* gene cluster (Curatti et al., 2007; Dos Santos et al., 2012; Shah, Allen, Spangler, & Ludden, 1994; Shah, Imperial, Ugalde, Ludden, & Brill, 1986; St John et al., 1975; Tal, Chun, Gavini, & Burgess, 1991). Most notably, the biosynthetic *nifB, nifE,* and *nifN* genes have been demonstrated, in addition to the catalytic *nifHDK* genes, to be — perhaps minimally — required for FeMo-cofactor assembly and Mo-nitrogenase function (Curatti et al., 2007; Dos Santos et al., 2012; Shah et al., 1994; Shah et al., 1986; St John et al., 1975; Tal et al., 1991). In *A. vinelandii,* *nifE* and *nifN* loci are located just downstream of the *nifHDK* cluster, whereas the *nifB* locus is located within a separate *nif* region near other regulatory and biosynthetic *nif* genes (Setubal et al., 2009). NifB catalyzes the formation of a Fe-S-C metallocluster, a precursor that forms the core of the mature FeMo-cofactor (Allen, Chatterjee, Ludden, & Shah, 1995; Y. Hu & Ribbe, 2011; St John et al., 1975). This precursor metallocluster is then transferred to a protein heterotetramer composed of NifE and NifN subunits (Allen et al., 1995; Roll, Shah, Dean, & Roberts, 1995), homologous to NifD and NifK, respectively, and likely having arisen by gene duplication (Boyd, Anbar, et al., 2011). Within NifEN, the precursor is further modified via the addition of homocitrate and Mo, and the mature cofactor is subsequently transferred to the nitrogenase NifDK catalytic protein component (Roll et al., 1995). Unlike that for the FeMo-cofactor, the biosynthetic pathways for the formation of the FeV- and FeFe-cofactors are relatively unknown. However, transcriptional profiling of the three nitrogenase systems in *A. vinelandii* suggests that FeV- and FeFe-cofactor synthesis relies on several *nif* genes in addition to *vnf* and *anf* genes, respectively (Hamilton et al., 2011; Joerger & Bishop, 1988; Kennedy & Dean, 1992). These include *nifBEN,* with the exception of certain taxa (including *A. vinelandii*) that possess *vnfEN* homologs of *nifEN* that likely perform a similar biosynthetic function (Boyd, Anbar, et al., 2011; Hamilton et al., 2011).
Paleobiological interest in nitrogenases has primarily centered on the coevolution of nitrogenase metal usage and the geochemical environment, with the possible implication that marine metal availabilities have significantly constrained nitrogenase evolution over geologic time (Anbar & Knoll, 2002; Boyd, Hamilton, et al., 2011; Canfield, Glazer, & Falkowski, 2010; Raymond et al., 2004). Inferences of ancient nitrogenase metal usage have relied on isotopic biosignatures (Stueken et al., 2015) and metal abundances (Anbar & Knoll, 2002) evidenced by the geologic record, as well as on phylogenetic reconstructions of both catalytic and cofactor biosynthesis proteins (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Raymond et al., 2004). High marine Fe concentrations and potential Mo scarcity prior to increased atmospheric oxygenation surrounding the ~2.3-2.5 Ga Great Oxidation Event (Anbar et al., 2007; Lyons, Reinhard, & Planavsky, 2014) has led to the hypothesis that Fe- or V- nitrogenases may have been dominant in early oceans (Anbar & Knoll, 2002; Canfield et al., 2010) and possibly predate Mo-nitrogenases (Raymond et al., 2004). More recent phylogenetic reconstructions have instead suggested that the evolution of Mo-nitrogenases, dated by time-calibrated phylogenies of Nif/Vnf/AnfDREN sequences to ~1.5-2.2 Ga (Boyd, Anbar, et al., 2011), preceded that of V- and Fe-nitrogenases (Boyd, Hamilton, et al., 2011). These phylogenetic inferences are also consistent with the observation that vnf and anf genes are only present in organisms that also harbor nif, and that V-/Fe-nitrogenase assembly relies on nif biosynthetic genes (Hamilton et al., 2011; Joerger & Bishop, 1988; Kennedy & Dean, 1992). However, ~3.2-Ga isotopic signatures of biological nitrogen fixation suggest an earlier origin of nitrogenase (Stueken et al., 2015), and, even though isotopically consistent with Mo-dependent N-fixation, predate age estimates of both Mo-nitrogenase (Boyd, Anbar, et al., 2011) and earliest marine Mo availability (Anbar et al., 2007;
Anbar & Knoll, 2002; Lyons et al., 2014). Thus, the evolutionary trajectory of nitrogenase metal usage — and by extension the link between nitrogenase evolution and marine metal availabilities over geologic time — is not yet known.

Here, we explored the indicators of nitrogenase metal usage history by a combinatorial method relying on ancestral sequence reconstruction, a method by which inferred, historical protein sequence information can be linked to functional molecular properties evidenced by computed structures or laboratory experiments (Aadland, Pugh, & Kolaczkowski, 2019; Benner, Sassi, & Gaucher, 2007; Thornton, 2004). These paleogenetic approaches have been increasingly applied in biogeochemically relevant molecular studies to offer insights into the coevolution of life and Earth (Garcia & Kacar, 2019; Gomez-Fernandez et al., 2018; Kacar, Hanson-Smith, Adam, & Boekelheide, 2017; Shih et al., 2016). We reconstructed the phylogenetic history of Mo-, V-, and Fe-nitrogenases in order to resurrect ancestral nitrogenases in silico, as well as to map the taxonomic distribution of cofactor biosynthetic components considered necessary for Mo-dependence (Curatti et al., 2007; Shah et al., 1994; Shah et al., 1986; St John et al., 1975; Tal et al., 1991). By this combined approach, we find phylogenetic and ancestral sequence features suggestive of Mo-dependence, potentially by an alternate pathway predating the origin of the FeMo-cofactor. We speculate that this unknown and possibly transient pathway may today be present in basal nitrogenase lineages. Integration of protein evolution and paleobiology is a unique melding of disparate data sets and may allow construct-and-build hypotheses that address interactions ranging from the external environment to the cellular environment, and from the cellular environment to the that maintained around the interacting protein. The exchange of materials across these different scales necessitates constraints on the flow and availability of
substrates that make such exchanges possible. It is the specific nature of these constraints and how they may change in response to external perturbations that enable us to develop completely new testable hypothesis that connect geochemical reservoirs with biological metabolisms — hypotheses that cannot be constructed from macroevolutionary or geological frameworks alone.

2 MATERIALS AND METHODS

2.1 Ancestral reconstruction of nitrogenase protein sequences

An initial dataset of extant nitrogenase Nif/Vnf/AnfHDK homologs was constructed by retrieving amino acid sequences from the National Center for Biotechnology Information non-redundant protein database, accessed September 2018 (O’Leary et al., 2016). Potential homologs were identified by BLASTp (Camacho et al., 2009) using query sequences from A. vinelandii (NifH: Avin_01380, NifD: Avin_01390, NifK: Avin_01400) and an Expect value cutoff of <1e-5. The dataset was then manually curated to remove partial and distantly related sequences. Additional nitrogenase sequences were manually retrieved from the Joint Genome Institute Integrated Microbial Genomes and Microbiomes database, accessed September 2018 (Chen et al., 2019). The nitrogenase sequence dataset was finalized to include NifHDK sequences from 256 taxa, AnfHDK sequences from 14 taxa, VnfHDK sequences from 14 taxa, and outgroup light-independent protochlorophyllide oxidoreductase (Bch/ChlLN B) sequences — sharing distant homology with nitrogenases (Boyd, Anbar, et al., 2011; Y. Hu & Ribbe, 2015; Raymond et al., 2004) — from 10 taxa (Appendix S1; additional analyses were performed with an expanded outgroup, Appendix S2). Only one Nif/Anf/VnfHDK sequence set was retained per genus to broaden taxonomic sampling. Equal sequence sampling for Anf and Vnf was made to remove the potential for oversampling bias in ancestral sequence inference. H-, D-, and K-subunit sequences corresponding
to each taxon were manually checked for synteny of their encoding genes. AnfHDK and VnfHDK sequences were identified by the proximity of each gene locus to \textit{anfG} or \textit{vnfG}, which encodes the additional G-subunit present in the VeFe or FeFe protein, respectively, but not present in the MoFe protein (Eady, 1996). Finally, the presence of the cofactor biosynthetic \textit{nifBEN} genes was investigated for all taxa represented in our dataset by BLASTp, as well as by manually inspecting the \textit{nif} genome region.

