

Non-cyanobacterial diazotrophs: global diversity, distribution, ecophysiology, and activity in marine waters

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Abstract

Biological dinitrogen (N₂) fixation supplies nitrogen to the oceans, supporting primary productivity, and is carried out by some bacteria and archaea referred to as diazotrophs. Cyanobacteria are conventionally considered to be the major contributors to marine N₂ fixation, but non-cyanobacterial diazotrophs (NCDs) have been shown to be distributed throughout ocean ecosystems. However, the biogeochemical significance of marine NCDs has not been demonstrated. This review synthesizes multiple datasets, drawing from cultivation-independent molecular techniques and data from extensive oceanic expeditions, to provide a comprehensive view into the diversity, biogeography, ecophysiology, and activity of marine NCDs. A NCD *nifH* gene catalog was compiled containing sequences from both PCR-based and PCR-free methods, identifying taxa for future studies. NCD abundances from a novel database of NCD *nifH*-based abundances were colocalized with environmental data, unveiling distinct distributions and environmental drivers of individual taxa. Mechanisms that NCDs may use to fuel and regulate N₂ fixation in response to oxygen and fixed nitrogen availability are discussed, based on a metabolic analysis of recently available Tara Oceans expedition data. The integration of multiple datasets provides a new perspective that enhances understanding of the biology, ecology, and biogeography of marine NCDs and provides tools and directions for future research.

Keywords: diazotrophs; marine nitrogen cycle; nitrogen fixation; non-cyanobacterial diazotrophs

Introduction

Biological dinitrogen (N₂) fixation is the microbial process, whereby otherwise inert N₂ gas is reduced to biologically available nitrogen (N) in the form of ammonia (NH₃). In the oceans, the N supplied through this process can support primary productivity, especially in N-limited surface waters and, consequently, the vertical transport (export) of carbon (C) to the deep ocean (Karl et al. 1997, Knapp et al. 2018, Zehr and Capone 2021). More broadly, the balance between N inputs from N₂ fixation and losses from denitrification and anaerobic ammonium (NH₄⁺) oxidation set the oceanic inventory of reactive N, impacting the global C cycle and Earth's climate system (Gruber and Galloway 2008). Rates of N₂ fixation in the oceans have strong biogeographical patterns, with the highest rates measured in surface open-ocean and coastal waters of the tropics, subtropics, and temperate zones (e.g. Capone et al. 2005, Berthelot et al. 2017, Tang et al. 2019b), and lower rates in polar regions (Blais et al. 2012, Sipler et al. 2017, Shiozaki et al. 2020) and the deep sea (reviewed by Moisander et al. 2017, Benavides et al. 2018a). Marine N₂ fixation has been best described in epipelagic environments, where rates are generally limited by temperature, nutrient [phosphorus (P) and iron (Fe)] concentra-

tions, and N:P and/or N:Fe ratios (Mills et al. 2004, Moore et al. 2009, Letelier et al. 2019, Tang et al. 2019a).

While the importance of N₂ fixation to marine biogeochemical cycling is indisputable, there is still uncertainty concerning which N₂-fixing microorganisms (diazotrophs) contribute to this process in marine euphotic waters. Early microscopy-based analyses established the importance of filamentous cyanobacterial diazotrophs (*Trichodesmium* and heterocyst-forming symbionts of diatoms) in tropical and subtropical surface waters (e.g. Dugdale et al. 1961, Mague et al. 1974, Venrick 1974, Carpenter and Price 1977, Saino and Hatori 1980). More recently, marine diazotrophic diversity has been investigated using molecular approaches targeting the *nifH* gene, which encodes the two identical subunits of the Fe protein of the nitrogenase enzyme complex and serves as a phylogenetic marker (Zehr and McReynolds 1989, Gaby and Buckley 2011). These approaches led to the discovery of diverse cyanobacterial diazotrophs in the open ocean, as well as diazotrophs from many lineages not within the phylum Cyanobacteria, which we refer to as non-cyanobacterial diazotrophs (NCDs; see Box 1; Zehr et al. 1998). Subsequent ocean surveys showed that *nifH* genes from NCDs are ubiquitous in marine waters and can reach higher

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relative abundances than their cyanobacterial counterparts (Riemann et al. 2010, Farnelid et al. 2011). However, very few marine NCDs have been cultivated and the few isolates that exist have not provided unequivocal evidence of N_2 fixation *in situ* (Bostrom et al. 2007, Farnelid et al. 2014, Bentzon-Tilia et al. 2015a, Martinez-Perez et al. 2018). To date, direct demonstrations of NCD N_2 fixation *in situ* using cultivation-independent approaches are extremely limited. Thus, the biogeochemical significance of NCDs remains unknown.

Box 1. The case for using the term ‘non-cyanobacterial diazotroph’

This review focuses on microorganisms referred to collectively as non-cyanobacterial diazotrophs, or NCDs. This term was previously defined by Moisander et al. (2017) and includes all diazotrophic bacteria and archaea not part of the phylum Cyanobacteria. We argue that the term NCD is more accurate than other terms used in the marine literature, such as ‘heterotrophic diazotrophs’ and ‘heterotrophic bacterial diazotrophs.’

Diazotrophic diversity is typically characterized by sequencing the *nifH* gene, which largely has congruence with 16S rRNA gene-based phylogeny (Zehr et al. 2003). This congruence is strong for cyanobacteria, such that if unknown sequences group within *nifH* Cluster 1B, they are phylogenetically distinguished as cyanobacteria. Most of these organisms are photoautotrophs, capable of using light as a source of energy and carbon dioxide as a C source, though there are some notable exceptions found among unicellular cyanobacterial symbionts of haptophytes and diatoms (Tripp et al. 2010, Nakayama and Inagaki 2017).

Unfortunately, *nifH* gene sequences are not easily used to predict the metabolic strategy a particular NCD may use to acquire energy and C. Although there is some congruence between *nifH*- and 16S rRNA gene-based phylogenies for non-cyanobacteria, there are exceptions for important marine phyla, including the proteobacteria, which are spread across multiple *nifH* clusters (Table S1, Supporting Information). More broadly, it can be challenging to infer metabolic potential from a single gene. Even well-established phylogenetic markers like the 16S rRNA gene are difficult to use for resolving species- and strain-level identities (Johnson et al. 2019), which are needed to infer metabolic strategies from genera containing high metabolic diversity, such as the γ -proteobacterial genus *Pseudomonas* (Jun et al. 2016).

