Origin and Evolution of Nitrogen Fixation in Prokaryotes

Hong-Wei Pi,†,1,2 Jinn-Jy Lin,†,2 Chi-An Chen,2,3 Po-Hsiang Wang,4,5 Yin-Ru Chiang,2 Chieh-Chen Huang,6 Chiu-Chung Young,7 and Wen-Hsiung Li*,2,8

1Ph.D. Program in Microbial Genomics, National Chung Hsing University and Academia Sinica, Taiwan
2Biodiversity Research Center, Academia Sinica, Taipei 11529, Taiwan
3Genome and Systems Biology Degree Program, Academia Sinica and National Taiwan University, Taipei 10617, Taiwan
4Graduate Institute of Environmental Engineering, National Central University, Taoyuan 32001, Taiwan
5Earth-Life Science Institute, Tokyo Institute of Technology, Tokyo 145-0061, Japan
6Department of Life Sciences, National Chung Hsing University, Taichung 402, Taiwan
7Department of Soil and Environmental Sciences, College of Agriculture and Natural Resources, National Chung Hsing University, Taichung 402, Taiwan
8Department of Ecology and Evolution, University of Chicago, Chicago 60637, IL, USA
†These authors contributed equally to this work.

*Corresponding author: E-mail: whli@uchicago.edu.
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Abstract

The origin of nitrogen fixation is an important issue in evolutionary biology. While nitrogen is required by all living organisms, only a small fraction of bacteria and archaea can fix nitrogen. The prevailing view is that nitrogen fixation first evolved in archaea and was later transferred to bacteria. However, nitrogen-fixing (Nif) bacteria are far larger in number and far more diverse in ecological niches than Nif archaea. We, therefore, propose the bacteria-first hypothesis, which postulates that nitrogen fixation first evolved in bacteria and was later transferred to archaea. As >30,000 prokaryotic genomes have been sequenced, we conduct an in-depth comparison of the two hypotheses. We first identify the six genes involved in nitrogen fixation in all sequenced prokaryotic genomes and then reconstruct phylogenetic trees using the six Nif proteins individually or in combination. In each of these trees, the earliest lineages are bacterial Nif protein sequences and in the oldest clade (group) the archaeal sequences are all nested inside bacterial sequences, suggesting that the Nif proteins first evolved in bacteria. The bacteria-first hypothesis is further supported by the observation that the majority of Nif archaea carry the major bacterial Mo (molybdenum) transporter (ModABC) rather than the archaeal Mo transporter (WtpABC). Moreover, in our phylogeny of all available ModA and WtpA protein sequences, the earliest lineages are bacterial sequences while archaeal sequences are nested inside bacterial sequences. Furthermore, the bacteria-first hypothesis is supported by available isotopic data. In conclusion, our study strongly supports the bacteria-first hypothesis.

Key words: nitrogen fixation, nitrogenase, molybdenum transporter, bacteria, archaea.

Introduction

Nitrogen (N), which accounts for almost 80% of the earth’s atmosphere, is essential for all living creatures but no organism can directly utilize nitrogen (N₂) (Erisman et al. 2008). Nitrogen fixation is the conversion of N₂ to bioavailable NH₃ (Boyd and Peters 2013; Mus et al. 2019). Only a small fraction of bacteria and archaea can fix nitrogen (Boyd and Peters 2013). However, nitrogen-fixing (Nif) bacteria are physiologically and ecologically diverse. Most of them are free-living, including aerobic chemooorganotrophs (Setubal et al. 2009); anoxygenic and oxygenic photosynthetic (Philippi et al. 2021; Watanabe and Horiike 2021); and anaerobic chemooorganotrophs (Li et al. 2019). Many of the remaining Nif bacteria are symbiotic with plants (Li et al. 2015; de Lajudie and Young 2017; Roy et al. 2020) or animals (Russell et al. 2009; König et al. 2016; Shukla et al. 2016). In contrast, most Nif archaea are exclusively free-living and belong to the chemolithotrophic methanogens (Mehta and Baross 2006; Boyd, Anbar, et al. 2011; Koirala and Brözel 2021). Free-living Nif species play an important role in nitrogen cycle in the modern biosphere as well as in the ancient environments.

Biological nitrogen fixation is catalyzed by nitrogenases, which have three variants, the Mo (molybdenum)-dependent nitrogenase (Nif), V (vanadium)-dependent nitrogenase (Vnf), and Fe-only nitrogenase (Anf) (Mus et al. 2018). Nitrogenases are composed of two distinct metalloproteins, namely dinitrogenase (the catalytic component)}
and dinitrogenase reductase (the electron transfer component) (Burén et al. 2020). Both components contain Fe–S clusters, and the dinitrogenase component contains a characteristic iron–molybdenum cofactor, FeMoco. Alternative nitrogenases contain either V + Fe or Fe-only in their dinitrogenase component (Mus et al. 2018). The nif (nitrogen-fixing) genes of conventional Mo-dependent nitrogenase, which is encoded by nifH (the dinitrogenase reductase subunit) and nifD and nifK (the dinitrogenase subunits) (Rubio and Ludden 2005; Hu and Ribbe 2016; López-Torrejón et al. 2016), have been well studied (Dos Santos et al. 2012; Mus et al. 2018, 2019; Zheng et al. 2018; Albright et al. 2019; Lee et al. 2019; Burén et al. 2020; Garcia et al. 2020). The nifD and nifK genes encode, respectively, the α- and β-subunit of dinitrogenase, which form a α3β2-tetramer (Rubio and Ludden 2005; Hu and Ribbe 2016; López-Torrejón et al. 2016). The nifB gene product functions in the biosynthesis of the Fe and S donors for the metal cofactor (Rubio and Ludden 2005; Dos Santos et al. 2012; Hu and Ribbe 2016; López-Torrejón et al. 2016). Although various sets of nif genes have been used as criteria to identify Nif species (Dos Santos et al. 2012; Poudel et al. 2018; Albright et al. 2019; Koirala and Brözel 2021), the entire set of the above six nif genes (i.e., nifHDKENB) has been proposed to be the definition of Nif species (Dos Santos et al. 2012; Mus et al. 2019). However, nifN or both nifE and nifN may not be essential because this gene is absent in the genomes of some Nif Chloroflexota (bacteria) (Ivanovsky et al. 2021), Firmicutes_A (bacteria) (Chen et al. 2021), Elusimicrobiota (bacteria) (Zheng, Dietrich, et al. 2016), and Nif methanogens (archaea) (Mehta and Baross 2006). We therefore will also consider the criterion of “the five-nif-gene set” (nifHDKENB), which includes no nifN, and that of “the four-nif-gene set” (nifHDKB), which includes no nifE. The molybdenum (Mo) is an essential element and participates in a variety of metalloenzymes in both bacteria and archaea (Zhang and Gladyshev 2008; Peng et al. 2018). Previous studies showed that the Mo-dependent enzymes (molybdoenzymes) evolved very early in prokaryotes, and it may have evolved their unique systems in bacteria and in archaea in very ancient times (Zhang and Gladyshev 2008; Peng et al. 2018). Moreover, in prokaryotes, nitrogenase is the only molybdoenzyme that contains the iron–Mo cofactor (FeMoco), while the remaining molybdoenzymes contain the molybdenum cofactor (Moco) (Zhang and Gladyshev 2008; Peng et al. 2018). The Mo transporter, which is the essential protein for prokaryotes to uptake Mo, may regulate nitrogenase (Demtröder et al. 2019). Therefore, the Mo transporter might have played an important role in the evolution of nitrogen fixation.