Reconstruction of ancestral nitrogenase sequences was performed by PhyloBot (Hanson-Smith & Johnson, 2016) (www.phylobot.com), which automates multiple sequence alignment, phylogenetic reconstruction, and ancestral sequence inference methods. The concatenated 294-sequence dataset of Nif/Anf/VnfHDK homologs (including 10 Bch/ChlLNB outgroup sequences) was aligned by MSAProbs v0.9 5r1 (Liu, Schmidt, & Maskell, 2010) and MUSCLE v3.8.31 (Edgar, 2004). Both alignment outputs were then used to perform phylogenetic reconstruction by RAxML v8.1.15 (Stamatakis, 2014) under 6 different combinations of amino acid substitution and rate heterogeneity models. Branch support was evaluated by the approximate likelihood ratio test (aLRT) (Anisimova & Gascuel, 2006), which assesses the gain in overall likelihood against a null hypothesis of branch length = 0. Additional phylogenetic reconstructions with an expanded outgroup were performed outside of Phylobot to resolve root positioning, but were not used in subsequent ancestral sequence inference (Appendix S2). Ancestral sequences were inferred by joint maximum likelihood using CODEML v4.2 (Z. Yang, 2007) at all nodes within the 12 Phylobot-constructed phylogenies (Tree-1 – Tree-12), with gaps inferred by parsimony. To assess ancestral sequence robustness to phylogenetic uncertainty (Hanson-Smith, Kolaczkowski, & Thornton, 2010), ancestors inferred from the top five phylogenies ranked by log likelihood scores
were selected for further analysis (Table 1, Appendix S2). Finally, to evaluate the effects of ambiguously reconstructed sites on subsequent structural analyses, Bayesian sampled ancestors were inferred from the maximum likelihood site posterior probabilities calculated by CODEML (Aadland et al., 2019). 100 random Bayesian sequence were generated for each of five ancestral nodes of interest across the top five phylogenies. Thus, 25 maximum likelihood and 2,500 Bayesian-sampled ancestral sequences were analyzed in total. All maximum likelihood reconstructed trees and ancestral sequences are available for view and download at http://phylobot.com/613282215/.
Table 1. Alignment and evolutionary model parameter combinations for the top five phylogenies, ranked by log likelihood scores.

| Phylogeny | Alignment method | Evolutionary model | Log likelihood | Maximum likelihood ancestors |
|-----------|------------------|--------------------|----------------|-----------------------------|
| Tree-1    | MSAProbs         | CAT + LG           | -300069.08     | AncA-1 – AncE-1             |
| Tree-2    | MSAProbs         | CAT + WAG          | -303296.08     | AncA-2 – AncE-2             |
| Tree-3    | MSAProbs         | Γ + LG             | -303951.21     | AncA-3 – AncE-3             |
| Tree-4    | MUSCLE           | CAT + LG           | -304457.52     | AncA-4 – AncE-4             |
| Tree-5    | MSAProbs         | Γ + WAG            | -305229.49     | AncA-5 – AncE-5             |

References: (Le & Gascuel, 2008; Quang, Gascuel, & Lartillot, 2008; Whelan & Goldman, 2001; Z. Yang, 1993)
2.2 Structural homology modeling of extant and ancestral nitrogenase D-subunits

Structural homology modeling of 33 extant and ancestral (25 maximum likelihood and 2,500 Bayesian-sampled) nitrogenase D-subunit proteins was performed by Modeller v9.2 (Sali & Blundell, 1993). Extant nitrogenase sequences, broadly sampled from the reconstructed nitrogenase phylogeny, were modeled to provide comparisons with ancestral models. D-subunit sequences from extant and ancestral nitrogenases were aligned to 38 NifD and 2 VnfD structural templates retrieved from the Protein Data Bank (Berman et al., 2000), accessed November 2018 (Appendix S5; published AnfD models not available at time of analysis). Information from all 40 templates was used to model each structure. All models were generated by specifying the inclusion of the FeMo-cofactor of the 3U7Q NifD structure (Spatzal et al., 2011), selected as the highest resolution Mo-nitrogenase template. To assess the effect of the template cofactor type on the generated structure, additional models were constructed by specifying the inclusion of the FeV-cofactor of the 56NY VnfD template (D. Sippel & Einsle, 2017) (Appendix S5). 100 modeling replicates were performed per sequence and assessed by averaging over the scaled Modeller objective function, Discrete Optimized Protein Energy, and high resolution Discrete Optimized Protein Energy scores, as previously described (Aadland et al., 2019). The ten best modeling replicates per extant sequence, ten best replicates per maximum likelihood ancestral sequence, and the single best replicate per Bayesian-sampled variant sequence were selected for further analysis, totaling 3,080 models with the FeMo-cofactor specified.

2.3 Active-site pocket volume calculation of extant and ancestral D-subunit models

Volumes of the modeled ancestral and extant D-subunit active-site cofactor pockets were calculated by POVME v2.0 (Durrant, Votapka, Sorensen, & Amaro, 2014). Spatial coordinates
and the inclusion region for volume calculation were specified manually. Pocket volumes were calculated with a grid spacing of 0.5 Å and a 1.09 Å distance cutoff from any receptor atom’s van der Waals radius. Volume outside of the modeled convex hull of the cofactor pocket as well as noncontiguous volume were removed. Statistical analysis of ancestral and extant pocket volume data was performed in R (R Core Team, 2014).

3 RESULTS

3.1 V- and Fe-nitrogenases diversified after Mo-nitrogenases

We reconstructed the phylogenetic history of Mo-, V-, and Fe-nitrogenases and to infer ancestral nitrogenase sequences and associated indicators of nitrogenase metal dependence. Nif/Anf/VnfHDK protein homologs curated from the National Center for Biotechnology Information and Joint Genome Institute databases represent 20 bacterial and archaeal phyla, 11 of which are known from experimental investigations to include diazotrophic taxa (Dos Santos et al., 2012; Ormeño-Orrillo, Hungria, & Martinez-Romero, 2013) (Appendix S1). The five most represented phyla in our dataset — Bacteroidetes, Cyanobacteria, Firmicutes, Proteobacteria, and Euryarchaeota — encompass ~80% of the curated sequences. Our dataset also presents genomic evidence of nitrogen fixation within the Acidobacteria, Actinobacteria, Aquificae, Chlorobi, Chloroflexi, Chrysiogenetes, Deferrribacteres, Elusimicrobia, Fusobacteria, Lentisphaerae, Candidatus Margulisbacteria, Nitrospirae, Planctomycetes, Spirochaetes, and Verrucomicrobia.

Our maximum likelihood nitrogenase phylogeny segregates Nif/Anf/VnfHDK sequences into two major lineages similarly observed in previous studies (Boyd, Anbar, et al., 2011; Raymond et al., 2004) (Tree-1; Figure 2): the first comprises Nif-I and Nif-II Mo-nitrogenases (aerobic/facultative
bacteria and anaerobic/facultative bacteria/archaea, respectively; highlighted in blue) (Raymond et al., 2004), and the second comprises V- and Fe-only-nitrogenases (Vnf and Anf, respectively; highlighted in red) as well as three clades of “uncharacterized” nitrogenases that lack extensive experimental characterization with regard to metal dependence (Mb-Mc, highlighted in purple; F-Mc, highlighted in green; Clfx, highlighted in yellow) (Boyd, Hamilton, et al., 2011; Dekas, Poretsky, & Orphan, 2009; Dos Santos et al., 2012; Mehta & Baross, 2006). We additionally analyzed four alternate phylogenies (ranked by log likelihood scores), reconstructed by varying alignment methods, amino acid substitution models, and rate heterogeneity models (Tree-2–Tree-5; Appendix S2). The aforementioned major nitrogenase clades are present across all alternate phylogenetic topologies. Branch ordering of these major clades is similarly consistent, with the exception of root position: in Tree-3 and Tree-5, the root is placed between Nif-I and Nif-II/uncharacterized/Anf/Vnf, rather than between Nif-I/Nif-II and uncharacterized/Anf/Vnf as in Tree-1, Tree-2, and Tree-4. Nevertheless, additional phylogenetic analyses incorporating an expanded outgroup provide stronger support for the root position of Tree-1, Tree-2, and Tree-4 (Appendix S2).

In all analyzed nitrogenase phylogenies, Vnf and Anf sequences (highlighted in red) form reciprocally monophyletic clades that branch immediately distal to Mb-Mc nitrogenases (highlighted in purple) (Figure 2, Appendix S2). The reciprocal monophyly of Vnf and Anf, as well as the clustering of Mb-Mc, Vnf, and Anf, is well-supported across all phylogenetic topologies (aLRT > 10^6). The Mb-Mc clade is composed of archaeal hydrogenotrophic methanogens within classes Methanobacteria and Methanococci. Because all Mb-Mc taxa possess nifBEN genes demonstrated to be necessary for the synthesis of the FeMo-cofactor (Curatti et al.,
2007; Dos Santos et al., 2012; Shah et al., 1994; Shah et al., 1986; St John et al., 1975; Tal et al., 1991), it is likely that Mb-Mc nitrogenases are Mo-dependent (Boyd, Hamilton, et al., 2011). Thus, the phylogenetic positioning of Vnf and Anf is consistent with previous suggestions that V- and Fe-nitrogenases diversified after Mo-nitrogenases (Boyd, Hamilton, et al., 2011).
Figure 2. Maximum likelihood phylogeny of concatenated Nif/Anf/VnfHDK nitrogenase and Bch/ChlLN1 outgroup protein sequences (Tree-1; see Table 1). Ancestral nodes analyzed in this study are labeled AncA–AncE. Known active-site cofactor metal content is listed on the right. Branch support is derived from the approximate likelihood ratio test (aLRT). Branch length scale is in units of amino acid substitutions per site. Outgroup branch break used to conserve space; true branch length = 5.578 substitutions per site. Phylogeny coloring is as follows: Clfx, yellow; F-Mc, green; Mb-Mc, purple; Anf/Vnf, red; Nif-I/-II, blue.