Since *nifH* gene sequences cannot be used to accurately infer the metabolic potential of all NCDs, the terms ‘heterotrophic bacterial diazotrophs (HBDs)’ (Delmont et al. 2018) or ‘heterotrophic diazotrophs’ (Gradoville et al. 2017a) could potentially be inaccurate. For example, γ -proteobacterial diazotrophs within Cluster 1G include cultivated microorganisms with divergent metabolic strategies, e.g. *M. purpuratum*, which uses light as a source of energy and organic C as the source of cell C (photoheterotrophy), some strains of *P. stutzeri*, which are facultative anaerobes that oxidize inorganic compounds as a source of energy for growth (chemolithotrophy), and *A. vinelandii* which are aerobes that use organic C as their major source of energy and C (chemoheterotrophy). Furthermore, NCDs include archaea, and are thus not all bacterial.

Ideally, NCDs should be subclassified according to their potential metabolic traits. This will become more feasible with advances the isolation and genome sequencing of marine NCDs (see ‘Diversity and ecophysiological features inferred from MAGs’). Here, we decided to use the term NCD as an accurate and inclusive way to refer to this group without implying a particular metabolic strategy.

Much of the ambiguity concerning the significance of marine NCDs stems from uncertainties about their ecophysiology, particularly in well-lit, oxygenated marine habitats. There are several major challenges that marine NCDs must overcome (reviewed by Bombar et al. 2016). N_2 fixation is energetically expensive, with high ATP and reductant requirements (Postgate 1982). Cyanobacterial diazotrophs can produce organic C and acquire energy through oxygenic photosynthesis, although there are some exceptions such as the photoheterotrophic symbiont UCYN-A (Tripp et al. 2010). In contrast, marine NCDs are thought to utilize organic substrates and/or alternative energy sources (including possibly light) to meet energy and C requirements. Similar to model NCDs from terrestrial systems (Dixon and Kahn 2004), various energy acquisition pathways have been observed in marine NCD genomes, which are speculated to be important strategies to fuel growth and N_2 fixation activity (Bentzon-Tilia et al. 2015a, Martinez-Perez et al. 2018, Acinas et al. 2021). Many marine NCDs are presumed to be chemoheterotrophic, thus may have difficulty acquiring sufficient energy for N_2 fixation in euphotic open-ocean waters, which are often depleted in labile organic C. Indeed, amendment experiments suggest that the abundances and/or N_2 fixation activity of NCDs are limited by the availability of dissolved organic C (DOC) in many pelagic marine environments (Moisander et al. 2012, Dekaezemacker et al. 2013, Rahav et al. 2016).

A major challenge facing NCDs (and all diazotrophs) is the irreversible inactivation of the nitrogenase enzyme by oxygen (O_2 ; Gallon 1992). The ecological advantage of N_2 fixation is hindered by the costs of cellular strategies needed to protect nitrogenase from O_2 . Cyanobacterial diazotrophs have evolved numerous mechanisms to separate N_2 fixation from O_2 either in space or in time (or both), including cellular differentiation, symbiosis, membranes to restrict O_2 diffusion, and increased rates of cellular respiration (reviewed by Zehr and Capone 2020). These mechanisms have energetic costs that would further exacerbate the C requirements for NCDs. Particles have been proposed as habitats for NCD N_2 fixation, since they can provide both a C source and the potential for low O_2 microzones (Paerl and Prufert 1987, Riemann et al. 2010, 2022, Bombar et al. 2016). However, active transcription of NCD *nifH* genes is also found in free-living size fractions (Salazar et al. 2019) and on particles too small to theoretically support low O_2 conditions (Farnelid et al. 2019). In the terrestrial environment, free-living *Azotobacter vinelandii* fixes N_2 under aerobic conditions through increased respiration to consume intracellular O_2 and by investing in chemical barriers (like exopolysaccharides) and larger cell sizes to reduce O_2 intrusion into the cell (Post et al. 1982, Poole and Hill 1997, Sabra et al. 2000). It is likely that marine NCDs inhabiting oxic environments may use similar strategies.

The presence of nitrate (NO_3^-) and NH_4^+ is generally thought to inhibit N_2 fixation, either by supporting the growth of fast-growing phytoplankton that outcompete diazotrophs for other limiting nutrients (Ward et al. 2013), or by providing fixed N that some diazotrophs can utilize instead of investing in N_2 fixation (although inhibition is variable, see Knapp 2012). However, fixed inorganic N is abundant in subeuphotic zone waters and coastal or upwelling ecosystems where N_2 fixation by NCDs has been implicated (reviewed by Moisander et al. 2017), as well as in sediment systems where N_2 fixation occurs (Capone 1983), underscoring that the sensitivity of NCDs to fixed N is not well-understood. Furthermore, in low O_2 environments, N_2 fixation in the presence of high NH_4^+ has been speculated to drive the balance of internal cellular redox states in *Rhodospseudomonas palustris* (isolate BAL398)

a model NCD from the Baltic Sea (reviewed by Bombar et al. 2016). However, it is unknown whether some marine NCDs may use N_2 fixation as a redox balancing strategy in low O_2 environments (such as on particles).

Despite these challenges, NCDs appear to be widely distributed and occasionally transcriptionally active in marine euphotic waters. Measuring the activity of NCDs and quantifying their importance to marine N_2 fixation is an active area of research. Fortunately, the past decade has seen a significant improvement in the tools and data available to study marine NCDs. High throughput sequencing (HTS) technologies and increased sampling efforts have exponentially increased the amount of *nifH* amplicon sequence data available for marine diazotrophs (e.g. Farnelid et al. 2011, Turk-Kubo et al. 2015, Shiozaki et al. 2017, Raes et al. 2020). Dozens of individual NCD *nifH* gene sequence types have been enumerated using quantitative PCR (qPCR), resulting in large datasets showing regional and seasonal abundance patterns (e.g. Langlois et al. 2015, Shiozaki et al. 2017, Cheung et al. 2021, Shao and Luo 2022). Furthermore, advances in sequencing technologies have enabled the detection of *nifH* genes from global ocean metagenomes and metatranscriptomes (Delmont et al. 2018, 2022, Salazar et al. 2019, Acinas et al. 2021, Pierella Karlusich et al. 2021, Poff et al. 2021), allowing for the validation of NCD distribution patterns without relying on the use of primers for the PCR-based amplification of NCD *nifH* gene sequences, and for the construction of metagenome-assembled genomes (MAGs) to evaluate the metabolic potential of uncultivated NCDs. Finally, advances in single-cell techniques are beginning to enable visualization and *in situ* single-cell N_2 fixation rate measurements of NCDs (e.g. Martinez-Perez et al. 2018, Geisler et al. 2019).