It has been proposed that nitrogen fixation first evolved in archaea and was later transferred to bacteria (Raymond et al. 2004; Boyd, Hamilton, et al. 2011; Boyd and Peters 2013; Mus et al. 2018, 2019; Koirala and Brözel 2021); we call this hypothesis “the archaea-first hypothesis”. This hypothesis was based on the observation that the earliest lineages in the reconstructed phylogeny of NiFH, D, K, E, and N sequences and in that of the radical S-adenosyl-L-methionine (SAM) domains of NifB proteins were archaeal sequences (Raymond et al. 2004; Boyd, Anbar, et al. 2011). As an alternative to the archaea-first hypothesis, it has been proposed that nitrogen fixation evolved in the last universal common ancestor (LUCA) (Raymond et al. 2004; Weiss et al. 2016); we call it the LUCA-first hypothesis. Here, we propose “the bacteria-first hypothesis”, which postulates that nitrogen fixation first evolved in bacteria and was later transferred to archaea. These three hypotheses are mutually exclusive, so that only one of them can be true.

In this study, we applied the criterion of the six-nif-gene set (nifHDKENB) to identify Nif species in the available genomic sequence data (Parks et al. 2018, 2020; Mendler et al. 2019); we also identified those species that carry nifHDKB and those that carry nifHDKEB. To distinguish between the archaea-first and the bacteria-first hypothesis, we reconstructed the following gene trees: The first one was the phylogeny of NiFH sequences because among the Nif proteins, NiFH is the widest spread in prokaryotes. The second tree was the phylogeny of NifD, K, E, and N sequences because these proteins are paralogous. The third tree was the phylogeny of radical SAM domains of NifB proteins. The fourth tree was the phylogeny of the concatenated sequences of NifH, D, and K sequences that are tightly linked in the genome. In each of these trees, our aim was to find out whether a gene under study first appeared in bacteria or archaea. In addition, we identified modABC- and wtpABC-gene-carrying species. It is known that ModABC is the bacterial Mo transporter while WtpABC is the major Mo transporter in archaea (Peng et al. 2018). Thus, studying the association of these two Mo-transporters with Nif species (i.e., the six-nif-gene set) is helpful for distinguishing between the two hypotheses. For this purpose, we also reconstructed the phylogeny of ModA and WtpA sequences. Using these phylogenetic trees, we argued for the bacteria-first hypothesis.

Results
Identifying nif Genes
We analyzed 30,238 bacterial and 1,672 archael genomes from AnnoTree and GTDB (Parks et al. 2018, 2020; Mendler et al. 2019), and found 3,485 bacterial species and 373 archaeal species that contain at least one of the nifH, nifD, nifK, nifE, nifN, and nifB genes (table 1). The number of species that carry the six-nif-gene set is ~18 times higher in bacteria (1,420) than in archaea (78) (table 1). Moreover, the number of species carrying the five-nif-gene set is ~4 times higher in bacteria (53) than in archaea (13); and the number of species carrying the four-nif-gene set is ~9 times higher in bacteria (171) than in archaea (18).
Thus, most known Nif species are bacteria. However, the proportion of bacterial species carrying the six-gene set (1,420/30,238 = 4.70%) is almost the same as that of archaeal species carrying the six-gene set (78/1,672 = 4.67%). The detailed information of nif-gene-harboring species in our collection and their genomic characteristics is given in supplementary data S1, Supplementary Material online.

Previous studies suggested that Mo-nitrogenase evolved earlier than V- and Fe-nitrogenase and most anf/ vnf-gene-harboring species also contain nif genes (Mus et al. 2018, 2019; Garcia et al. 2020). Our collection identified conventional Mo-nitrogenase, so it included species that also contain the V- or Fe-nitrogenase. The additional γ-subunit (encoded by vnfG or anfG), of which the function has been unraveled only recently, is only present in V- and Fe-nitrogenases (Mus et al. 2018; Garcia et al. 2020), and the γ-subunit (vnfG/anfG) is attached to the α-subunit (vnf/anfD). So, to identify V- and Fe-nitrogenases, we used the location of vnf/anfG genes in the genomes to identify vnf/anfH, D, K genes. We identified 25 bacteria and 8 archaea that contain vnfG, and 101 bacteria and 6 archaea that contain anfG (table 1 and supplementary data S1, Supplementary Material online). In total, our collection includes 114 bacteria and 5 archaea that contain Mo-nitrogenase and V- or Fe-nitrogenase, and 6 bacteria and 4 archaea that contain all three types of nitrogenases. In total, we identified 130 Nif species that contain V- or Fe-nitrogenase or both.

### Phylogenetic Tree of Nif/Vnf/AnfH Proteins

The nifH gene encodes the dinitrogenase reductase and many studies used nifH to investigate the diversity of Nif species in environments (Zehr and McReynolds 1989; Affourtit et al. 2001; Hamelin et al. 2002; Langlois et al. 2005; Hamilton et al. 2011). Therefore, we first reconstruct the phylogeny of Nif/Vnf/AnfH protein sequences (fig. 1). Our study includes 3,535 Nif/Vnf/AnfH protein sequences and uses 376 Bch/ChlN and BchX sequences, which are NifH homologous proteins (Nomata et al. 2006; Tahon et al. 2016; Garcia et al. 2020), as the outgroup. We use the IQ-TREE method to reconstruct phylogenetic trees (Minh et al. 2020) and ModelFinder (Kalyaanamoorthy et al. 2017), which is implemented in IQ-TREE, to select the best-fit model of protein sequence evolution with ultrafast bootstrap (1,000 replicates) (Hoang et al. 2018). The model was selected according to the Bayesian information criterion (BIC) (Posada and Buckley 2004) is LG + R10.