3.2 Basal uncharacterized nitrogenases lack associated genes necessary for FeMo-cofactor synthesis

In addition to investigating the phylogenetic relationships between Mo-, V-, and Fe-nitrogenase isozymes, we mapped the presence of the biosynthetic nifB, nifE, and nifN genes — necessary for FeMo-cofactor assembly (Curatti et al., 2007; Dos Santos et al., 2012; Shah et al., 1994; Shah et al., 1986; St John et al., 1975; Tal et al., 1991) — among taxa represented in our dataset. All analyzed taxa possess the full complement of nifBEN biosynthetic genes, with the exception of two uncharacterized clades: Clfx (highlighted in yellow) and F-Mc (highlighted in green) (Figure 2). Within the lineage containing Vnf and Anf nitrogenases, Clfx and F-Mc clades are most basal. These branching positions of Clfx and F-Mc clades within the Vnf/Anf lineage are consistently observed across all phylogenetic topologies (Trees-1–5; Appendix S2) and are well-supported (aLRT > 10^3 for Clfx, aLRT > 10^{18} for F-Mc). In Tree-1-, 2-, and -4, as well as in trees reconstructed with an expanded outgroup (Appendix S2), Clfx and F-Mc clades also branch immediately distal to the root.
Both basal Clfx and F-Mc nitrogenases are mostly derived from thermophilic hosts, and lack one or more associated nifEN cofactor biosynthesis genes. Clfx nitrogenases, constituting the most basal uncharacterized clade within the uncharacterized/Vnf/Anf lineage, are in the present dataset derived from three mesophilic or thermophilic Chloroflexi species (Figure 2, highlighted in yellow). The nif clusters of Clfx taxa lack both nifE and nifN genes, and the typically continuous nifHDK genes observed in other taxa are instead interrupted by nifB (arranged nifHBDK)(Setubal et al., 2009). F-Mc nitrogenases branch immediately distal to the Clfx clade and represent eight thermophilic Firmicutes and archaeal methanogen (class Methanococci) taxa (Figure 2, highlighted in green). F-Mc species possess biosynthetic nifB and nifE genes, but not nifN, with the exception of Methanothermococcus thermolithotrophicus that retains nifN. A previous study found that sequence features and modeled structural features of F-Mc nitrogenases closely resemble those of Mo-nitrogenases (McGlynn, Boyd, Peters, & Orphan, 2012). However, the absence of nifN in most F-Mc taxa suggests that these strains are not capable of synthesizing the FeMo-cofactor. Though the lack of nifE and/or nifN genes in Clfx and F-Mc taxa might additionally suggest that such strains cannot express functional nitrogenases, some have been experimentally observed to fix nitrogen: the Clfx species Oscillochloris trichoides (lacking nifEN) (Keppen, Lebedeva, Troshina, & Rodionov, 1989; Kuznetsov et al., 2011) and the F-Mc species Methanocaldococcus sp. FS406-22 (lacking nifN) (Keppen et al., 1989; Kuznetsov et al., 2011; Mehta & Baross, 2006), in addition to uncharacterized anaerobic methane-oxidizing archaea not included in the present study (Dekas et al., 2009). The ability of early-branching Clfx and F-Mc nitrogenases to fix nitrogen in the absence of nifEN genes may indicate an atypical pathway for cofactor assembly and incorporation not used for extant Mo-, V-, and Fe-nitrogenases.
3.3 High statistical support for ancestral nitrogenase active-site residues

We inferred ancestral sequences for each of the H-, D-, and K-subunits that constitute the nitrogenase enzyme complex (Figure 1) across five phylogenetic topologies (Tree-1–5; Appendix S2). Ancestral nitrogenase sequences were inferred for five well-supported internal nodes along a phylogenetic transect between Mo- (highlighted in blue) and V-/Fe-nitrogenases (highlighted in red) (Figure 2). The five targeted nodes are: AncA (ancestral to Anf and Vnf), AncB (ancestral to AncA and Mb-Mc), AncC (ancestral to AncB and F-Mc), AncD (ancestral to AncC and Clfx), and AncE (ancestral to Nif-I and Nif-II). Thus, AncA–D are nested, whereas AncE lies along a divergent lineage toward Nif-I and Nif-II Mo-nitrogenases. For further analyses, we selected the maximum likelihood ancestral sequence per target node from each of the five phylogenies (Tree-1–5), totaling 25 sequences. Due to differences in root position, identical AncE nodes were not present across all topologies and analogous nodes were instead selected (Appendix S2). Ancestral sequences are hereafter labeled with the tree likelihood rank from which they were inferred (e.g., AncA from Tree-1 is labeled AncA-1). All tree and ancestral sequence information can be found at http://phylobot.com/613282215/.

Mean site posterior probabilities for ancestral nitrogenase HDK sequences across all phylogenies range between ~0.83 and 0.91, and for the highest-likelihood phylogeny (Tree-1), between ~0.84 and 0.90 (Appendix S3). Ancestral sequence support generally decreases with increasing phylogenetic node age. For example, within the uncharacterized/V-Fe-nitrogenase lineage, AncA-1 has the highest mean posterior probability (0.90 ± 0.18) and AncD-1 has the lowest mean posterior probability (0.84 ± 0.22). Mean ancestral sequence probability for each node also does not deviate by more than ~0.02 across each of the five phylogenetic topologies (Tree-1–5;
Appendix S3). These observations suggest that sequence support for ancestral nitrogenases is more sensitive to ancestral node position than to topological differences between the analyzed trees.

In addition to surveying total ancestral HDK sequence support, we analyzed support for 30 active-site residues, defined as those residing within 5 Å of any atom in either the FeMo-cofactor of the *A. vinelandii* NifD protein (PDB 3U7Q (Spatzal et al., 2011)) or the FeV-cofactor of the *A. vinelandii* VnfD protein (PDB 5N6Y (D. Sippel & Einsle, 2017)) (Figure 3). These active-site residues are not contiguous but are instead scattered throughout the D-subunit sequence. Mean posterior probabilities of ancestral active-site residues, which range between 0.92 to 0.98 across all phylogenies, are consistently greater than those of entire reconstructed nitrogenase HDK sequences (0.83–0.91) (Appendix S3). Of the 30 active-site residues, only five sites have, in one or more ancestral sequences, plausible alternative reconstructions with posterior probabilities > 0.30: sites 59, 69, 358, 360, 425, 441 (site numbering both here and hereafter based on *A. vinelandii* NifD). No ancestral sequences have more than three active-site residues with such plausible alternative reconstructions. Ten active-site residues are conserved across all analyzed extant nitrogenases: Val-70, Gln-191, His-195, Cys-275, Arg-277, Ser-278, Gly-356, Phe-381, Gly-424, His-442. These conserved residues are thus reconstructed in all ancestral nitrogenases unambiguously (site posterior probability = 1.00). Statistical support for ancestral active-site residues (greater than 0.92) underpins subsequent analyses of ancestral active-site properties that may inform inferences of nitrogenase metal dependence.
**Figure 3.** Active-site protein environment of representative ancestral and extant nitrogenases. All residues located within 5 Å of any atom in either the FeMo-cofactor of the *A. vinelandii* NifD protein (PDB 3U7Q (Spatzal et al., 2011)) or the FeV-cofactor of the *A. vinelandii* VnfD protein (5N6Y (D. Sippel & Einsle, 2017)). Residue numbering from aligned *A. vinelandii* NifD. Cys-275 and His-442 residues that coordinate the cofactor are indicated by red arrows. Phylogeny coloring is as follows: Clfx, yellow; F-Mc, green; Mb-Mc, purple; Anf/Vnf, red; Nif-I/-II, blue.

### 3.4 Active-site structural features are uninformative for inferring ancestral metal dependence

To investigate metal-specific features of ancestral nitrogenase structures, we generated homology models of both extant and ancestral nitrogenase D-subunits that house the active site (**Figure 4a**). First, we modeled 33 broadly sampled extant nitrogenase NifD, VnfD, and AnfD sequences to benchmark classifications of ancestral nitrogenase models. Second, we calculated structural models of 25 nitrogenase ancestors inferred by maximum likelihood and of 2,500 ancestors inferred by random Bayesian sampling of maximum likelihood site posterior probabilities (100 Bayesian samples per maximum likelihood ancestor). We generated ten model replicates per extant sequence and maximum likelihood sequence, and one model per Bayesian-sampled sequence. All structures were modeled with the FeMo-cofactor included (additional modeling runs were...
executed with the FeV-cofactor included; Appendix S5). In total, 3,080 models were generated with the FeMo-cofactor.

Figure 4. Structural and active-site pocket modelling of ancestral nitrogenases (a) Modeled D-subunit protein structures of ancestral nitrogenases inferred from the highest-likelihood phylogeny (Tree-1; Figure 2) aligned to an *A. vinelandii* Nif structural template (PDB 3U7Q (Spatzal et al., 2011)). (b) Example of a modeled active-site pocket for ancestral nitrogenase AncA-1. The 0.5-Å-resolution point field generated for pocket volume calculation is shown in pink.

For each of the 3,080 extant and ancestral D-subunit nitrogenase models, we calculated the volume of the active-site pocket (Figure 4b), a parameter previously used to classify the metal dependence of extant uncharacterized nitrogenases (McGlynn et al., 2012). These pocket volume values are plotted in Figure 5. Among modeled extant nitrogenases, mean pocket volumes are $1175.12 \pm 51.93$ Å$^3$ for Mo-nitrogenases, $1121.86 \pm 36.36$ Å$^3$ for V-nitrogenases, and $963.39 \pm 75.80$ Å$^3$ for Fe-nitrogenases.
Figure 5. Extant and ancestral nitrogenase active-site pocket volumes. Pocket volumes calculated for ancestral and representative extant nitrogenase D-subunit structures modeled with the FeMo-cofactor. Each ancestral plot contains 110 volume calculations (ten model replicates per maximum likelihood sequence plus one model for each of 100 Bayesian-sampled sequences) and each extant plot contains 10 volume calculations (10 model replicates per extant sequence). Median values are indicated by bars, mean values by points, the range (excluding outliers) by whiskers, and outliers by crosses. Phylogeny coloring is as follows: Clfx, yellow; F-Mc, green; Mb-Mc, purple; Anf/Vnf, red; Nif-I/-II, blue.