This review synthesizes these new data to better understand the role of NCDs in marine waters. In 'NCD diversity: *nifH* gene catalog,' we introduce a novel catalog of marine NCD gene diversity that compiles available *nifH* sequence data from the NCBI Genbank non-redundant (nr) database and selected *nifH* HTS studies, qPCR targets, and global ocean metagenomes. 'Habitats and environments of NCDs in marine systems' reviews the known environments and habitats for NCDs in marine systems. In 'Environmental drivers of NCD biogeography, activity, and presumed N_2 fixation,' we discuss known environmental controls on their abundances and activity, introduce a novel database of available NCD *nifH* gene abundances, and discuss the global distributions and environmental drivers of individual phylotypes. 'NCD biogeography and ecophysiology from metagenomes and metatranscriptomes' explores the potential ecophysiological features of marine NCDs using newly available MAGs from the global ocean (Delmont et al. 2022). In 'Moving from genes to rates: are NCDs fixing N_2 in the pelagic oceans?,' we consider the current state of knowledge on NCD N_2 fixation activity and in 'Future perspectives' we discuss remaining knowledge gaps and future perspectives for assessing the contribution of NCDs to N_2 fixation in marine waters.

NCD diversity: *nifH* gene catalog

Knowledge of NCD diversity is based on the nucleic acid sequences of the genes that encode the nitrogenase enzyme. Nitrogenase is composed of dinitrogenase reductase and dinitrogenase, which are the Fe and molybdenum (Mo)-Fe (MoFe) containing metalloproteins, respectively, in the most common form. The MoFe protein-containing nitrogenase is encoded by the *nifHDK* operon; *nifDK* encodes the dinitrogenase alpha and beta subunits (containing MoFe) and *nifH* encodes dinitrogenase reductase (containing Fe). Less common nitrogenases substitute Fe or vanadium

for Mo and are encoded by the *vnfHDKG* and *anfHDKG* genes, respectively (Seefeldt et al. 2009, Newton 2015). N_2 fixation requires the involvement of many proteins and factors beyond nitrogenase, thus is a highly regulated and complex process ultimately set by intracellular N status and O_2 , Mo, and Fe concentrations, as well as available energy sources (Dixon and Kahn 2004, Leigh and Dodsworth 2007, Masepohl 2017).

The *nifH* gene is the most widely used proxy for N_2 fixation potential in the marine environment. While lateral gene transfer has been observed in some taxa (Bolhuis et al. 2010), *nifH* gene-based phylogeny is broadly congruent with 16S rRNA gene-based phylogeny and four major *nifH* gene clusters have been defined (Zehr et al. 2003; Table S1, Supporting Information). Early application of 'universal' *nifH* PCR assays using pelagic marine samples by Zehr et al. (1998) established that there were multiple lineages of marine unicellular cyanobacteria along with α -, β -, and γ -proteobacteria, sulfate reducers (δ -proteobacteria), and *Clostridia* among the picoplankton population. Over the past quarter-century, additional diazotrophic lineages have been identified, including marine representatives from each of the four major *nifH* clusters. Applications of new molecular techniques, including HTS of *nifH* amplicons and metagenomic-/transcriptomic-based approaches, have greatly increased the known *nifH*-based diversity of marine diazotrophs; however, most individual studies have focused on cyanobacterial diazotrophs, and no comprehensive compilations of the currently known NCD diversity exist.

We compiled a catalog of marine NCD *nifH* gene sequences (Table S2, Supporting Information; dataset doi: 10.5281/zenodo.6537451). This catalog is a new community resource that enables analysis of *nifH* datasets in the context of prior studies and identifies NCD sequences that have been recovered using different approaches. The NCD *nifH* gene catalog includes sequences recovered from: (1) studies using *nifH* PCR and cloning-based approaches (archived in the NCBI nr database); (2) select studies employing HTS of *nifH* PCR amplicons; (3) metagenomic/transcriptomic datasets from recent large-scale ocean surveys; and (4) targets for qPCR and digital droplet PCR (ddPCR) quantitative methods.

Briefly, marine-derived sequences from the NCBI nr database were selected from a curated *nifH* database (Heller et al. 2014 updated in June 2017) using available metadata. All marine nr-derived sequences ($n = 23\ 848$) were clustered at 97% amino acid identity using CD-HIT (Huang et al. 2010), and NCD operational taxonomic units (OTUs) with <100 sequences were removed. This resulted in 34 OTUs (representing 9360/23 848 of the total marine-derived sequences). Additionally, sequences representing the top three most abundant NCD OTUs (or amplicon sequence variants, ASVs) from *nifH* HTS gene studies that sampled the North Pacific (Shiozaki et al. 2017, Farnelid et al. 2019, Cabello et al. 2020, Gradoville et al. 2020, Sato et al. 2021, Turk-Kubo et al. 2021), South Pacific (Turk-Kubo et al. 2015), South Atlantic (Ribeiro et al. 2018), Indian Ocean (Wu et al. 2021), and polar regions (Shiozaki et al. 2018b, 2020) were also included. These studies were selected based on the accessibility of reference sequences for reported NCDs; sequences have been renamed according to the region, author, and specified name from the original publication (e.g. 'Npac-Shio-otu00004' for otu00004 from the North Pacific in Shiozaki et al. 2017; Table S1, Supporting Information). The catalog also includes NCD *nifH* sequences obtained using PCR-free approaches from Tara Oceans metagenomes, metatranscriptomes, and MAGs (Delmont et al. 2018, 2022, Salazar et al. 2019, Cornejo-Castillo and Zehr 2021, Pierella Karlusich et al. 2021) and the Polar Microbe R Gene Catalog (PM-RGC; Cao et al. 2020). PM-RGC *nifH* sequences

were identified via blastp against a curated database containing genome-derived *nifH* sequences (jzehrlab.com/nifh). Finally, the catalog includes 55 reference sequences from NCDs targeted by published *nifH* qPCR/ddPCR assays, since these were identified as recurrently present diazotrophs from independent studies.