In figure 1, bacterial sequences (labeled by blue color) form well-supported earliest lineages while archaeal sequences (labeled by red color) are all nested inside bacterial sequences. This result suggests that NifH first evolved in bacteria. Moreover, the Vnf/AnfH sequences appear in the latest lineages, consistent with the previous inference that V- and Fe-nitrogenases evolved later than Mo-nitrogenase (Mus et al. 2018, 2019; Garcia et al. 2020).

To show more details of the phylogeny of Nif/Vnf/AnfH proteins, we classify the protein sequences under study into Groups I, II, III, IV, Vnf, and Anf according to previous studies (Raymond et al. 2004; Howard et al. 2013; Zheng, Dietrich, et al. 2016; North et al. 2020) (fig. 2, supplementary fig. S1, Supplementary Material online). Figure 2 indicates that Group IV is the oldest group and Groups II, III/Vnf/Anf, and I are younger. As the sequences of archaea-1 and -2 are nested inside bacterial Groups IV sequences, none of them are likely the ancestral sequences. The archaeal sequences in archaea-3 are split into five subgroups but all belong to Halobacteriota. One of Halobacteriota subgroups is the early lineage in Group II, while the other four are nested inside Group II bacterial sequences, implying five different HGT events between bacteria and Halobacteriota because in each case the gene tree is drastically different from the species tree. Bacterial sequences form the early lineage in Group III, and the archaea-4 sequences, most of which belong to Methanobacteriota, are nested inside Group III bacterial sequences. Group I contains only bacteria, so its common ancestor should be a bacterium. In Group Vnf/ AnfH, bacterial sequences form the earliest lineages and the archaeal Vnf/AnfH sequences are nested inside bacterial Vnf/AnfH sequences, implying that Vnf/AnfH first evolved in bacteria. Additionally, most Vnf/AnfH sequences are located next to Group III while some Vnf/AnfH sequences are nested inside Groups I and II NifH sequences (the red arrows in fig. 2) as in a previous study (Angel et al. 2018). This finding suggests that Vnf/AnfH evolved from the duplication of NifH as previously proposed (Mus et al. 2018; Garcia et al. 2020).

### Phylogenetic Tree of Nif/Vnf/AnfH, K, E, and N Proteins

It has been proposed that NifDK were the ancestral sequences and that NifE were derived from a duplication of NifDK (Fani et al. 2000; Raymond et al. 2004; Boyd, Anbar, et al. 2011; Dos Santos et al. 2012; Garcia et al. 2020). We call this hypothesis the NifDK-first hypothesis. This hypothesis can be depicted by figure 3, in which two linked proto-proteins evolved in one lineage into Bch/ChINB and BchYZ (Muraki et al. 2010; Boyd, Anbar, et al. 2011; Garcia et al. 2020), and in another lineage...
**Fig. 1.** Simplified phylogeny of 3,535 Nif/Vnf/AnfH sequences. The scale bar denotes 0.1 amino acid substitutions per residue site. The solid and empty black circles denote the bootstrap supports of 90–100% and 80–89%, respectively, and the empty triangle denotes a bootstrap value of 70–79%. The outgroup is labeled by black color. The bacterial group is labeled by blue color and archaeal groups by dark red color. We use the light-independent protochlorophyllide reductase subunit L (Bch/ChlL) and chlorophyllide a reductase subunit X (BchX) sequences as the outgroup. We use the IQ-TREE method to reconstruct phylogenetic trees and ModelFinder to select the best-fit model of protein sequence evolution with ultrafast bootstrap (1,000 replicates). The model selected according to the BIC is LG + R10.
into NifDK. NifDK then expanded into a group of NifDK sequences (Group IV), one of which was duplicated to become NifDKEN, which then evolved into Groups I, II, III, and IV. Boyd, Anbar, et al. (2011) constructed a phylogeny of ∼150 Nif/Vnf/AnfD, K, E, and N protein sequences from 40 genomic sequences and proposed the archaea-first hypothesis. We reconstruct a phylogeny of 7,421 Nif/Vnf/AnfD, K, E, and N sequences from 1,489 genomes (fig. 4A), using the IQ-tree software (Minh et al. 2020) and using 20 Bch/ChlN and Bch/ChlB sequences as the outgroup as in a previous study (Garcia et al. 2020). The best-fit model of protein sequence evolution is LG + F + R10. Supplementary figure S2, Supplementary Material online is a detailed version of figure 4.

In figure 4A, we label NifD by light green, NifK by dark green, NifE by light blue and NifN by dark blue. To make it easier to understand, we divided the Nif protein sequences into Groups I, II, III, and IV as in previous studies (Raymond et al. 2004; Howard et al. 2013; Zheng, Dietrich, et al. 2016; Méheust et al. 2020; North et al. 2020; Pan et al. 2022). Inside each group, the bacterial sequences are labeled by dark blue while the archaeal sequences by dark red (fig. 4B). The earliest lineages in Group I (NifD, K, E, and N), Group II (NifD, K, E, and N), and Group III-2 (NifE) are bacterial sequences while archaeal sequences appear as the earliest lineages only in Group III (NifD and K), Group III-3 (NifE), and Group III (NifN). These observations suggest that NifD, K, E, and N all first evolved in bacteria.

It has been proposed that the Vnf/AnfDK were derived from NifDK by gene duplication (Mus et al. 2018; Garcia et al. 2020). This hypothesis is supported by our data, because NifDK and Vnf/AnfDK are found in the same genome (supplementary data S1, Supplementary Material online) and many bacterial species in Group I-DK and Group II-DK also carry Vnf/AnfDK. Moreover, in Group Vnf/Anf bacterial sequences form the earliest lineages and the Vnf/Anf sequences of archaea are nested inside Vnf/Anf sequences of bacteria (fig. 4B and supplementary fig. S2,
Supplementary Material online). These observations suggest that Vnf/AnfDK first evolved in bacteria.

Figure 4A largely supports the NifDK-first hypothesis because Group IV (NifD and K) sequences are the ancestral sequences, as suggested by many previous studies (Raymond et al. 2004; Staples et al. 2007; Howard et al. 2013; North et al. 2020) and because the NifD, K, E, N of Groups III, II, and I were derived from NifD, K of Group IV. Indeed, the lower half of figure 4A suggests that an ancestral K (in Group IV) was duplicated into NifK and NifN, which were then expanded into Groups I, II, and III. The only problem with figure 4A is that Groups I, II, and III NifD sequences are clustered with Groups II and III NifE sequences. Only this part of figure 4A supports the NifEN-first hypothesis, which postulates that NifEN were the ancestral sequences and that NifDK were a duplication of NifEN (GBE 2022; Garcia et al. 2022).