We observe less difference between mean pocket volumes of extant V-nitrogenases (1121.86 ± 36.36 Å³) and Nif-II Mo-nitrogenases (1141.13 ± 46.30 Å³) than between Nif-I (1209.11 ± 30.79 Å³) and Nif-II Mo-nitrogenases. A statistical nonparametric test of volume median differences also suggests greater similarity between V- and Nif-II Mo-nitrogenases than between Nif-I and Nif-II nitrogenases (Appendix S5). All V-nitrogenase and Nif-II Mo-nitrogenase volume values range
between 1002.13 and 1220.63 Å³, which problematically overlap with the volume ranges of AncB (917.38–1256.25 Å³), AncC (933.75–1269.50 Å³), AncD (968.50–1272.75 Å³), and AncE (966.75–1225.25 Å³) ancestral models, as well as ranges for uncharacterized Clfx (988.25–1218.13 Å³) and F-Mc (908.00–1277.75 Å³) nitrogenases. The volume range of AncA models (919.00-1191.13 Å³) lies between the ranges of both V- (1020.75–1197.63 Å³) and Fe-nitrogenases (821.38–1177.00 Å³). These same overall patterns are observed when comparing only maximum likelihood ancestral models or Bayesian-sampled ancestral models, as well as alternate modeling runs with the FeV-cofactor (Appendix S5). Greater similarities between V- and Nif-II Mo-nitrogenases than between Nif-I and Nif-II Mo-nitrogenases suggests that, contrary to previous analyses (McGlynn et al., 2012), modeled pocket volume may be uninformative for inferring ancestral metal dependence. At the very least, the overlap in volume ranges between ancestors and extant isozymes of varying metal dependence in our analyses preclude the unambiguous classification of ancestral metal dependence by these structural features.

3.5 Oldest ancestral nitrogenase active-site sequences resemble those of Mo-nitrogenases

We analyzed sequence features of both ancestral and extant nitrogenases to identify those correlated with metal dependence. In particular, we focused on nitrogenase active-site sequences for three reasons: (1) active-site residues are known to affect catalytic efficiency and substrate specificity (Brigle et al., 1987; Christiansen, Cash, Seefeldt, & Dean, 2000; Fixen et al., 2016; Kim, Newton, & Dean, 1995; Sarma et al., 2010; Z. Y. Yang, Moure, Dean, & Seefeldt, 2012) and thus may be tuned to nitrogenase metal dependence (2) active-site sequence features have previously been used to classify the metal dependence of extant uncharacterized nitrogenases
an ancestral active-site residues are reconstructed here with high statistical support as compared to the total HDK sequence (see Section 3.3).

We first assessed the sensitivity of ancestral sequence variation to phylogenetic uncertainty and ancestral statistical support. Overall, mean identities for ancestral sequences compared across different nodes range from ~55 to 90%. Ancestral sequences inferred from the same node across alternate phylogenies (Tree-1–5) have relatively high mean identities, ranging from ~93 to 95% across the total HDK sequence and from ~96 to 100% within the active site. These high mean identities suggest that topological differences among the alternate phylogenies used for ancestral sequence inference do not contribute to a high degree of ancestral sequence variation. Identities among sequences inferred from the same node also do not appear to be correlated with statistical support. For example, though full AncA HDK sequences are reconstructed with the highest mean statistical support (~0.89–0.91), they exhibit neither the lowest nor the highest mean identities as compared with sequences inferred from other nodes (Appendix S3).

We next identified specific residues within the nitrogenase active site that are unique to particular isozymes of known metal dependence in order to survey their occurrence in ancestral sequences. We found that three active-site residues are unique to Mo-nitrogenases, six are unique to V-nitrogenases, five are unique to Fe-nitrogenases, and six are unique to V- and Fe-nitrogenases (Appendix S4). Surprisingly, most nitrogenase ancestors exhibit comparable numbers of residues unique to either V-/Fe-nitrogenases or Mo-nitrogenases, and thus their occurrence does not appear informative for inferring ancestral metal dependence (e.g., ancestral AncC sequences contain two residues unique to V-/Fe-nitrogenases and two residues unique to Mo-nitrogenases). An exception
is AncA sequences, which contain a preponderance of active-site residues that are unique to extant V- and Fe-nitrogenases. One of the residues unique to V-nitrogenases, Thr-355, has recently been suggested to interact directly with a proposed carbonate ligand only present in the FeV-cofactor (D. Sippel & Einsle, 2017). This carbonate ligand lies within a protein loop that also contains Pro-358, unique to V-nitrogenases, and Leu-360, unique to V- and Fe-nitrogenases. These Thr-355, Pro-358, and Leu-360 residues are observed across all AncA sequences, reflecting specific sequence resemblances that may be associated with V-dependence.

In addition to analyzing specific active-site residues, we compared the total active-site sequence composition of 25 ancestral nitrogenases (inferred across Tree-1–5) and all 284 extant nitrogenase sequences used for phylogenetic reconstruction. Our analysis shows clear distinctions between the active-site compositions of extant Mo- versus V-/Fe-nitrogenases (Figure 6a). Specifically, the mean identity between Vnf/Anf and Nif-I/Nif-II is ~50%, as compared with the mean identity between Vnf and Anf (~71%) and between Nif-I and Nif-II (~77%). The difference between Vnf/Anf and Nif-I/Nif-II sequences is not seen as distinctly when comparing total HDK sequences (Figure 6b), suggesting that the active-site sequence differences in particular may be better indicators of metal dependence than whole sequence differences due to greater catalytic tuning toward the cofactor.
Figure 6. Active-site and full HDK sequence comparisons between extant and ancestral nitrogenases. (a) Active-site sequence identities of ancestral and extant nitrogenases. Active-site residues include 30 amino acids positioned within 5 Å of the active-site cofactor. (b) Total HDK sequence identities of ancestral and extant nitrogenases. Percentage identity values have been averaged within each field of comparison. All 25 maximum likelihood ancestors and 284 extant sequences included in this study were used for sequence identity calculation. Phylogeny coloring is as follows: Clfx, yellow; F-Mc, green; Mb-Mc, purple; Anf/Vnf, red; Nif-I/-II, blue.

Because we observed active-site sequence distinctions between extant Vnf/Anf and Nif-I/Nif-II nitrogenases, we compared active-site sequences of ancestral versus extant nitrogenases to provide clues regarding ancestral metal dependence. Nearly all ancestral nitrogenases, including those inferred for the oldest ancestral nodes, share greater active-site identity with Mo-nitrogenases than with V-/Fe-only-nitrogenases (Figure 6a). Mean active-site sequence identities between AncB–AncE and Anf/Vnf nitrogenases range between ~50 and 63%, whereas those between AncB–AncE and Nif-I/Nif-II nitrogenases range between ~69 and 87%. An exception is AncA (ancestral to Vnf
and Anf), which has higher mean identity to Anf/Vnf nitrogenases (~85%) than to Nif-I/Nif-II nitrogenases (~51%). Because active-site sequence identity can reliably differentiate extant Mo- from V-/Fe-nitrogenases, the resemblance of most ancestral active sites to those of Mo- nitrogenases is suggestive of Mo-dependence.

4 DISCUSSION

Nitrogenase mediates the reduction of N₂ to NH₃, a key step in nitrogen fixation (Anbar & Knoll, 2002; Canfield et al., 2010; Falkowski, 1997). The metal dependence of nitrogenase, which impacts both catalytic properties (Eady, 1996; Harris, Yang, et al., 2018; Y. Hu et al., 2011; Lee et al., 2018; Rebelein et al., 2018; Zheng et al., 2018) and ecological distribution (McRose, Zhang, Kraepiel, & Morel, 2017; Zhang et al., 2016), suggests a potential role for marine geochemical constraints on its evolution (Anbar & Knoll, 2002; Boyd, Hamilton, et al., 2011; Canfield et al., 2010; Raymond et al., 2004). Thus, understanding ancestral nitrogenase metal dependence can help resolve the early history of biological nitrogen fixation, and, in a broader sense, the impact that ancient metal availabilities have had on the evolution of biologically essential metabolisms over Earth history (Anbar & Knoll, 2002; Moore, Jelen, Giovannelli, Raanan, & Falkowski, 2017).

Previous phylogenetic work has established that Mo-, V-, and Fe-nitrogenases, though genetically distinct, are evolutionarily homologous (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Raymond et al., 2004). Most recent phylogenetic analyses also indicate that V- and Fe-nitrogenases are derived from Mo-nitrogenases, the latter having originated following the gene duplication event that produced nifE and nifN (Boyd, Anbar, et al., 2011). However, the precise trajectory of metal-binding evolution in the nitrogenase family is not completely known, and
discrepancies between current phylogenetics-based models and the geochemical record of nitrogen fixation remain (Stueken et al., 2015).

We used both phylogenetic reconstruction and ancestral sequence inference to explore these outstanding questions of early nitrogenase evolution and metal dependence. Though the tree reconstructions presented here are largely in congruence with previous phylogenetic analyses, certain topological differences, particularly with regard to basal uncharacterized nitrogenases lacking associated nifE and/or nifN genes, suggest important deviations from previous narratives of early metal dependence (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Boyd & Peters, 2013; Raymond et al., 2004). The reconstruction of ancestral nitrogenase sequences in silico provides the means to directly infer ancient metal-binding properties from molecular information.

4.1 Inferred metal dependence of ancestral nitrogenases

The active-site protein environment is known, in addition to the metal content of the cofactor, to contribute to the variable catalytic properties of different nitrogenase isozymes (Brigle et al., 1987; Christiansen et al., 2000; Fixen et al., 2016; Kim et al., 1995; Sarma et al., 2010; Z. Y. Yang et al., 2012). We therefore explored active-site features of ancestral nitrogenases for correlations with metal dependence.