Together, the NCD *nifH* gene catalog contains 204 total sequences (Table S2, Supporting Information; dataset doi: 10.5281/zenodo.6537451). OTUs comprised of sequences with >99% amino acid identity were identified using CD-HIT (Huang et al. 2010) to explore which sequence types have been recovered using different methodological approaches. OTUs containing sequence types detected using both *nifH* PCR-based and PCR-free (derived from metagenomics/metatranscriptomics) methods are of particular interest, some of which are discussed below.

The Gamma A and Gamma 4 OTUs (1G) contain sequences from both PCR and non-PCR-based approaches (Fig. 1) and are arguably the best-studied marine NCD groups. The widespread distribution of Gamma A throughout tropical and subtropical surface waters has been demonstrated in qPCR-based studies (see 'Environmental drivers of NCD biogeography, activity, and presumed N₂ fixation'), *nifH* gene HTS studies (e.g. Npac-Shio-otu00004 from Shiozaki et al. 2017, SATl_Ribe_otu006 from Ribeiro et al. 2018), and in a metagenome-based study (Cornejo-Castillo and Zehr 2021). Gamma A *nifH* transcripts have been detected in the environment (e.g. Bird et al. 2005, Moisander et al. 2014, Cornejo-Castillo and Zehr 2021) and this organism is hypothesized to be associated with small particles or picophytoplankton (Benavides et al. 2016b, Cornejo-Castillo and Zehr 2021; Fig. 2a). Gamma 4 is another γ -proteobacterium emerging as a potentially important marine NCD. Originally described as a qPCR target by Halm et al. (2012), and later observed in the Eastern Tropical South Pacific (P8; Löscher et al. 2014), this group now includes a MAG assembled from North Pacific samples collected during the Tara Oceans expedition (TARA_PON_109_MAG_00010, also referred to as HBD-06; Delmont et al. 2018, 2022) and can reach high abundances (5.8×10^6 *nifH* gene copies l⁻¹) in the North Pacific (Cheung et al. 2021).

Other potentially important γ -proteobacterial OTUs emerged from this meta-analysis. γ -ETSP1 was first described as a qPCR target in the South Pacific (Turk-Kubo et al. 2014), but also includes a MAG (TARA_AOS_82_MAG_00008), which was detected mainly in the Atlantic Ocean in <5 μ m size fractions (Delmont et al. 2022). In total, three additional γ -proteobacterial OTUs contained both MAGs and abundant sequence types from *nifH* gene HTS studies: TARA_PON_109_MAG_00047 included a sequence from Shiozaki et al. (2017), and TARA_PSE_93_MAG_00126 and OM-RGC.v2.009033127 included sequences from Gradoville et al. (2020; Table S2, Supporting Information). The recovery of numerous γ -proteobacterial sequences from PCR-based studies is notable, given recent assertions that primer mismatches could result in lack of amplification of this group (Delmont et al. 2018, 2022). Our analysis reinforces that γ -proteobacteria are recovered by both approaches and are among the most diverse NCD groups in the marine pelagic environment.

Several OTUs affiliated with *nifH* cluster 1J/1K were also found in both PCR-based and PCR-free studies (Fig. 1). The α -24809A06 OTU contains sequences related to putative α -proteobacteria reported in the South China Sea (SCS; α -24809A06; Moisander et al. 2007) and mesopelagic waters (Azosp_1; Hewson et al. 2007), and also includes a MAG (TARA_PSW_86_MAG_00238; Delmont et al. 2022) detected predominantly in the Pacific. A second putative α -proteobacterial OTU, Alpha 2, contained a sequence type targeted

by qPCR in the Northern SCS (Alpha 2; Chen et al. 2019) and a MAG (TARA_AON_82_MAG_00070; Delmont et al. 2022) observed in the <20 μ m size fraction in several ocean basins including the Atlantic, Indian, and Pacific.

Unlike the proteobacteria, no putative δ -proteobacterial OTUs (cluster 3 or 1A) were found using both PCR-based and PCR-free approaches. Cluster 3 contains sequences from PCR-based studies and from MAGs including Planctomycetes lineages speculated to be important NCDs (Delmont et al. 2018, 2022). However, sequences affiliated with cluster 1A were recovered only by PCR-based techniques. In this dataset, cluster 1A sequences are predominant in coastal ecosystems (e.g. Short and Zehr 2007, Shiozaki et al. 2018b) that were not heavily sampled in the Tara Oceans expedition, which may partially explain this discrepancy.

Caveats and considerations for *nifH* sequence analysis

Although PCR-based approaches have been foundational in the study of marine diazotrophs, there are several challenges and caveats associated with using *nifH* sequences to explore the diversity of diazotrophs. Numerous sets of primers have been developed to amplify *nifH* genes, many of which have a high degree of degeneracy and are biased towards certain taxa (Gaby and Buckley 2011). The most widely applied assay in the marine environment (*nifH*1-4; Zehr and McReynolds 1989, Zani et al. 2000) has known biases, especially within the cyanobacteria (Caputo et al. 2018). However, recent assertions that *nifH*1-4 primers may not amplify some of the NCDs recovered using 'omics-based approaches, e.g. *nifH* genes from Planctomycetes (Delmont et al. 2018, 2022), are primarily based on a single mismatch against the *nifH*4 primer located at the 5' end of the priming site (Table S2, Supporting Information), which does not impact PCR amplification (Bru et al. 2008). Our NCD gene catalog shows that several MAG-derived sequences asserted to have incompatibilities with the *nifH*1-4 primers have been recovered from PCR-based surveys, underscoring that the 5' mismatch does not prevent PCR amplification (Table S3, Supporting Information). Therefore, the *nifH*1-4 primers do not appear to be broadly incompatible with pelagic marine NCD taxa.

There are several additional technical limitations of PCR-based approaches. Due to the generally low abundances of diazotrophs, amplification typically requires a nested PCR approach and many rounds of amplification, which introduces bias in relative abundances (Turk et al. 2011). Recovery of *nifH* gene fragments can also be influenced by contamination from numerous sources; differentiating between marine- and contaminant-sourced *nifH* genes can be challenging, particularly for NCDs (Zehr et al. 2003). This has occasionally been addressed in individual studies by processing reagent blanks or comparing data to known contaminant sequences (Bostrom et al. 2007, Farnelid et al. 2011, Blais et al. 2012, Moisander et al. 2014, Langlois et al. 2015, Fernandez-Mendez et al. 2016, Cheung et al. 2021), so that putative contaminant sequences can be removed during analysis.