To see why Groups I, II, and III NifD sequences are clustered with Groups II and III NifE sequences in figure 4A, we construct a tree of NifD and NifE sequences (fig. 5A) and a tree of NifK and NifN sequences (fig. 5B), using the full-version maximum likelihood (ML) method (Mega X tool) (Le and Gascuel 2008; Kumar et al. 2018) instead of the IQ-tree method. To reduce the computational load, we select only 215 species (supplementary data S2, Supplementary Material online); we retain most of the archaea species in figure 4A that contain NifDK and eliminate mainly Group I sequences, which are all bacterial. The best-fit model of protein evolution is LG + G. Figure 5A and B shows that Group IV NifD and NifK sequences are ancestral to the other sequences in the tree, in agreement with the NifDK-first hypothesis and with previous studies (Raymond et al. 2004; Boyd, Hamilton, et al. 2011; Mus et al. 2019).

Figure 5A is consistent with the NifDK-first hypothesis, except that a small group of Group III NifE sequences (indicated by *) is clustered with the NifD sequences. This group includes Firmicutes_A, Firmicutes_B, Chloroflexota, and Methanobacteriota, which all lack NifN. A previous study found that the NifE* of *Caldicellulosiruptor* spp. (Firmicutes_A) showed low similarities with other known NifE’s (Chen et al. 2021). To see why this happened, we inferred the ancestral sequence at each branching node of the Nif/Vnf/AnfD, E tree (fig. 5A) and found that the common ancestor of NifE* (node 765) is more similar to NifD than to the other NifE sequences. To illustrate this point, figure 5C shows the alignment of ancestral sequences at 30 amino acid positions, which, according to previous studies (Raymond et al. 2004; Howard et al. 2013; Zheng, Dietrich, et al. 2016; North et al. 2020), are functionally important. Note that sites 188, 191, 277, and 383 of NifE* are the same as all NifD sequences but different from all other NifE sequences. This observation suggests that the substrate coordination and P-cluster ligand of NifE* are more similar to NifD than NifE. Because the NifE* sequences are more similar to NifD sequences than NifE sequences, they were clustered with the Groups III, II, and I NifD sequences in figure 5A.

Figure 5B is completely consistent with the NifDK-first hypothesis as depicted in figure 3. It provides strong evidence against the NifEN-first hypothesis because it clearly indicates that the NifN sequences were derived from a NifK sequence. The NifEN-first hypothesis is contrary to the observations that Group IV NifD and NifK were the ancestral sequences of Groups I, II, and III NifD, K, E, N sequences (fig. 5A and B) and that in both figure 5A and B Group IV includes no NifE and NifN sequences. These two observations were not made by Garcia et al. (2022), because their study included no Group IV NifD and NifK sequences.
Phylogenetic Tree of the Radical SAM Domains of NifB Proteins

Boyd, Anbar, et al. (2011) inferred that the earliest lineages in the phylogeny of the 30 radical SAM domains of NifB proteins were archaeal sequences (Boyd, Anbar, et al. 2011). Here, we reconstruct a phylogeny of 1,712 radical SAM domains of NifB proteins (fig. 6 and supplementary fig. S3, Supplementary Material online); we use 13 MoaA sequences as the outgroup as in Boyd, Anbar, et al. (2011); we use the IQ-tree software (Minh et al. 2020). The best-fit model of protein sequence evolution is LG + R10.

Fig. 4. Phylogeny of 7,421 Nif/Vnf/AnfD, K, E, and N protein sequences. The scale bar denotes 0.5 amino acid substitutions per residue site. The solid and empty black circles denote the bootstrap values of 90–100% (solid circle) and 80–89% (empty circle) and the empty triangle denotes a bootstrap value of 70–79%. The light-independent protochlorophyllide reductase subunit N (Bch/ChlN) and light-independent protochlorophyllide reductase subunit B (Bch/ChlB) are used as the outgroup. The protein sequences are classified into Groups I, II, III, IV, Vnf, and Anf. (A) The outgroup is labeled by black color, NifD by light green, NifK by dark green, NifE by light blue, and NifN by dark blue. (B) is an expanded version of each subtree contained both bacteria and archaea in (A). The bacterial group is labeled by blue color and archaeal groups by dark red color. We use the IQ-TREE method to reconstruct the phylogenetic tree and ModelFinder to select the best-fit model of protein sequence evolution with ultrafast bootstrap (1,000 replicates). The model selected is LG + F + R10.
In figure 6, many of the nodes have low bootstrap values. However, it is clear that the earliest lineages are bacterial sequences (blue color) and the archaeal sequences (red color) are all nested inside bacterial sequences. This result suggests that NifB first evolved in bacteria. Moreover, our result indicates that many NifB proteins of Firmicutes_B species,
which contain only the SAM domain (supplementary fig. S3, Supplementary Material online), form the earliest lineages. A previous study inferred that *Methanococcus* spp. (Methanobacteriota phylum) formed the earliest lineages in the phylogeny of radical SAM domains of NifB proteins (Boyd, Anbar, et al. 2011) and the bacterial species, which

![Phylogeny of the radical SAM domains of 1,712 NifB sequences.](image-url)
contain the fused "SAM–NifX" proteins (the orange small box in supplementary fig. S3, Supplementary Material online), appeared in later lineages. In our species collection, almost every bacterial phylum contains both types of NifB (with only the SAM domain or with the fused "SAM–NifX" protein) as in a previous study (Arragain et al. 2017). Moreover, in our inferred phylogeny, the SAM domain sequences of Methanococcus spp. are nested inside the NifB sequences of bacteria, which contain only the SAM domain (supplementary fig. S3, Supplementary Material online).

Phylogenetic Tree of Concatenated Nif/Anf/VnfHDK Proteins

We concatenate the NifH, D, K sequences (structural components of nitrogenase) because concatenation increases the number of amino acid positions to infer the tree (Boyd and Peters 2013; Boyd et al. 2015; Garcia et al. 2020). We use the ML method with LG + G + I as the model of protein sequence evolution to construct the phylogeny of 326 concatenated Nif/Vnf/AnfHDK sequences (fig. 7). We also construct an IQ-tree using 1,487 concatenated Nif/Vnf/AnfHDK sequences (supplementary fig. S4, Supplementary Material online) and find that the two trees have similar topologies.

In figure 7, Group IV is the oldest group and Groups III, II, and I are younger, as in the NifH tree (figs. 1 and 2) and the NifD, K, E, N trees (figs. 4A, 5A and 8). Note that in each group (I, II, III, IV, Vnf and Anf), bacterial sequences form well-supported earliest lineages while archaeal sequences are nested inside bacterial sequences. Thus, these three nif genes likely first evolved in bacteria. Most of the bacterial species that appear in the earliest lineages belong to Firmicutes_A or Firmicutes_B phyla (fig. 7 and supplementary fig. S4, Supplementary Material online). Moreover, Group IV species and the earliest lineages of Group III Chloroflexota (indicated by #) contain no NifE and NifN, implying that NifDK is older than NifEN. Thus, this tree also supports the NifDK-first hypothesis.