We find that modeled active-site structural features are not informative for the inference of ancestral metal dependence. Pocket volumes of modeled extant nitrogenases do not appear to be strongly correlated with metal cofactor content. Problematically, the volume ranges of oldest ancestors (as well as those of uncharacterized nitrogenases) overlap with both V- and Mo-
nitrogenases (Figure 5). We acknowledge that these homology modeling results may not precisely
reflect true biological differences (for which a comprehensive analysis is not yet possible due the
limited availability of V- and Fe-nitrogenase structures). This is illustrated by the difference
between pocket volumes of published structures and those of homology models (e.g., A. vinelandii
PDB 3U7Q NifD structure pocket volume ≈ 994.750 and mean A. vinelandii NifD modeled
structure pocket volume ≈ 1186.412). However, it is not surprising that active-site pocket volume
does not predict metal dependence, given the probable similar sizes and structures of the FeMo,
FeV-, and FeFe-cofactors (Eady, 1996; Krahn et al., 2002; Daniel Sippel et al., 2018; Spatzal et
al., 2011). These findings contrast with a previous study by McGlynn and coworkers (2012) in
which pocket volume comparisons were used to classify the Mo-dependence of uncharacterized
nitrogenases. In this previous study, the volume means for Mo-nitrogenases were sufficiently
distinct from V- and Fe-nitrogenases as to provide unambiguous classification of Mo-dependence.
Our analyses differ from this previous study in several ways: (1) we incorporated V-nitrogenase
structural templates (D. Sippel & Einsle, 2017; Daniel Sippel et al., 2018) for homology modeling,
not available at the time of the previous study (2) we modeled 20 extant sequences of known metal
dependence rather than 12 (3) we used the same explicit parameters for both homology modeling
and pocket volume calculation across all sequences (4) we included modeling replicates for pocket
volume calculation. Thus, an expanded modeling analysis appears to reduce the efficacy of the
pocket volume parameter for inference of ancestral (and uncharacterized) nitrogenase metal
dependence.

Unlike structure-based analyses, we find that active-site sequence features reliably differentiate
Mo-nitrogenases and V-/Fe-nitrogenases. Regarding AncA (ancestral to V- and Fe-nitrogenases),
we identified specific active-site residues that have previously been suggested to interact with a proposed carbonate ligand unique to the FeV-cofactor. These residues form a $^{355}$TGGPRL$^{360}$ loop conserved only among V-nitrogenases and homologous to $^{355}$IGGLRP$^{360}$ in *A. vinelandii* NifD (D. Sippel & Einsle, 2017). This substitution of Thr-355 for Ile-355, as well as the exchange of Leu and Pro positions may permit the inclusion of the FeV-cofactor carbonate ligand by VnfD that is not possible by the NifD protein (D. Sippel & Einsle, 2017). Thr-355 and Pro-358 are unique to V-nitrogenases, and Leu-360 is unique to V- and Fe-nitrogenases. All AncA sequences conserve the $^{355}$TGGPRL$^{360}$ residue loop capable of accommodating the FeV-cofactor. Furthermore, AncA sequences generally exhibit greater numbers of residues unique to V-nitrogenases than those unique to Fe-nitrogenases, and the mean identity of AncA active-site sequences is highest for V-nitrogenases (Figure 6a). Together, these observations suggest that AncA is V-dependent.

Our comparisons of ancestral and extant sequence features indicate that the active-sites of oldest nitrogenase ancestors (AncB–AncE) resemble those of extant Mo-nitrogenases more than V- or Fe-nitrogenases (Figure 6a). This observation is particularly significant given that these same patterns are not observed across the total HDK sequence (Figure 6b). Specifically, this discrepancy supports the notion that the nitrogenase active-site has been tuned to the catalytic properties of each metal cofactor over its evolutionary history (Harris, Yang, et al., 2018), and that this tuning has manifested in active-site sequence differences that stand apart from baseline phylogenetic distance. Though we are not able to identify specific residues that may functionally relate to metal dependence (as with AncA), the resemblance of the early ancestral nitrogenase active site to those of Mo-nitrogenases is highly suggestive of ancient Mo-dependence.
4.2 A proposed model for the evolution of nitrogenase metal dependence over geologic time

Despite the lack of information provided by structural analyses, the active-site sequence features of oldest ancestral nitrogenases (i.e., AncB–E) support the inference of early Mo-dependence. The observation that ancestral AncC and AncD active sites in particular most resemble those of extant Mo-nitrogenases is at odds with the phylogenetic distribution of nifE and nifN genes, which suggest that early-branching uncharacterized Clfx and F-Mc taxa (for which AncC and AncD are ancestral) are not capable of assembling the FeMo-cofactor. The placement of Clfx and F-Mc clades in our analyses differs from previous phylogenetic reconstructions. For example, the phylogenetic tree presented by Boyd and coworkers nests Clfx and F-Mc clades within Mo-nitrogenases, which notably branch more recently than V-, Fe-, and Mb-Mc Mo-nitrogenases (Boyd, Hamilton, et al., 2011). A subsequently published topology is more similar to the tree presented here, though lacking in Clfx sequences (Boyd, Costas, Hamilton, Mus, & Peters, 2015; Boyd & Peters, 2013). It is possible that the larger sequence dataset used here has refined the placement of these uncharacterized clades, which is supported by our analyses with an expanded outgroup that maintains the positions of Clfx and F-Mc sequences (Appendix S2). Given the well-supported placement of these uncharacterized clades in our reconstruction, we find that the presence of nifE and nifN genes decreases stepwise with divergence age within the uncharacterized/V-/Fe-nitrogenase lineage: Mb-Mc taxa, most recently branched, have both nifEN, most F-Mc taxa only have nifE, and Clfx taxa, earliest branched, have neither. One may thus parsimoniously conclude that uncharacterized nitrogenase AncC–D ancestors similarly lacked the genetic requirements for FeMo-cofactor synthesis.
At least two possible models might resolve this discrepancy: (1) The last common nitrogenase ancestor represented in our tree was in fact hosted by an organism that possessed \textit{nifEN} genes and was able to synthesize the FeMo-cofactor. However, this capability was lost in both Clfx and F-Mc taxa, but retained for all ancestors (AncA–E) as well as for Nif-I, Nif-II, and Mb-Mc taxa. The resemblance of oldest ancestral active sites (AncB–E) to those of Mo-nitrogenases indicate ancient FeMo-cofactor dependence, and the same resemblance of Clfx and F-Mc nitrogenases to Mo-nitrogenases may be inherited from these ancestors. (2) The last common nitrogenase ancestor was hosted by an organism that did not possess \textit{nifEN} genes and was incapable of synthesizing the FeMo-cofactor. The presence of \textit{nifEN} genes increased stepwise through the uncharacterized/V- /Fe-nitrogenase lineage, until the full FeMo-cofactor pathway was completed for AncB nitrogenase at the earliest, but at least in Mb-Mc nitrogenases (\textit{nifEN} genes may have been transferred between ancestors of Mb-Mc, Nif-I, and Nif-II taxa). However, the resemblance of ancestral nitrogenase sequences to those of extant Mo-nitrogenases may evidence a currently unknown, alternative pathway for Mo-usage still present in extant Clfx and F-Mc taxa. This alternative, and possibly unrefined pathway for Mo-usage, may represent a transition state between ancient Mo-independence and full FeMo-cofactor usage.

We prefer the second model for several reasons:

(1) It is challenging to envision a scenario in which the FeMo-cofactor biosynthetic pathway would be lost in Clfx and F-Mc nitrogenases that appear to otherwise be capable of nitrogen fixation (Dekas et al., 2009; Keppen et al., 1989; Kuznetsov et al., 2011; Mehta & Baross, 2006). Mo-nitrogenases are far more efficient at reducing nitrogen than other isozymes (Eady, 1996; Harris
et al., 2019; Harris, Yang, et al., 2018), and the majority of all extant nitrogenases are Mo-dependent across both anoxic and oxic environments (Boyd et al., 2015; Mus, Colman, Peters, & Boyd, 2019; Raymond et al., 2004). Even those organisms that have additional V- or Fe-nitrogenases still retain and preferentially express Mo-nitrogenases (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Dos Santos et al., 2012; Hamilton et al., 2011; Raymond et al., 2004). Aside from Clfx and F-Mc clades, no other nitrogenases are known to lack associated nifEN genes.

(2) It has previously been proposed that early nitrogenases may have been capable of reducing nitrogen prior to the origin of nifEN, and thus of the FeMo-cofactor (Boyd, Hamilton, et al., 2011; Boyd & Peters, 2013; Mus et al., 2019; Soboh, Boyd, Zhao, Peters, & Rubio, 2010). Prior to the evolution of Mo-usage, an ancient Mo-independent nitrogenase may have been capable of — perhaps inefficiently — reducing nitrogen by a cofactor resembling the Fe-S-C cluster assembled by NifB, which constitutes the biosynthetic precursor to the FeMo-cofactor (Boyd & Peters, 2013; Mus et al., 2019; Soboh et al., 2010). Though our sequence analyses cannot assess ancestral nitrogenase dependence for a NifB-cofactor, it is likely that the NifB-cofactor resembles the structure and composition of the FeFe-cofactor, excepting homocitrate (Harris, Lukoyanov, et al., 2018). The greater similarity of AncC–D active sites to those of Mo-nitrogenases than Fe-nitrogenases likely suggests ancestral dependence on a cofactor incorporating Mo rather than only Fe. It is possible that, lacking NifEN, an alternative pathway for Mo-usage may have acted as a transition state between Mo-independence and full FeMo-cofactor usage. It is thus reasonable to speculate that this transition state of Mo-usage may be exhibited by AncC–D ancestors for which the lack of one or both nifEN genes might be parsimoniously inferred.
(3) Given greater catalytic efficiency of Mo-nitrogenases, even an unrefined pathway for Mo-usage may have been of selective advantage if Mo was transiently available prior to ~2.3–2.5 Ga Earth surface oxygenation (Anbar et al., 2007; Anbar & Knoll, 2002; Canfield et al., 2010; Lyons et al., 2014; Raymond et al., 2004). An early generalist behavior with regard to metal usage has previously been proposed (Raymond et al., 2004); such limited and possibly non-specific Mo-usage early in nitrogenase history would have provided an evolutionary pathway for increased Mo-dependence via the addition of NifEN biosynthetic components.

(4) ~3.2-Ga nitrogen isotopic signatures have been interpreted to reflect ancient Mo-dependent nitrogenase activity, due to similarities in isotopic fractionation by extant Mo-nitrogenases (Stueken et al., 2015). However, this interpretation conflicts with age estimates of both nifEN (and the FeMo-cofactor) (Boyd, Anbar, et al., 2011) and of earliest marine Mo availability (Anbar et al., 2007; Lyons et al., 2014). It is possible that alternative Mo-usage in early nitrogenases may have resulted in similar isotopic fractionations. Such possibilities may resolve this discrepancy and remain to be tested.