Detection of *nifH* transcripts in environmental samples is sometimes used as a proxy for active N₂ fixation, and in early studies was the standard for choosing targets for qPCR assays. However, detection of *nifH* transcripts may be heavily dependent on the time of sampling due to the diel pattern in *nifH* gene expression (observed in diazotrophs including marine cyanobacteria and terrestrial NCDs; Wyman et al. 1996, You et al. 2005), size fraction (which can reflect lifestyle, e.g. symbiont, particle-attached), and environmental controls (e.g. O₂ concentration, presence of re-

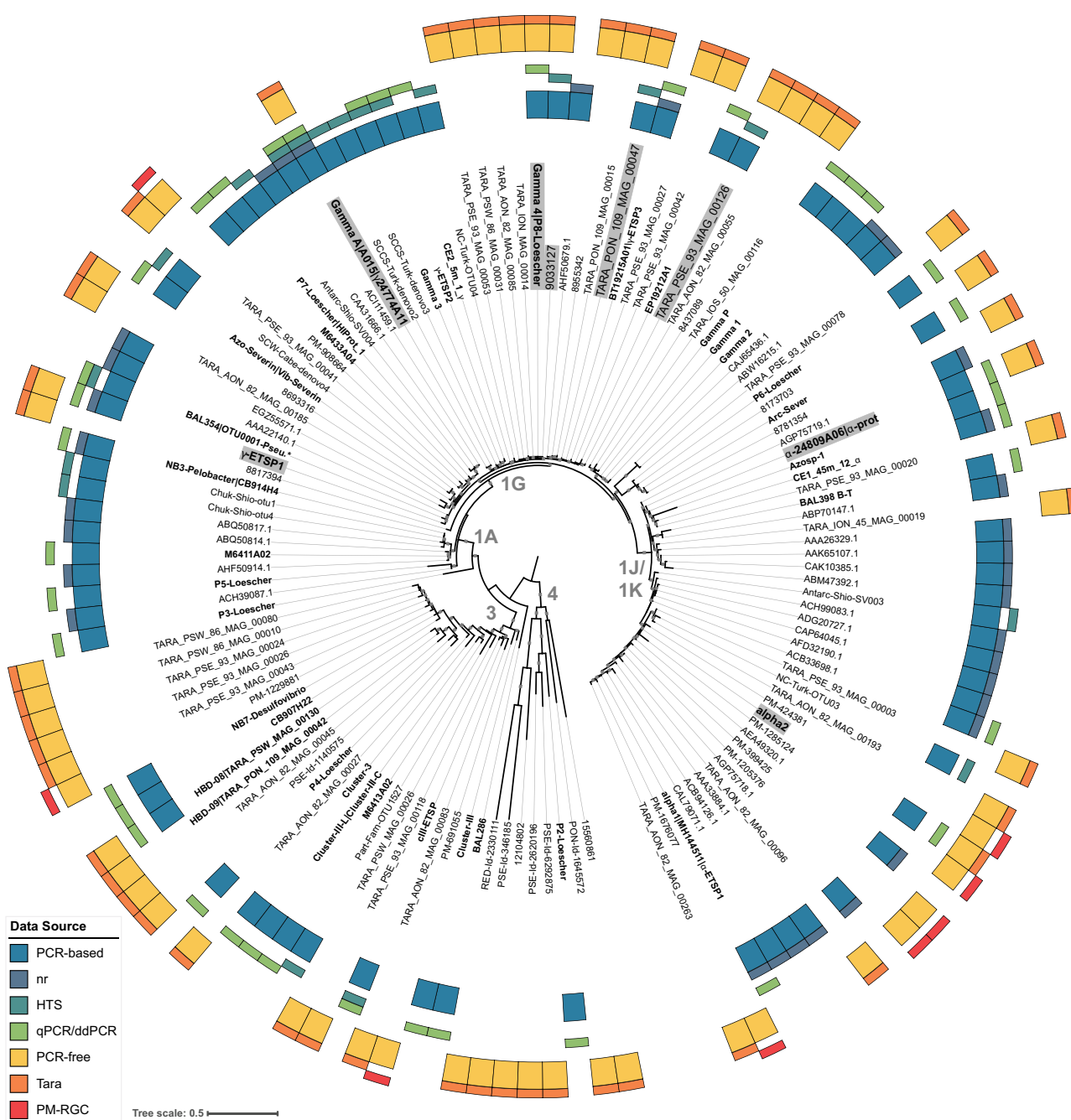


Figure 1. NCD diversity includes taxa found using PCR-based and PCR-free approaches. Phylogenetic tree represents the NCD *nifH* gene catalog based on amino acid sequences from OTU representatives. Sequences were aligned to the NifH/frxC family (Fer4_NifH; PF00142) using HMMalign in HMMER software v2.4 (Finn et al. 2011). Tree topology was calculated using FastTree 2.1.11 (Price et al. 2010) using maximum-likelihood rearrangements and the JTT model for nucleotide evolution. Branch support was determined using the Shimodaira–Hasegawa test (>50% support indicated with small gray squares on branches). iTOL 6.5.2 (Letunic and Bork 2021) was used to visualize the tree and display the source(s) of sequences in each cluster. *nifH* clusters are defined according to the convention established in Zehr et al. (2003) and are indicated in gray text in the center of the tree. Representative sequences affiliated with NCD qPCR/ddPCR assays are in bold. OTUs that contain sequences derived from both PCR-based and PCR-free approaches are in shaded boxes. Branches with multiple names indicate OTUs that contain sequences targeted by more than one qPCR assay. Thick outer color bars show if the sequences were acquired through PCR-based (teal) or PCR-free (yellow) methods, while the thinner color bars correspond to the specific method (PCR-based) or sampling campaign (PCR-free). Interactive tree publicly available at <https://itol.embl.de/shared/1HrZrblPr7p4s>. See Table S2 (Supporting Information) for supporting data.

duced N, and availability of organic C and Fe). Furthermore, while the detection of *nifH* transcripts may reflect active transcription of the *nif* operon, active N₂ fixation is also under post-transcriptional control. Therefore, *nifH* transcript detection (or lack thereof) as evidence of active/inactive N₂ fixation should be interpreted with caution.