Group I contains only bacteria, so its common ancestor should be a bacterium (Firmicutes_B form the early lineages, supplementary fig. S4, Supplementary Material online). The majority of bacterial species in the earliest lineages of Group II belong to Firmicutes_A or Firmicutes_B (fig. 7 and supplementary fig. S4, Supplementary Material online).

Association of the Mo Transporter With Nitrogen Fixation

We identify the modABC- and wtpABC-gene-harbor species (supplementary data S3, Supplementary Material online) and study their association with nitrogen fixation (i.e., the six-nif-gene set). ModABC is the major Mo transporter in both Nif bacteria and Nif archaea (fig. 8A). Among the 1,420 bacterial species that carry the six-nif-gene set (table 1), 960 (941 + 19) are associated with ModABC whereas only 22 (19 + 3) are associated with WtpABC (see fig. 8A). Similarly, among the 78 archaeal species that carry the six-nif-gene set (table 1), 54 (36 + 18) are associated with ModABC whereas only 21 (3 + 18) are associated with WtpABC (fig. 8A). Moreover, most wtpABC-gene-harbor Nif archaea also harbor modABC genes (18/21). Under the archaea-first hypothesis, the six-nif-gene set first evolved in archaea, so it should be associated more often with wtpABC than with modABC, but the opposite is true in both archaea and bacteria. A recent study suggested that purple sulfur bacteria (Proteobacteria) could fix nitrogen and used ModABC transporter as early as 2.5–0.5 Ga (Philippi et al. 2021). As purple sulfur bacteria are not the oldest Nif species (Group I), there could be even older Nif bacteria that carried the modABC genes. The above observations strongly support the bacteria-first hypothesis.

We reconstruct a phylogeny of 10,119 ModA and 829 WtpA sequences, which represent all available ModA and WtpA sequences in our collection (fig. 8B). ModA (WtpA) is the molybdate-binding protein component of the ModABC (WtpABC) transporter and is the first key protein involved in the absorption of molybdate by prokaryotes. We use the IQ-tree software (Minh et al. 2020) and the best-fit model (Q.Pfam + R7) of protein sequence evolution to construct the tree. Our tree topology is similar to that of a previous tree of 4,623 ModA/WtpA sequences (Ge et al. 2020). In our tree (fig. 8B), the branches are represented by red and black lines, which indicate bootstrap support of ≥95% and <95%, respectively. Thus, the many clades that are preceded by a red branch are supported by a bootstrap value of ≥95% and so are likely reliable. In Circle b, each gray segment represents a group of WtpA sequences. As all of the five gray segments are deeply nested inside the green segments (ModA sequences), a simple explanation is that the WtpA sequences were derived from ModA sequences. A previous study also suggested that WtpA evolved later than ModA (Aguilar-Barajas et al. 2011). In Circle a, each dark red segment represents an archaeal clade. As all of the 13 dark red (archaeal) clades that carry ModA are nested inside the bacterial ModA sequences (blue segments), the archaeal ModA sequences were likely derived from bacterial ModA sequences, probably via horizontal gene transfers (HGTs). Thus, figure 8 supports the bacteria-first hypothesis.

The dispersal of the gray segments on Circle b might have three possible non-exclusive explanations. First, the dispersal might be partly due to tree reconstruction errors. This is, however, unlikely the case for those segments that are far separated on the phylogeny because such separations are supported by very high bootstrap values and so are unlikely to have occurred by chance. Second, some gray segments had evolved independently from green segments. Third, gene conversion had occurred between gray segments. Thus, the many clades that are preceded by a red branch are supported by a bootstrap value of ≥95% and so are likely reliable. In Circle b, each gray segment represents a group of WtpA sequences. As all of the five gray segments are deeply nested inside the green segments (ModA sequences), a simple explanation is that the WtpA sequences were derived from ModA sequences. A previous study also suggested that WtpA evolved later than ModA (Aguilar-Barajas et al. 2011). In Circle a, each dark red segment represents an archaeal clade. As all of the 13 dark red (archaeal) clades that carry ModA are nested inside the bacterial ModA sequences (blue segments), the archaeal ModA sequences were likely derived from bacterial ModA sequences, probably via horizontal gene transfers (HGTs). Thus, figure 8 supports the bacteria-first hypothesis.
**Fig. 7.** The phylogeny of 326 concatenated Nif/Vnf/AnfHDK sequences. The protein sequences are classified into Groups I, II, III, IV, Vnf, and Anf. We use the full-version ML method to construct the tree and ModelFinder to select the best-fit model of protein sequence evolution with bootstrap (500 replicates). The model selected is LG + G + I. The scale bar denotes 0.5 amino acid substitutions per residue site. The solid and empty black circles denote the bootstrap values of 90–100% and 80–89%, respectively, and the empty triangle denotes a bootstrap value of 70–79%. The bacteria are labeled by blue color and the archaea by dark red color. The Vnf/Anf groups contain both bacteria and archaea, labeled by earthy yellow. The four-nif-gene set is labeled by #, the five-nif-gene set by *, and the six-nif-gene set by $. We use 6 concatenated BchXYZ sequences and 10 concatenated Bch/Chl LNB sequences as the outgroup.
Fig. 8. The distribution of ModABC and WtpABC in bacteria and archaea, and the phylogeny of ModA and WtpA sequences. (A) The Venn diagrams of Nif species (red color), modABC-gene-harboring species (gray color) and wtpABC-gene-harboring species (olive green). (B) The phylogeny of 10,119 ModA and 829 WtpA sequences. In Circle a, the bacterial groups are labeled by blue and the archaeal groups by dark red. In Circle b, ModA groups are labeled by green and WtpA groups by gray. In Circle c, the Nif species are labeled by small red squares. The tree is unrooted, but for simplicity of presentation we took one group of ModA sequences as the outgroup. The number of ultrafast bootstrap replicates was 1,000. A red line (branch) means a bootstrap value of \( \geq 95\% \), whereas a black line means a bootstrap value of \(< 95\%\).
others by bacteria suggests that HGTs had occurred between archaea and bacteria. For example, the fourth gray segment contains a subsegment of bacterial WtpA sequences (blue color in Circle a) that lies in between two archaeal gray subsegments (dark red in Circle a), so it was likely derived from archaeal sequences via a HTG.