The preferred model of an ancestral, alternative pathway for Mo-usage, mapped to our phylogenetic reconstruction and inferred ancestors, is illustrated in Figure 7. In the first stage, represented by AncD and extant Clfx nitrogenases, nitrogen fixation is achieved by a possible alternative pathway for Mo-usage in the absence of NifEN, incorporating an unknown “proto-cofactor” (Figure 7a). Such unrefined Mo-usage is of advantage during transient Mo availabilities prior to ~2.3–2.5 Ga Earth surface oxygenation (Anbar et al., 2007). In the second stage, represented by AncC and extant F-Mc nitrogenases, a gene duplication event forms NifE, though...
potentially not sufficient for FeMo-cofactor synthesis (Figure 7b). Alternative Mo-usage may still be possible. In the third stage, represented by AncB, AncE, and extant Mb-Mc nitrogenases, a subsequent gene duplication event forms NifN (Figure 7c). Together, NifE and NifN are capable of synthesizing the FeMo-cofactor, resulting in canonical Mo-dependence. In the fourth stage, gene duplication of nif results in, first, vnf, followed by anf (Figure 7d). These V- and Fe-dependent enzymes rely on NifBEN biosynthetic components. Though the greater efficiency of Mo-nitrogenase results in widespread diversification, V- and Fe-nitrogenases provide selective advantage in microenvironments deficient in Mo. This model, built from the phylogenetic and ancestral sequence inferences provided here, as well as from decades of previous geobiological investigations of nitrogenase evolution, helps resolve outstanding questions regarding ancient metal dependence, and, importantly, provides testable hypotheses for future investigations. Such investigations may, for example, seek to clarify the nitrogen capability and metal dependence of uncharacterized Clfx and F-Mc nitrogenases, as well as experimentally resurrect and characterize ancestral nitrogenase sequences in the laboratory.
Figure 7. Proposed model for the evolution of nitrogenase metal specificity from an ancestral, alternative pathway for Mo-usage. Full description of stages (a)–(d) in the evolution of nitrogenase metal dependence is provided in the main text (Section 4.2). Possible alternative pathway for nitrogenase Mo-usage indicated in stages (a)–(b). Phase (a)–(d) are also mapped to analyzed ancestors within the nitrogenase phylogeny. Presence of NifB, NifE, and NifN in represented taxa indicated next to each clade of the phylogeny (F-Mc clade has only one species that harbors NifN).

4.3 Potential association of early nitrogen fixation with phototrophic metabolism

The relatively early-branching position of Clfx nitrogenases, hosted by strains in class Chloroflexi (green non-sulfur bacteria), suggests that biological nitrogen fixation may have had early associations with anoxygenic phototrophy. It has previously been proposed that biological nitrogen fixation originated in hydrogenotrophic methanogens due to the basal positioning of Mb-Mc (Methanobacteria and Methanococcus) or F-Mc (Firmicutes and Methanococcus) sequences in phylogenetic analyses of catalytic (i.e., NifHDK) and biosynthetic (i.e., NifBEN) nitrogenase
sequences (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Boyd & Peters, 2013; Raymond et al., 2004). By contrast, in our phylogenetic reconstruction, Clfx NifHDK sequences diverge earlier than both F-Mc and Mb-Mc sequences, with at least one represented strain having been observed to fix nitrogen (Keppen et al., 1989; Kuznetsov et al., 2011). An early association of biological nitrogen fixation with anoxygenic phototrophy would not be unexpected given the well-studied homology between nitrogenases and dark-operative protochlorophyllide oxidoreductases (Bch/ChlLNb), enzymes that catalyze part of the bacteriochlorophyll biosynthetic pathway (Y. Hu & Ribbe, 2015; Moser & Brocker, 2011; Raymond et al., 2004). Furthermore, Chloroflexi strains that possess uncharacterized Clfx nitrogenases also harbor BchLNb genes. Though the phylogeny presented here may still be consistent with the origin of the full FeMo-cofactor biosynthetic pathway (NifHDKBEN) in early methanogens (Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011; Boyd & Peters, 2013; Raymond et al., 2004), the basal position of Clfx sequences may evidence an earlier metabolic context for biological nitrogen fixation prior to the origin of the FeMo-cofactor. Future investigations of nitrogen fixation by other phototrophic Chloroflexi strains and additional taxa that harbor basal uncharacterized nitrogenases may resolve these aspects of nitrogenase origins.

CONCLUSION

We reconstructed the phylogenetic history of nitrogenase proteins, as well as inferred ancestral nitrogenase sequences, in order to explore the evolutionary trajectory of nitrogenase metal dependence. We find that, whereas modeled structural features of ancestral nitrogenases do not offer conclusive indications of ancient metal usage, active-site sequence features of ancestors most resemble those of extant Mo-nitrogenases. The absence of associated cofactor biosynthesis
proteins, considered necessary for FeMo-cofactor assembly, in several early-branching uncharacterized nitrogenase lineages evidences a possible alternative pathway for Mo-usage. We speculate that this alternative pathway may have preceded the evolution of the FeMo-cofactor and may today be present in extant uncharacterized nitrogenases, and we propose a model wherein canonical Mo-usage evolved via the stepwise introduction of FeMo-cofactor biosynthetic components following the divergence of more basal uncharacterized lineages. V-nitrogenases subsequently diversified, followed by Fe-nitrogenases, in agreement with previous phylogenetic inferences that Mo-dependence evolved first (Boyd, Hamilton, et al., 2011). This model helps to reconcile phylogenetic and geobiological explanations of nitrogenase evolution (Anbar & Knoll, 2002; Boyd, Anbar, et al., 2011; Boyd, Hamilton, et al., 2011). Future studies, particularly those that integrate experimental assessments of laboratory-resurrected ancestral nitrogenases, may continue to refine our understanding of nitrogenase and environmental coevolution.

**ACKNOWLEDGEMENTS**

This work was supported by a NASA Astrobiology Postdoctoral Fellowship (Garcia), the Harvard Origins of Life Initiative (Kacar, McShea), the National Science Foundation (Kacar, #1724090) and the John Templeton Foundation (Kacar, #58562 and #61239). We thank Andrew Knoll for constructive feedback.
REFERENCES

Aadland, K., Pugh, C., & Kolaczkowski, B. (2019). High-Throughput Reconstruction of Ancestral Protein Sequence, Structure, and Molecular Function. Methods Mol Biol, 1851, 135-170. doi:10.1007/978-1-4939-8736-8_8

Allen, R. M., Chatterjee, R., Ludden, P. W., & Shah, V. K. (1995). Incorporation of iron and sulfur from NifB cofactor into the iron-molybdenum cofactor of dinitrogenase. J Biol Chem, 270(45), 26890-26896. doi:10.1074/jbc.270.45.26890

Anbar, A. D., Duan, Y., Lyons, T. W., Arnold, G. L., Kendall, B., Creaser, R. A., . . . Buick, R. (2007). A whiff of oxygen before the great oxidation event? Science, 317(5846), 1903-1906. doi:10.1126/science.1140325

Anbar, A. D., & Knoll, A. H. (2002). Proterozoic ocean chemistry and evolution: a bioinorganic bridge? Science, 297(5584), 1137-1142. doi:10.1126/science.1069651

Anisimova, M., & Gascuel, O. (2006). Approximate likelihood-ratio test for branches: A fast, accurate, and powerful alternative. Syst Biol, 55(4), 539-552. doi:10.1080/10635150600755453

Benner, S. A., Sassi, S. O., & Gaucher, E. A. (2007). Molecular paleoscience: systems biology from the past. Adv Enzymol Relat Areas Mol Biol, 75, 1-132, xi.

Berman, H. M., Westbrook, J., Feng, Z., Gilliland, G., Bhat, T. N., Weissig, H., . . . Bourne, P. E. (2000). The Protein Data Bank. Nucleic Acids Res, 28(1), 235-242.

Boyd, E. S., Anbar, A. D., Miller, S., Hamilton, T. L., Lavin, M., & Peters, J. W. (2011). A late methanogen origin for molybdenum-dependent nitrogenase. Geobiology, 9(3), 221-232. doi:10.1111/j.1472-4669.2011.00278.x

Boyd, E. S., Costas, A. M., Hamilton, T. L., Mus, F., & Peters, J. W. (2015). Evolution of molybdenum nitrogenase during the transition from anaerobic to aerobic metabolism. J Bacteriol, 197(9), 1690-1699. doi:10.1128/JB.02611-14

Boyd, E. S., Hamilton, T. L., & Peters, J. W. (2011). An alternative path for the evolution of biological nitrogen fixation. Front Microbiol, 2, 205. doi:10.3389/fmicb.2011.00205

Boyd, E. S., & Peters, J. W. (2013). New insights into the evolutionary history of biological nitrogen fixation. Front Microbiol, 4, 201. doi:10.3389/fmicb.2013.00201

Brigle, K. E., Setterquist, R. A., Dean, D. R., Cantwell, J. S., Weiss, M. C., & Newton, W. E. (1987). Site-directed mutagenesis of the nitrogenase MoFe protein of Azotobacter vinelandii. Proc Natl Acad Sci U S A, 84(20), 7066-7069.