Recovering *nifH* gene sequences from the environment, regardless of the approach, does not guarantee that they are sourced from organisms capable of fixing N₂ as some taxa contain *nifH* genes but lack other genes required for a functional nitrogenase (Dos Santos et al. 2012, Mise et al. 2021). Furthermore, alternative nitrogenases in marine diazotrophs are not well-understood,

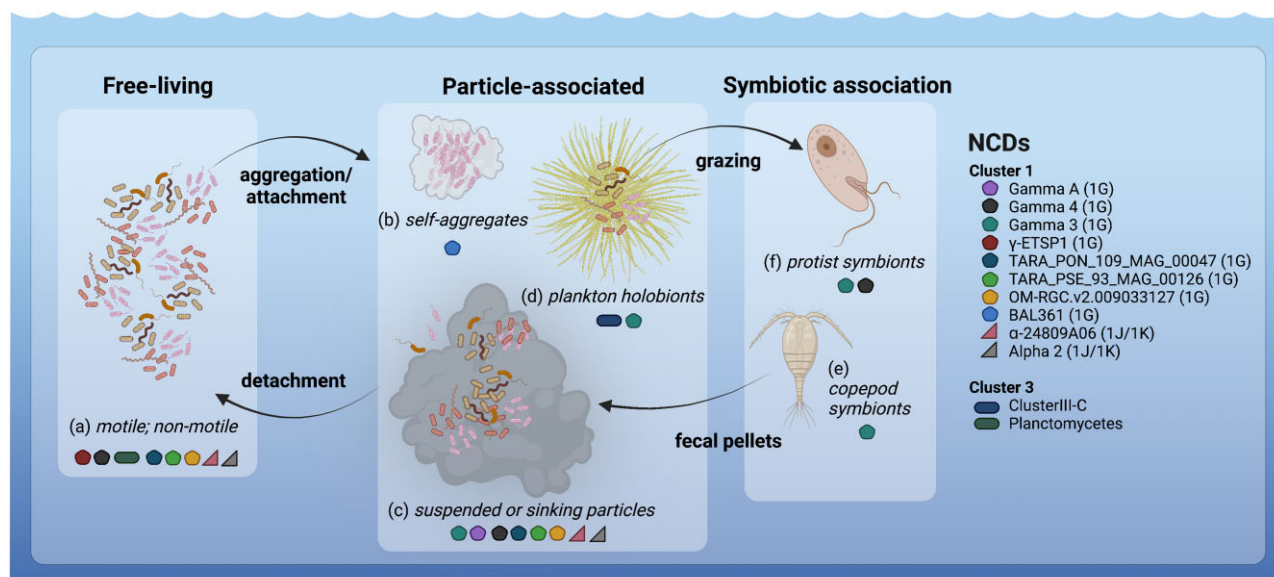


Figure 2. NCDs occupy diverse marine habitats. NCDs in marine waters may be free-living and motile or nonmotile (a), associated with various particles including self-aggregates (b), suspended or sinking particles (c), plankton holobionts (d), or live in symbiosis with copepods (e) or other protists (f). Presumed habitats of some NCD taxa which are discussed throughout this review are indicated using symbols described in the legend. Created with BioRender.com.

but also signify the potential to fix N_2 (McRose et al. 2017, Reeder and Löscher 2022), which is not captured in environmental *nifH* gene surveys. Nevertheless, approaches targeting partial *nifH* gene sequences have generated the majority of available marine NCD data and have enabled numerous insights into their diversity, distribution, and environmental drivers.

Habitats and environments of NCDs in marine systems

Marine euphotic waters are heterogeneous environments fostering free-living NCDs, NCDs attached to living or detrital particulate material, and NCDs likely living in symbiosis with protists (Fig. 2). These contrasting habitats can be found in close proximity, thus the scale of standard oceanographic sampling (typically milliliters–liters) can conceal complex life strategies and interactions. For example, some NCD taxa likely cycle through free-living (Fig. 2a) and particle-associated lifestyles (Fig. 2b–d), evidenced by their occurrence across size fractions (Pierella Karlusich et al. 2021) as well as their potential for motility (Hallstrøm et al. 2022c) and the formation of self-aggregates (Bentzon-Tilia et al. 2015a). Other taxa may form facultative or obligate symbioses with other planktonic taxa (Fig. 2f and e). Rates and magnitudes of NCD N_2 fixation in oxygenated euphotic waters are likely influenced by these different lifestyles. For example, N_2 fixation by free-living or particle-associated NCDs would be dependent on a given taxa's metabolic potential and ability to respond to a changing environment, while N_2 fixation by NCDs living in symbiosis may be under host control like in other marine and terrestrial symbioses (Smercina et al. 2019, Landa et al. 2021, Mohr et al. 2021). Teasing apart the habitats and lifestyles of NCDs in specific environments remains a challenge but is key to linking diversity to active N_2 fixation.

NCDs are also present in many marine environments beyond the euphotic zone (Table 1). Since the first reports of NCDs in the surface ocean (Paulsen et al. 1991, Coyer et al. 1996, Zehr et al. 1998), technological advances and ambitious sampling cam-

paigns have highlighted a diversity of water column environments harboring NCDs, including coastal waters and estuaries, polar seas, aphotic environments, and O_2 minimum zones (OMZs). Diverse benthic ecosystems also harbor NCDs and may seed them into the water column, where they become part of the rare biosphere (Pedros-Alio 2012, Troussellier et al. 2017). Here, we review known water column NCD habitats and environments ('Habitats and environments of NCDs in marine systems') before discussing global NCD distributions and environmental drivers ('Environmental drivers of NCD biogeography, activity, and presumed N_2 fixation').

Marine particles

Although little is known about the physiology of NCDs and how they fix N_2 in oxygenated surface waters, for over two decades it has been theorized that NCDs residing in sunlit surface waters have a particle- or aggregate-associated lifestyle (Fig. 2b–d; Paerl and Prufert 1987, Church et al. 2005a, Riemann et al. 2010, Bombar et al. 2013, Rahav et al. 2016, Pedersen et al. 2018). Marine particles are an amalgamation of living and detrital material formed throughout the water column, with highest concentrations in the euphotic zone, especially in areas of high productivity, such as coastal and upwelling regions (Simon et al. 2002, Azam and Malfatti 2007). Particles provide substrates for attachment as well as access to organic C that could fuel chemoheterotrophic N_2 fixation, while promoting the formation of low O_2 microenvironments that provide protection for the O_2 sensitive nitrogenase enzyme (Paerl 1985, Ploug et al. 1997, Ploug 2001, Ploug and Bergkvist 2015). Cyanobacterial aggregates (Klawonn et al. 2015) or self-produced NCD aggregates bound together by O_2 -impermeable exopolysaccharides may likewise form anoxic microniches for NCDs as exemplified by a *Pseudomonas stutzeri* strain (BAL361) isolated from the Baltic Sea (Fig. 2b; Bentzon-Tilia et al. 2015a, Paerl et al. 2018). A recent cellular model of *A. vinelandii* showed that the energetic costs to maintain low intracellular O_2 for a free-living chemoheterotrophic diazotroph through increased respiration, synthesis of thicker cell membranes, or polysaccharide production sur-