**Discussion**

In this study, we have used a large number of bacterial and archaeal genomes to conduct an in-depth study of the origin and evolution of nitrogen fixation. We identified many more Nif prokaryotes than did previous studies (Poudel et al. 2018; Albright et al. 2019; Mus et al. 2019; Koirala and Brözel 2021). Our data suggest that many HGT events have occurred between bacteria and archaea but the majority of them were apparently from bacteria to archaea because most archaeal nif genes are phylogenetically nested inside bacterial nif genes (figs. 1, 2 and 4–7). Moreover, we found that while the majority of archaea use WtpABC as the Mo transporter, the majority of Nif archaeal species, like bacteria, use the ModABC as the Mo transporter (fig. 8A). This finding strongly supports the bacteria-first hypothesis. Furthermore, our phylogeny of ModA and WtpA protein sequences shows that the archaeal sequences are nested inside bacterial sequences (fig. 8B), providing further support for the bacteria-first hypothesis.

In constructing the concatenated NifHDK tree, we used only linked HDK sequences, so that the three genes have the same evolutionary history. The NifHDK tree (fig. 7) is congruent with the NifH, NifD and NifK trees (figs. 1, 5A and 8) in that Group IV is older than Groups I, II, and III. It is also congruent with the NifD and NifK trees (fig. 5A and 8) in that Groups I and II are joined together before they are jointed to Group III, that is ([Group I, Group II, Group III), which is the same as in previous studies (Enkh-Amgalan et al. 2006; Boyd and Peters 2013; Garcia et al. 2020; Chen et al. 2021). It is incongruent with the NifH tree in that the NifH tree shows ([Group I, Group III], Group II), which is the same as in previous studies (Angel et al. 2018; Nishihara, Thiel, et al. 2018; Pan et al. 2022). The observation that concatenation of NifH, NifD, and NifK leads to ([Group I, Group II], Group III) suggests that NifD and NifK together provide stronger phylogenetic signals for the branching order of Groups I, II, and III than does NifH. Note that in all of these trees, Group IV is the oldest and that in Group IV the archaeal sequences are all nested inside bacterial sequences, so that no archaeal sequence was likely the ancestral sequence of Groups III, II, and I. As most genomes of the NifHDK tree (fig. 7) are supported by a higher than 90% bootstrap value, the tree appears to be largely reliable. We therefore will include this tree in the following inferences.

The NifH, NifDE, and NifK trees (figs. 1, 5A and 8) suggest that NifH, NifD, and NifK first evolved in bacteria (Group IV). Moreover, the NifB tree (fig. 6) suggests that NifB first evolved in bacteria because bacterial NifB sequences form the earliest lineages. Thus, the four-nif-gene set (nifHDKB) should have first evolved in bacteria. This view is supported by the NifHDK tree (fig. 7) because all Group IV genomes contain NifB as well as NifHDK. Note that *Endomicrobium proavitum*, which is a Group IV-Nfa bacterium, has been experimentally shown to be able to fix nitrogen (Zheng, Dietrich, et al. 2016). Moreover, several studies found that only bacteria can use Group IV-Nfa NifHDKB to fix nitrogen (Zheng, Dietrich, et al. 2016; Méheust et al. 2020; North et al. 2020; Koirala and Brözel 2021). Note further that *Oscillochloris trichoides* has also been shown experimentally to fix nitrogen (Ivanovsky et al. 2021). As *O. trichoides* belongs to Group III (Chloroflexota) while *E. proavitum* belongs to Group IV, there might be other species with the four-nif-gene set (nifHDKB) that can fix nitrogen. However, there seems to be no experimental evidence to show that an archaeon with nifHDKB can fix nitrogen. In our phylogenies of NifH, NifD, K, E, N, and NifHDK (supplementary figs. S1, S2 and S4, Supplementary Material online), Firmicutes_A species or Firmicutes_B species form the earliest lineages in Group IV-Nfa and *E. proavitum* is nested inside Group IV-Nfa sequences of Firmicutes_A (e.g., genus *Clostridium*). Thus, Firmicutes_A might be the first species to use Group IV-Nfa proteins (NifHDKB) for nitrogen fixation.

We now argue that the six-nif-gene set (nifHDKENB) first evolved in bacteria. In the above, we have provided evidence for the NifDK-first hypothesis. According to this hypothesis, NifEN was derived from the duplication of a Group IV NifDK (fig. 3). This NifDK should be in a bacterium because Group IV archaeal NifDK sequences were all nested inside bacterial NifDK sequences, so that no archaeal NifDK could be the ancestral sequence of Groups I, II, and III; that is, there is no branch connecting an archaeon in Group IV to the common ancestor of Groups III, II, and I. Note further that when NifDK was duplicated in a bacterium, that bacterium immediately possessed the six-nif-gene set because before the duplication it already possessed the four-nif-gene set. Thus, we propose that the six-nif-gene set first arose in bacteria.

How the five-nif-gene set (nifHDKENB) arose is less certain. It could have arisen from a duplication of NifD without the duplication of NifK, but then the gene order would have been DEK instead of DKE. A simpler scheme is that the five-nif-gene set was derived from the six-nif-gene set by losing NifN. This view is supported by the following inference. As most Groups I genomes carry the six-nif-gene set, their common ancestor (node 15 in fig. 7) would carry the six-nif-gene set. The same comment applies to the common ancestor of Group II genomes (node 14). Therefore, the common ancestor of Groups I and II (node 13) should also carry the six-nif-gene set. The nif-gene set at the common ancestor (node 4) of Group III is less certain because Group III contains species that carry 4 (denoted by #), 5 (denoted by *) or 6 (denoted by $) nif genes. In particular, the subgroup denoted as Methanobacteriota-1 contains a clade of 4 species carrying nifHDKEB and a clade of 2 species carrying nifHDKENB; we denote these two clades by Methanobacteriota-1-1 and Methanobacteriota-1-2. First,
assume that the common ancestor (node 4) of Group III carried nifHDKB. Then this gene set gained nifEN to become nifHDKENB at node 6 (the assumption of nifHDKEB at node 6 can be shown to be less parsimonious). The nifHDKENB gene set then lost nifN at node 8, at node 10 and in the common ancestor of Methanobacteriota-1-1 to become nifHDKEB at these three nodes. Moreover, this scheme implies that node 3 carried nifHDK, so it requires the gain of nifEN to explain the nifHDKENB at node 13. Thus, this scheme requires two gains of nifEN and three losses of nifN. Second, assume that node 4 carried nifHDKEB (the five gene set). As this scheme requires that nifHDKEB was derived from nifHDK in Group IV, it requires the gain of nifE at node 4 and also a gain of nifEN at node 13 to explain nifHDKENB at node 13. As mentioned above, the gain of nifE from nifD without the simultaneous duplication of nifK would produce the gene order of DEK instead of DKE. Moreover, it requires a gain of nifN in the common ancestor of Methanobacteriota-1-2 and another gain of nifN at node 12 and a loss of nifEN at node 5. Thus, this scheme requires one gain of nifE, one gain of nifEN, two gains of nifN and one loss of nifEN. Therefore, this scheme is complex and not necessarily more parsimonious than the first scheme. Third, assume that node 4 carried nifHDKENB. This scheme requires the loss of nifEN at node 5 and the loss of nifN at node 8, node 10 and in the common ancestor of Methanobacteriota-1-1. Thus, this scheme requires one loss of nifEN, three losses of nifN and also one gain of nifEN (at node 3). Compared to the first scheme, it requires one additional loss of nifEN, but one fewer gain of nifEN. As losing nifEN is simpler than gaining nifEN at a specific location (Albalat and Cañestro 2016), the third scheme is more plausible than the first. In conclusion, it seems that nifHDKEB was derived from nifHDKENB due to a loss of nifN. Note that Caldicellulosiruptor spp, which possesses the five-nif-gene set (nifHDKEB) and has been experimentally shown to fix nitrogen (Chen et al., 2021), is a Firmicutes_A species in Group III.