Bulen, W. A., & LeComte, J. R. (1966). The nitrogenase system from Azotobacter: two-enzyme requirement for N2 reduction, ATP-dependent H2 evolution, and ATP hydrolysis. Proc Natl Acad Sci U S A, 56(3), 979-986. doi:10.1073/pnas.56.3.979

Camacho, C., Coulouris, G., Avagyan, V., Ma, N., Papadopoulos, J., Bealer, K., & Madden, T. L. (2009). BLAST+: architecture and applications. BMC Bioinformatics, 10, 421. doi:10.1186/1471-2105-10-421

Canfield, D. E., Glazer, A. N., & Falkowski, P. G. (2010). The evolution and future of Earth's nitrogen cycle. Science, 330(6001), 192-196. doi:10.1126/science.1186120

Chen, I. A., Chu, K., Palaniappan, K., Pillay, M., Ratner, A., Huang, J., . . . Kyrpides, N. C. (2019). IMG/M v.5.0: an integrated data management and comparative analysis system for microbial genomes and microbiomes. Nucleic Acids Res, 47(D1), D666-D677. doi:10.1093/nar/gky901
Christiansen, J., Cash, V. L., Seefeldt, L. C., & Dean, D. R. (2000). Isolation and characterization of an acetylene-resistant nitrogenase. *J Biol Chem, 275*(15), 11459-11464.

Curatti, L., Hernandez, J. A., Igarashi, R. Y., Soboh, B., Zhao, D., & Rubio, L. M. (2007). In vitro synthesis of the iron-molybdenum cofactor of nitrogenase from iron, sulfur, molybdenum, and homocitrate using purified proteins. *Proc Natl Acad Sci U S A, 104*(45), 17626-17631. doi:10.1073/pnas.0703050104

Dekas, A. E., Poretsky, R. S., & Orphan, V. J. (2009). Deep-sea archaea fix and share nitrogen in methane-consuming microbial consortia. *Science, 326*(5951), 422-426. doi:10.1126/science.1178223

Dos Santos, P. C., Fang, Z., Mason, S. W., Setubal, J. C., & Dixon, R. (2012). Distribution of nitrogen fixation and nitrogenase-like sequences amongst microbial genomes. *BMC Genomics, 13*, 162. doi:10.1186/1471-2164-13-162

Durrant, J. D., Votapka, L., Sorensen, J., & Amaro, R. E. (2014). POVME 2.0: An Enhanced Tool for Determining Pocket Shape and Volume Characteristics. *J Chem Theory Comput, 10*(11), 5047-5056. doi:10.1021/ct500381c

Eady, R. R. (1996). Structure-function relationships of alternative nitrogenases. *Chemical Reviews, 96*(7), 3013-3030. doi:10.1021/cr950057h

Edgar, R. C. (2004). MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res, 32*(5), 1792-1797. doi:10.1093/nar/gkh340

Falkowski, P. G. (1997). Evolution of the nitrogen cycle and its influence on the biological sequestration of CO2 in the ocean. *Nature, 387*(6630), 272-275. doi:10.1038/387272a0

Fixen, K. R., Zheng, Y., Harris, D. F., Shaw, S., Yang, Z. Y., Dean, D. R., . . . Harwood, C. S. (2016). Light-driven carbon dioxide reduction to methane by nitrogenase in a photosynthetic bacterium. *Proc Natl Acad Sci U S A, 113*(36), 10163-10167. doi:10.1073/pnas.1611043113

Garcia, A. K., & Kacar, B. (2019). How to resurrect ancestral proteins as proxies for ancient biogeochemistry. *Free Radic Biol Med*. doi:10.1016/j.freeradabiomed.2019.03.033

Gomez-Fernandez, B. J., Garcia-Ruiz, E., Martin-Diaz, J., Gomez de Santos, P., Santos-Moriano, P., Plou, F. J., . . . Alcalde, M. (2018). Directed -in vitro- evolution of Precambrian and extant Rubiscos. *Sci Rep, 8*(1), 5532. doi:10.1038/s41598-018-23869-3

Hageman, R. V., & Burris, R. H. (1978). Nitrogenase and nitrogenase reductase associate and dissociate with each catalytic cycle. *Proc Natl Acad Sci U S A, 75*(6), 2699-2702. doi:10.1073/pnas.75.6.2699

Hales, B. J., Case, E. E., Morningstar, J. E., Dzeda, M. F., & Mauterer, L. A. (1986). Isolation of a new vanadium-containing nitrogenase from Azotobacter vinelandii. *Biochemistry, 25*(23), 7251-7255. doi:10.1021/bi00371a001

Hamilton, T. L., Ludwig, M., Dixon, R., Boyd, E. S., Dos Santos, P. C., Setubal, J. C., . . . Peters, J. W. (2011). Transcriptional profiling of nitrogen fixation in Azotobacter vinelandii. *J Bacteriol, 193*(17), 4477-4486. doi:10.1128/JB.05099-11

Hanson-Smith, V., & Johnson, A. (2016). PhyloBot: A Web Portal for Automated Phylogenetics, Ancestral Sequence Reconstruction, and Exploration of Mutational Trajectories. *PLoS Comput Biol, 12*(7), e1004976. doi:10.1371/journal.pcbi.1004976

Hanson-Smith, V., Kolaczkowski, B., & Thornton, J. W. (2010). Robustness of ancestral sequence reconstruction to phylogenetic uncertainty. *Mol Biol Evol, 27*(9), 1988-1999. doi:10.1093/molbev/msq081
Harris, D. F., Lukoyanov, D., Kallas, H., Trncik, C., Yang, Z., Compton, P., . . . Seefeldt, L. C. (2019). Mo-, V-, and Fe-nitrogenases use a universal eight-electron reductive-elimination mechanism to achieve N₂ reduction. *Biochemistry*.

Harris, D. F., Lukoyanov, D. A., Shaw, S., Compton, P., Tokmina-Lukaszewska, M., Bothner, B., . . . Seefeldt, L. C. (2018). Mechanism of N2 Reduction Catalyzed by Fe-Nitrogenase Involves Reductive Elimination of H2. *Biochemistry, 57*(5), 701-710. doi:10.1021/acs.biochem.7b01142

Harris, D. F., Yang, Z. Y., Dean, D. R., Seefeldt, L. C., & Hoffman, B. M. (2018). Kinetic Understanding of N2 Reduction versus H2 Evolution at the E4(4H) Janus State in the Three Nitrogenases. *Biochemistry, 57*(39), 5706-5714. doi:10.1021/acs.biochem.8b00784

Hoffman, B. M., Lukoyanov, D., Yang, Z. Y., Dean, D. R., & Seefeldt, L. C. (2014). Mechanism of nitrogen fixation by nitrogenase: the next stage. *Chem Rev, 114*(8), 4041-4062. doi:10.1021/cr400641x

Hu, B., Harris, D. F., Dean, D. R., Liu, T. L., Yang, Z. Y., & Seefeldt, L. C. (2018). Electrocatalytic CO₂ reduction catalyzed by nitrogenase MoFe and FeFe proteins. *Bioelectrochemistry, 120*, 104-109. doi:10.1016/j.bioelechem.2017.12.002

Hu, Y., Lee, C. C., & Ribbe, M. W. (2011). Extending the Carbon Chain: Hydrocarbon Formation Catalyzed by Vanadium/Molybdenum Nitrogenases. *Science, 333*(6043), 753-755. doi:10.1126/science.1206883

Hu, Y., & Ribbe, M. W. (2011). Biosynthesis of Nitrogenase FeMoco. *Coord Chem Rev, 255*(9-10), 1218-1224. doi:10.1016/j.ccr.2010.11.018

Hu, Y., & Ribbe, M. W. (2015). Nitrogenase and homologs. *J Biol Inorg Chem, 20*(2), 435-445. doi:10.1007/s00775-014-1225-3

Joerger, R. D., & Bishop, P. E. (1988). Bacterial alternative nitrogen fixation systems. *Crit Rev Microbiol, 16*(1), 1-14. doi:10.3109/10408418809104465

Kacar, B., Hanson-Smith, V., Adam, Z. R., & Boekelheide, N. (2017). Constraining the timing of the Great Oxidation Event within the Rubisco phylogenetic tree. *Geobiology, 15*(5), 628-640. doi:10.1111/gbi.12243

Kennedy, C., & Dean, D. (1992). The nifU, nifS and nifV gene products are required for activity of all three nitrogenases of Azotobacter vinelandii. *Mol Gen Genet, 231*(3), 494-498.

Keppen, O. I., Lebedeva, N. V., Troshina, O. Y., & Rodionov, Y. V. (1989). The nitrogenase activity of filamentous phototrophic green bacterium. *Mikrobiologiya, 58*, 520-521.

Kim, C. H., Newton, W. E., & Dean, D. R. (1995). Role of the MoFe protein alpha-subunit histidine-195 residue in FeMo-cofactor binding and nitrogenase catalysis. *Biochemistry, 34*(9), 2798-2808.

Krahn, E., Weiss, R., Krockel, M., Groppe, J., Henkel, G., Cramer, P., . . . Muller, A. (2002). The Fe-only nitrogenase from Rhodobacter capsulatus: identification of the cofactor, an unusual, high-nuclearity iron-sulfur cluster, by Fe K-edge EXAFS and 57Fe Mossbauer spectroscopy. *J Biol Inorg Chem, 7*(1-2), 37-45. doi:10.1007/s007750100263

Kuznetsov, B. B., Ivanovsky, R. N., Keppen, O. I., Sukhacheva, M. V., Bumazhkin, B. K., Patutina, E. O., . . . Skryabin, K. G. (2011). Draft genome sequence of the anoxygenic filamentous phototrophic bacterium Oscillochloris trichoides subsp. DG-6. *J Bacteriol, 193*(1), 321-322. doi:10.1128/JB.00931-10

Le, S. Q., & Gascuel, O. (2008). An improved general amino acid replacement matrix. *Mol Biol Evol, 25*(7), 1307-1320. doi:10.1093/molbev/msn067
Lee, C. C., Tanifuji, K., Newcomb, M., Liedtke, J., Hu, Y., & Ribbe, M. W. (2018). A Comparative Analysis of the CO-Reducing Activities of MoFe Proteins Containing Mo- and V-Nitrogenase Cofactors. ChemBioChem, 19(7), 649-653. doi:10.1002/cbic.201800035

Liu, Y., Schmidt, B., & Maskell, D. L. (2010). MSAProbs: multiple sequence alignment based on pair hidden Markov models and partition function posterior probabilities. Bioinformatics, 26(16), 1958-1964. doi:10.1093/bioinformatics/btq338