Table 1. Marine NCD environments

	Environment	Known or theorized NCD habitat/lifestyle ^a	Representative references
Water column	Open ocean: euphotic	Free-living; suspended and sinking particles; plankton holobionts	Proctor (1997), Zehr et al. (1998), Langlois et al. (2005, 2008, 2015), Church et al. (2005a, b), Fong et al. (2008), Moisander et al. (2008, 2014), Farnelid et al. (2010, 2011) Bombar et al. (2011), Kong et al. (2011), Zhang et al. (2011, 2016), Halm et al. (2012), Shiozaki et al. (2014, 2017) Sunagawa et al. (2015), Azimuddin et al. (2016), Benavides et al. (2016b), Gradoville et al. (2017a), Delmont et al. (2018), Chen et al. (2019), Wu et al. (2019), Yang et al. (2019), Cheung et al. (2020, 2021), Raes et al. (2020), Pierella Karlusich et al. (2021), Hallstrøm et al. (2022a)
	Open ocean: aphotic	Free-living; suspended and sinking particles	Hewson et al. (2007), Bonnet et al. (2013), Benavides et al. (2015, 2018c), Selden et al. (2019), Acinas et al. (2021)
	Oxygen minimum zones	Free-living; suspended and sinking particles	Fernandez et al. (2011), Hamersley et al. (2011), Löscher et al. (2014), Jayakumar et al. (2017), Chang et al. (2019), Reeder and Löscher (2022)
	Temperate coastal ecosystems	Free-living; suspended and sinking particles; plankton holobionts; resuspended sediments; terrestrial particles	Short et al. (2004), Moisander et al. (2007), Rees et al. (2009), Mulholland et al. (2012), Messer et al. (2015, 2021), Scavotto et al. (2015), Shiozaki et al. (2015), Bentzon-Tilia et al. (2015b), Gradoville et al. (2017a), Pedersen et al. (2018), Cabello et al. (2020), Turk-Kubo et al. (2021), Hallstrøm et al. (2022b)
	Inland seas	Free-living; suspended and sinking particles	Bostrom et al. (2007), Man-Aharonovich et al. (2007), Farnelid et al. (2013), Rahav et al. (2013, 2016), Benavides et al. (2016a), Kirkpatrick et al. (2018), Geisler et al. (2020), Ridame et al. (2022)
	Polar seas	Free-living; suspended and sinking particles; sediment resuspension; terrestrial input	Blais et al. (2012), Fernandez-Mendez et al. (2016), Shiozaki et al. (2018b, 2020)
Benthic	Coral reefs and sponges	Associated with released mucus, coral tissues, coral skeletons, associated macroalgae, and microbial mats, surrounding seawater	Mohamed et al. (2008), Olson et al. (2009), Lema et al. (2012, 2014), Ribes et al. (2015), Yang et al. (2016), Liang et al. (2020), Moynihan et al. (2022)
	Macroalgae	Living and decomposing tissue, surrounding seawater	Hamersley et al. (2015), Zhang et al. (2015), Raut et al. (2018), Raut and Capone (2021)
	Coastal and continental shelf sediments	Sediment-associated	Fulweiler et al. (2013), Brown and Jenkins (2014), Newell et al. (2016), Jabir et al. (2018)
	Oxygen-deficient zone sediments	Sediment-associated	Gier et al. (2016, 2017)
	Abyssal pelagic	Associated with methane seeps, hydrothermal vents, whale falls	Mehta et al. (2003), Mehta and Baross (2006), Pernthaler et al. (2008), Dang et al. (2009), Dekas et al. (2009, 2014, 2016, 2018), Miyazaki et al. (2009), Kapili et al. (2020)
	Microbial mats	Microbial mat associated, surrounding water	Zehr et al. (1995), Steppe et al. (1996), Olson et al. (1999)
	Seagrass rhizosphere	Sediments, roots	McGlathery et al. (1998), Mohr et al. (2021)

^aAllocation of NCDs between free-living and particle-association is not well-resolved.

passed the cost of N₂ fixation (Inomura et al. 2017), underscoring the potential importance of particle- or aggregate-associated lifestyles.

To better understand the potential for particles as N₂ fixation hot spots, Chakraborty et al. (2021) designed a mathematical model to investigate N₂ fixation by NCDs bound to large (ca. 250 μm) sinking marine particles (reviewed by Riemann et al. 2022). Cells were modeled as facultative diazotrophs that acquire C and N from particle-supplied polysaccharides and polypeptides and only supplement cellular N requirements with N₂ fixation once other N sources are exhausted, producing high C:N ratios favorable for heterotrophic N₂-fixers. Microbial respiration depletes O₂, however, the anoxic or microanoxic conditions created in the particle interior is temporary. Cells eventually become C-limited and O₂ diffusion into the particle again exceeds microbial respiration, which in turn inhibits N₂ fixation. NCD N₂ fixation in the

modeled particle interior thus depends on the surrounding water O₂ concentration, initial composition and ratio of C to N in the particle, particle size (minimum of 0.6 mm diameter), and sinking speed. Based on this simulation, the amount of time a particle may harbor an anoxic or low O₂ microzone and support N₂ fixation can be short-lived (<1 day), which may make this process challenging to validate with experimental data.