In conclusion, regardless of whether the four-nif-gene set, the five-nif-gene set or the six-nif-gene is used as the criterion for nitrogen fixation, nitrogen fixation appears to have first evolved in bacteria. Thus, our study strongly favors the bacteria-first hypothesis over the archaea-first hypothesis and the LUCA-first hypothesis. We note that a previous phylogenetic analysis of protein families including nitrogenase did not support the existence of Nif LUCA (Berkemer and McGlynn 2020).

Garcia et al. (2020) proposed the following sequential evolutionary events of the nitrogenase complex formation: 1) NifHDKB, 2) NifHDKEB, 3) NifHDKENB, and 4) Vnf/AnfHDK. Step 1 is supported by our observation that Group IV is the oldest group and most of its members possess the four-nif-gene set (node 1 of fig. 7). However, Step 2 is less certain because as discussed above the five-nif-gene set could have been derived from the six-nif-gene set; in this case, Steps 2 and 3 are reversed. The NifHDK tree suggests that Vnf/AnfHDK (Step 4) was derived from a duplication of NifHDK (fig. 7).

Previous studies suggested that the Group IV-Cfb protein, which is a methane biogenesis nickel-containing tetrapyrrole (known as coenzyme F₄₃₀) (Zheng, Ngo, et al. 2016; Moore et al. 2017), is the common ancestor of both Nif proteins and Bch/Chl proteins (Raymond et al. 2004; Boyd, Hamilton, et al. 2011). This hypothesis was proposed because Group IV-Cfb separated the Nif and Bch/Chl lineages in the phylogeny (Raymond et al. 2004; Boyd, Hamilton, et al. 2011). The “Cfb” is reclassified here as a subgroup of Group IV-CfbC (fig. 2) according to previous studies (Zheng, Ngo, et al. 2016; Moore et al. 2017; North et al. 2020). Because only archaea were found to contain Cfb, the archaea-first hypothesis was proposed (Boyd, Hamilton, et al. 2011; Mus et al. 2019). However, a recent study (North et al. 2020) found that another subgroup of Group IV-Mar (fig. 2) can biosynthesize methionine, ethylene, and methane, which contribute to sulfur and carbon recycling. Moreover, MAR has been found only in bacteria. With the new homologous Group IV protein sequences, our figure 2 suggests that bacterial sequences form the early lineages and archaeal sequences are nested inside bacterial sequences. Moreover, there are many bacteria that contain multiple types of these homologous proteins in their genomes; for example, Rhodospseudomonas palustris CGA009 contains Bch/Chl, IV-Mar, and Nif/Vnf/Anf proteins in its genome (North et al. 2020), implying that these homologous proteins originally evolved in bacteria. As only bacterial species contain Group IV-Nfa and -Mar sequences, this observation implies that the ancient bacteria play an important role not only in the nitrogen cycle but probably also in the carbon or even sulfur cycle.

In our phylogenetic trees, anaerobic firmicutes form the earliest lineages (supplementary figs. S1–S4, Supplementary Material online), and many of them can fix CO₂ through the Wood–Ljungdahl pathway (also termed the acetyl CoA pathway) and possess the sulfate-reducing ability (supplementary data S4, Supplementary Material online). The acetyl CoA pathway has been considered the earliest metabolic pathway on earth (Martin 2020). Moreover, carbon isotope evidence consistent with the presence of the acetyl CoA pathway was found in rocks dated ∼3.95 billion years old (Tashiro et al. 2017). Therefore, we suggest that the earliest Nif species is an anaerobic firmicutes. Consistently, anaerobic firmicutes were found to play a major role in nitrogen fixation in environments analogous to the early earth. For example, thermophilic Caldicellulosiruptor spp. carrying nifH were highly abundant in alkaline sulfidic hot springs in Japan (Nishihara, Thiel, et al. 2018). Furthermore, the nitrogenase activity in an alkaline sulfidic hot spring was not affected by the addition of 2-bromo-ethane sulfonate, a specific inhibitor of methanogenesis, but was apparently inhibited by the addition of molybdate, a specific inhibitor of assimilatory sulfate reduction and sulfur disproportionation (biotransformation of elemental sulfur or thiosulfate into sulfide and sulfate), suggesting that sulfate-reducing/sulfur-metabolizing bacteria, rather than methanogenic archaea, were responsible for nitrogen fixation in the early-earth-like environments (Nishihara, Haruta, et al. 2018).
According to isotopic records, ancient anaerobic Fe-oxidizing bacteria might have evolved around 3.77–4.28 Ga (Dodd et al. 2017; Lepot 2020; Papineau et al. 2022). As Fe-oxidizing bacteria are phylogenetically nested inside Firmicutes (and Nitrospirota, Proteobacteria, Bacteroidota) (Hedrich et al. 2011; Quaiser et al. 2014; Scott et al. 2015), the common ancestor of firmicutes should be older than the Fe-oxidizing bacteria; that is, it is at least 3.77 Ga. It is thus possible that firmicutes had evolved nitrogen fixation very early during the evolution of bacteria.

The isotopic data obtained from pyrite suggested the occurrence of bacterial sulfate reduction around 3.47 billion years ago (Ga) (Reitner and Thiel 2011). From a thermodynamic perspective, the use of sulfate as an electron acceptor allows chemolithotrophic bacteria to utilize H₂ at a threshold level much lower than that in chemolithotrophic methanogenic archaea (Lovley 1985) and to acquire more free energy for ATP generation and nitrogen fixation. Although Archaeoglobus spp. are sulfate-reducing archaea, they are mostly thermophilic aerotolerant anaerobes (Abreu et al. 2000) and are not considered an ancient lineage. Moreover, we did not find any Nif Archaeoglobus spp. as in two previous studies (Dahl et al. 1994; Steinsbu et al. 2010).