Lyons, T. W., Reinhard, C. T., & Planavsky, N. J. (2014). The rise of oxygen in Earth's early ocean and atmosphere. Nature, 506(7488), 307-315. doi:10.1038/nature13068

MacKay, B. A., & Fryzuk, M. D. (2004). Dinitrogen coordination chemistry: on the biomimetic borderlands. Chem Rev, 104(2), 385-401. doi:10.1021/cr020610c

McGlynn, S. E., Boyd, E. S., Peters, J. W., & Orphan, V. J. (2012). Classifying the metal dependence of uncharacterized nitrogenases. Front Microbiol, 3, 419. doi:10.3389/fmicb.2012.00419

McRose, D. L., Zhang, X., Kraepiel, A. M., & Morel, F. M. (2017). Diversity and Activity of Alternative Nitrogenases in Sequenced Genomes and Coastal Environments. Front Microbiol, 8, 267. doi:10.3389/fmicb.2017.00267

Mehta, M. P., & Baross, J. A. (2006). Nitrogen fixation at 92 degrees C by a hydrothermal vent archaeon. Science, 314(5806), 1783-1786. doi:10.1126/science.1134772

Moore, E. K., Jelen, B. I., Giovannelli, D., Raanan, H., & Falkowski, P. G. (2017). Metal availability and the expanding network of microbial metabolisms in the Archaean eon. Nature Geoscience, 10(9), 629-636. doi:10.1038/ngeo3006

Mus, F., Alleman, A. B., Pence, N., Seefeldt, L. C., & Peters, J. W. (2018). Exploring the alternatives of biological nitrogen fixation. Metallomics, 10(4), 523-538. doi:10.1039/c8mt00038g

Mus, F., Colman, D. R., Peters, J. W., & Boyd, E. S. (2019). Geobiological feedbacks, oxygen, and the evolution of nitrogenase. Free Radic Biol Med. doi:10.1016/j.freeradbiomed.2019.01.050

O'Leary, N. A., Wright, M. W., Brister, J. R., Ciufo, S., Haddad, D., McVeigh, R., . . . Pruitt, K. D. (2016). Reference sequence (RefSeq) database at NCBI: current status, taxonomic expansion, and functional annotation. Nucleic Acids Res, 44(D1), D733-745. doi:10.1093/nar/gkv1189

Ormeño-Orrillo, E., Hungria, M., & Martinez-Romero, E. (2013). Dinitrogen-Fixing Prokaryotes. In The Prokaryotes (pp. 427-451).

Quang, L. S., Gascuel, O., & Lartillot, N. (2008). Empirical profile mixture models for phylogenetic reconstruction. Bioinformatics, 24(20), 2317-2323. doi:10.1093/bioinformatics/btn445

R Core Team. (2014). R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing.

Raymond, J., Siebert, J. L., Staples, C. R., & Blankenship, R. E. (2004). The natural history of nitrogen fixation. Mol Biol Evol, 21(3), 541-554. doi:10.1093/molbev/msh047

Rebelein, J. G., Lee, C. C., Newcomb, M., Hu, Y., & Ribbe, M. W. (2018). Characterization of an M-Cluster-Substituted Nitrogenase VFe Protein. MBio, 9(2). doi:10.1128/mBio.00310-18
Roll, J. T., Shah, V. K., Dean, D. R., & Roberts, G. P. (1995). Characteristics of NIFNE in Azotobacter vinelandii strains. Implications for the synthesis of the iron-molybdenum cofactor of dinitrogenase. *J Biol Chem*, 270(9), 4432-4437. doi:10.1074/jbc.270.9.4432

Rubio, L. M., & Ludden, P. W. (2008). Biosynthesis of the iron-molybdenum cofactor of nitrogenase. *Annu Rev Microbiol*, 62, 93-111. doi:10.1146/annurev.micro.62.081307.162737

Sali, A., & Blundell, T. L. (1993). Comparative protein modelling by satisfaction of spatial restraints. *J Mol Biol*, 234(3), 779-815. doi:10.1006/jmbi.1993.1626

Sarma, R., Barney, B. M., Keable, S., Dean, D. R., Seefeldt, L. C., & Peters, J. W. (2010). Insights into substrate binding at FeMo-cofactor in nitrogenase from the structure of an alpha-70(Ile) MoFe protein variant. *J Inorg Biochem*, 104(4), 385-389. doi:10.1016/j.jinorgbio.2009.11.009

Schmid, B., Einsle, O., Chiu, H. J., Willing, A., Yoshida, M., Howard, J. B., & Rees, D. C. (2002). Biochemical and structural characterization of the cross-linked complex of nitrogenase: comparison to the ADP-AlF4(-)-stabilized structure. *Biochemistry*, 41(52), 15557-15565.

Setubal, J. C., dos Santos, P., Goldman, B. S., Ertesvag, H., Espin, G., Rubio, L. M., . . . Wood, D. (2009). Genome sequence of Azotobacter vinelandii, an obligate aerobe specialized to support diverse anaerobic metabolic processes. *J Bacteriol*, 191(14), 4534-4545. doi:10.1128/JB.00504-09

Shah, V. K., Allen, J. R., Spangler, N. J., & Ludden, P. W. (1994). In vitro synthesis of the iron-molybdenum cofactor of nitrogenase. Purification and characterization of NifB cofactor, the product of NIFB protein. *J Biol Chem*, 269(2), 1154-1158.

Shah, V. K., Imperial, J., Ugalde, R. A., Ludden, P. W., & Brill, W. J. (1986). In vitro synthesis of the iron-molybdenum cofactor of nitrogenase. *Proc Natl Acad Sci U S A*, 83(6), 1636-1640. doi:10.1073/pnas.83.6.1636

Shih, P. M., Occhialini, A., Cameron, J. C., Andralojc, P. J., Parry, M. A., & Kerfeld, C. A. (2016). Biochemical characterization of predicted Precambrian RuBisCO. *Nat Commun*, 7, 10382. doi:10.1038/ncomms10382

Sippel, D., & Einsle, O. (2017). The structure of vanadium nitrogenase reveals an unusual bridging ligand. *Nat Chem Biol*, 13(9), 956-960. doi:10.1038/nchembio.2428

Sippel, D., Rohde, M., Netzer, J., Trncik, C., Gies, J., Grunau, K., . . . Einsle, O. (2018). A bound reaction intermediate sheds light on the mechanism of nitrogenase. *Science*, 359(6383), 1484-1489. doi:10.1126/science.aar2765

Soboh, B., Boyd, E. S., Zhao, D., Peters, J. W., & Rubio, L. M. (2010). Substrate specificity and evolutionary implications of a NifDK enzyme carrying NifB-co at its active site. *FEBS Lett*, 584(8), 1487-1492. doi:10.1016/j.febslet.2010.02.064

Spatzal, T., Aksoyoglu, M., Zhang, L., Andrade, S. L., Schleicher, E., Weber, S., . . . Einsle, O. (2011). Evidence for interstitial carbon in nitrogenase FeMo cofactor. *Science*, 334(6058), 940. doi:10.1126/science.1214025

St John, R. T., Johnston, H. M., Seidman, C., Garfinkel, D., Gordon, J. K., Shah, V. K., & Brill, W. J. (1975). Biochemistry and genetics of Klebsiella pneumoniae mutant strains unable to fix N2. *J Bacteriol*, 121(3), 759-765.

Stamatakis, A. (2014). RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics*, 30(9), 1312-1313. doi:10.1093/bioinformatics/btu033
Stueken, E. E., Buick, R., Guy, B. M., & Koehler, M. C. (2015). Isotopic evidence for biological nitrogen fixation by molybdenum-nitrogenase from 3.2 Gyr. *Nature, 520*(7549), 666-669. doi:10.1038/nature14180

Tal, S., Chun, T. W., Gavini, N., & Burgess, B. K. (1991). The delta nifB (or delta nifE) FeMo cofactor-deficient MoFe protein is different from the delta nifH protein. *J Biol Chem, 266*(16), 10654-10657.

Thornton, J. W. (2004). Resurrecting ancient genes: experimental analysis of extinct molecules. *Nat Rev Genet, 5*(5), 366-375. doi:10.1038/nrg1324

Whelan, S., & Goldman, N. (2001). A general empirical model of protein evolution derived from multiple protein families using a maximum-likelihood approach. *Mol Biol Evol, 18*(5), 691-699. doi:10.1093/oxfordjournals.molbev.a003851

Yang, Z. (1993). Maximum-likelihood estimation of phylogeny from DNA sequences when substitution rates differ over sites. *Mol Biol Evol, 10*(6), 1396-1401. doi:10.1093/oxfordjournals.molbev.a040082

Yang, Z. (2007). PAML 4: phylogenetic analysis by maximum likelihood. *Mol Biol Evol, 24*(8), 1586-1591. doi:10.1093/molbev/msm088

Yang, Z. Y., Moure, V. R., Dean, D. R., & Seefeldt, L. C. (2012). Carbon dioxide reduction to methane and coupling with acetylene to form propylene catalyzed by remodeled nitrogenase. *Proc Natl Acad Sci U S A, 109*(48), 19644-19648. doi:10.1073/pnas.1213159109

Zhang, X., McRose, D. L., Darnajoux, R., Bellenger, J. P., Morel, F. M. M., & Kraepiel, A. M. L. (2016). Alternative nitrogenase activity in the environment and nitrogen cycle implications. *Biogeochemistry, 127*(2-3), 189-198. doi:10.1007/s10533-016-0188-6

Zheng, Y., Harris, D. F., Yu, Z., Fu, Y., Poudel, S., Ledbetter, R. N., . . . Harwood, C. S. (2018). A pathway for biological methane production using bacterial iron-only nitrogenase. *Nat Microbiol, 3*(3), 281-286. doi:10.1038/s41564-017-0091-5