Recent studies reported cases of aerobic bacterial methane synthesis from methyl-phosphonate, methylamine, and betaine by cyanobacteria and aerobic proteobacteria (Bižić et al. 2020; Lepot 2020; Wang et al. 2021). Therefore, the 3.48 Ga carbon isotopic data might be from aerobic bacteria rather than archaea (Lepot 2020). Aerobic nitrogen fixation evolved only after Nif bacteria had evolved the defense against oxygen (Boyd et al. 2015; Li et al. 2019). A previous study hypothesized that members of Cyanobacteria (aerobes) evolved nitrogen fixation after archaea (Mus et al. 2019). However, recent findings indicate that the ancestor of Cyanobacteria already existed >3.63 Ga (Garcia-Pichel et al. 2019; Bižić et al. 2020; Ohmoto 2020), which is older than the record of microbial methanogenesis (3.48 Ga). Moreover, cyanobacterial nitrogen fixation can be traced back to 2.7–3.2 Ga (Thomazo et al. 2018; Yang et al. 2019). Since Cyanobacteria is only a phylum in bacteria, aerobic nitrogen fixation in bacteria might have evolved before 2.7–3.2 Ga, which is as old as the estimated minimum age of 2.77 Ga for archaea (Javaux 2019; Lepot 2020). As Nif archaea are a small fraction of archaea, nitrogen fixation could have evolved considerably later than the common ancestor of archaea. From these arguments and those above, we propose that nitrogen fixation evolved first in anaerobic bacteria and later at similar times in aerobic bacteria and archaea.

Identification of nif Genes

In our BLASTp search, we set the e-value threshold = 1e−5, and cutoffs of 30% percent amino acid sequence identity, 70% subject alignment sequence coverage and 70% query sequence alignment, which are more stringent than a previous study (Poudel et al. 2018). The genes collected are listed in supplementary data S1, Supplementary Material online. However, to more accurately identify nifH, nifD, nifK, nifE, nifN, and nifB genes in a genome, we increased the amino acid sequence identity from 30% to 60%. The role of each nif gene in nitrogen fixation has been described in previous studies (Rubio and Ludden 2005; Burén et al. 2020).

Because it is difficult to distinguish among the three types of nitrogenase (nif, vnf, and anf) by structural protein sequences (Hu and Ribbe 2015), we selected the major difference between conventional and alternative nitrogenases as follows. We used the γ-subunit genes (vnfG and anfG) to detect alternative nitrogenases. We also identified the vnfHDK and anfHDK genes by comparing the KEGG and TIGRfams databases of AnnoTree v1.2.0. In some cases, it was difficult to distinguish between vnfHDK and anfHDK genes, and we distinguished them by their clustering with vnfG or anfG in the genome.

Boyd, Anbar, et al. (2011) identified the VnfE and VnfN genes and included them in the phylogeny of NifD, NifK, NifE, and NifN sequences. These genes are named “Nitrogenase_VnfE/N_like,” “nitrogenase associated protein E/N,” or “Nitrogenase molybdenum-iron protein like” in the NCBI database. Therefore, we label these genes as “-like” in figure 4.

Identification of Mo Transporter

To collect the Mo transporter ModABC and WtpABC, we set the E-value threshold = 1e−5, and cutoffs of 60% amino acid sequence identity, 70% subject alignment sequence coverage and 70% query sequence alignment. The 60% identity cutoffs strength is same as a previous study (Ge et al. 2020). These data are also collected from AnnoTree (v1.2.0) (Mendler et al. 2019) and the GTDB (Release R95) database (Parks et al. 2018, 2020).

Materials and Methods

Data Collection

We downloaded the protein sequences from AnnoTree (v1.2.0) (Mendler et al. 2019) and the GTDB (Release R95) database (Parks et al. 2018, 2020). The genomic characteristics and the taxonomic classification of each selected species were downloaded from GTDB (Parks et al. 2020) (supplementary data S1, Supplementary Material online). The nitrogenase genes and other genes were downloaded from the AnnoTree database based on the functional annotation of TIGRfam (v15.0) and the homology information of KEGG (UniRef100) (supplementary data S5, Supplementary Material online). The genes of the Wood–Ljungdahl pathway were identified according to previous studies (Adam et al. 2018, 2019; Jiao et al. 2021) (supplementary data S4, Supplementary Material online).

Phylogenetic Tree Reconstruction

Amino acid sequences were aligned using MAFFT (v7.487) (Katoh and Standley 2013). Pairwise alignments of NifH, NifD, NifK, NifE, and NifN sequences and the radical
SAM domains of NifB protein sequences are listed in supplementary data S6, Supplementary Material online. The pairwise alignments were imported into a MS Excel® spreadsheet (supplementary data S6, Supplementary Material online); the sequences were numbered to correspond to the *Azotobacter vinelandii* proteins; and the key conserved residues of each protein sequence are according to previous studies (Howard et al. 2013; Fay et al. 2015; North et al. 2020).

We use the IQ-TREE tool (Minh et al. 2020) to reconstruct phylogenetic trees and ModelFinder (Kalyaanamoorthy et al. 2017), which is implemented in IQ-TREE, to select the best-fit model of protein sequence evolution with ultrafast bootstrap (1,000 replicates) (Hoang et al. 2018). We also use the MEGA X tool (Kumar et al. 2018) to reconstruct full-version ML (Le and Gascuel 2008) trees and ModelFinder, which is implemented in MEGA X (Kumar et al. 2018), to select the best-fit model of protein sequence evolution with bootstrap (500 replicates). The model is selected according to the BIC (Posada and Buckley 2004). Each phylogeny was annotated by the tool “Interactive tree of life (iTOL) v4” (Letunic and Bork 2019). We classify the nitrogenase genes into different groups by the pairwise sequence alignments and the phylogenetic relationships as in previous studies (Raymond et al. 2004; Howard et al. 2013; North et al. 2020).

**Venn Diagrams**

The area-proportional Venn diagrams in figure 8A are created by the online tool “BioVenn” (Hulsen et al. 2008). We used the GTDB IDs and names of species from supplementary data S1 and S3, Supplementary Material online to create figure 8A. The data collection is mentioned above.

**Supplementary Material**

Supplementary data are available at Molecular Biology and Evolution online.

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**Author Contributions**

H.-W.P., J.-J.L., and W.-H.L. designed the study. H.-W.P. and J.-J.L. collected the data. H.-W.P. and C.-A.C., and J.-J.L. performed the bioinformatic analyses. H.-W.P. wrote the first manuscript draft with the help of W.-H.L. W.-H.L., P.-H.W., and Y.-R.C. advised the study. W.-H.L. supervised the study. All authors participated in revising the manuscript.

**Data Availability**

All our relevant data are provided in Supplementary files of our paper.

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