A few recent studies show the most direct evidence to date of N₂ fixation by NCDs bound to particles. In eutrophic waters of the Qishon estuary (Israel), which flow into the Mediterranean Sea, immunolabeling of the nitrogenase protein was used to demonstrate that NCDs colonized aggregates (Geisler et al. 2020). Furthermore, active N₂ fixation by NCDs was implicated based on the detection of N₂ fixation in particle-enrichment incubations where cyanobacterial photosynthesis was inhibited. This immunolabeling approach requires intact nitrogenase proteins but provides no

phylogenetic information making interpretation challenging. Another recent study surveyed well-lit oxygenated pelagic waters of the North Pacific and measured the incorporation of $^{15}\text{N}_2$ into putative NCDs (defined as cells with ^{15}N enrichment, but not ^{13}C enrichment after 24h $^{15}\text{N}_2/^{13}\text{C}$ -bicarbonate incubations) residing on small particles ($<210\ \mu\text{m}$, smaller than those predicted to host anaerobic microzones by Chakraborty et al. 2021) using particle-targeted nanoSIMS (Harding et al. 2022). Single cell N_2 fixation rates ranged from low to quite high (0.02 to 8.61 fmol N cell $^{-1}$ d $^{-1}$) and in some cells could entirely fulfill cellular N requirements. Complementary *nifH* gene HTS indicated γ -proteobacterial NCDs, including Gamma A, were prevalent in the water column. However, particle-targeted nanoSIMS does not provide phylogenetic information needed to identify taxonomic affiliations of the active NCDs.

Despite this recent progress, most current knowledge of particle-associated NCDs is based on molecular surveys. Accounts of NCDs from large size fractions or marine particles, based on both PCR-based and PCR-free approaches (Turk-Kubo et al. 2014, Benavides et al. 2016b, Gradoville et al. 2020, Pierella Karlusich et al. 2021), collectively suggest diverse NCD communities attached to suspended particles, sinking particles (marine snow), and fecal pellets (Riemann et al. 2010), as well as forming self-produced aggregates (Bentzon-Tilia et al. 2015a). Notably, NCDs hypothesized to be associated with larger organisms either as symbionts or as prey items (Farnelid et al. 2009, Scavotto et al. 2015, Pierella Karlusich et al. 2021) could also be recovered from large size fractions. Techniques linking specific taxa to active N_2 fixation are needed to assess the biogeochemical importance of these diverse particle-associated NCDs in the euphotic zone.

NCDs are also found on particles sinking out of the euphotic zone (Farnelid et al. 2019) and in the deep ocean (Rahav et al. 2013, Acinas et al. 2021, Poff et al. 2021). Particles (20–500 μm) sampled from floating net traps deployed at a depth of 150 m in the North Pacific often contained NCD sequences from Cluster 1 (including Part-Farn-OTU1012 and Part-Farn-OTU1107, which are Gamma 3 and Gamma A, respectively; Fig. 2c) and Cluster 3 (Farnelid et al. 2019; Table S2, Supporting Information). Notably, some sequences from individual particles ($>20\ \mu\text{m}$) were not well-represented in the whole water column diazotrophic community in this study, suggesting that preference for particle attachment may be taxa-specific. Additionally, three novel NCD MAGs have recently been described from mesopelagic particles including a member of the Micavibrionaceae family (Poff et al. 2021), a putative sulfur-oxidizing lithotroph (α -proteobacteria MAG0509), and a Gamma 4 MAG (MAG0081; Acinas et al. 2021). It remains unclear whether these sinking particle-associated NCDs are active N_2 -fixers, but collectively these studies suggest that particles delivered to the deep ocean harbor both distinct NCD lineages and those known to also reside in the surface ocean.

There remain large knowledge gaps about the role of marine particles in N_2 fixation by NCDs. Most critically, although NCDs are clearly present on marine particles and there is some evidence of NCDs fixing N_2 , no studies have demonstrated N_2 fixation by a taxonomically defined, particle-bound NCD. Further work is needed to determine the magnitude of potential fixed N inputs from particle-bound NCDs given the high concentration of particles in marine systems (Riemann et al. 2022).

Plankton holobionts

NCDs are often found associated with planktonic organisms, although the exact nature of these interactions is not well-

described. NCDs affiliated with α -, β -, and γ -proteobacteria, Clusters 2 and 3 have been found in association with heterotrophic dinoflagellate–cyanobacteria consortia (Farnelid et al. 2010). Cheung et al. (2021) noted that one of the prevalent γ -proteobacterial sequences found with several dinoflagellate genera described in Farnelid et al. (2010) was Gamma 4 (Fig. 2f); the lack of host specificity suggests a facultative symbiotic interaction, or possibly grazing on Gamma 4. *Trichodesmium* colonies also contain diverse assemblages of Cluster 3 and α -, β -, and γ -proteobacteria, notably including ClusterIII-C (Fig. 2d; Gradoville et al. 2014, 2017b). More generally, δ -proteobacterial (1A) and γ -proteobacterial taxa (1G) have been found actively transcribing *nifH* in net tow samples ($>100\ \mu\text{m}$) (defined as ‘plankton associated diazotrophs’; Yang et al. 2019), including a putative *Marichromatium*-like sequence type closely related to Gamma 3 (KY774962.1, 98.4% nucleotide identity). NCDs are also associated with small photosynthetic picocoeukaryotes, but these observations are sparse (Bombar et al. 2013).

Pelagic copepods have long been suspected to form symbiotic associations with N_2 -fixing bacteria, originally based on the visible growth of purple sulfur bacteria from copepod enrichment cultures allowed to develop anoxia, after which N_2 fixation could be detected (Proctor 1997). Subsequent studies have described diverse NCD taxa presumed to be affiliated with copepods (Braun et al. 1999, Scavotto et al. 2015, Azimuddin et al. 2016). In addition, Scavotto et al. (2015) reported distinct NCD community compositions between full-gut and starved copepods that suggest δ -proteobacteria diazotrophs may be grazed from particles, while γ -proteobacteria may form more permanent symbioses with copepods.

Beyond the presence of NCD *nifH* genes, very little is known about these associations. Unique challenges exist in sampling plankton holobionts. For example, plankton nets collect (and concentrate) various types of aggregates in addition to plankton, thus do not allow for confident differentiation between plankton- and particle-associated lifestyles. Nonetheless, holobionts are understudied habitats for marine NCDs and more research is needed to better characterize these associations, and to elucidate the role a host or partner may have in regulating N_2 fixation in plankton-NCD symbioses.

Coastal ecosystems

The availability of fixed inorganic N in coastal waters has been argued to select against N_2 fixation, as diazotrophs are generally considered to be poor competitors for acquiring P and Fe compared to faster growing photoautotrophs such as diatoms (Ward et al. 2013, Landolfi et al. 2015). However, some cyanobacterial diazotrophs (most notably the UCYN-A symbiosis, e.g. Short and Zehr 2007, Mulholland et al. 2012, Turk-Kubo et al. 2021) and NCDs are now recognized as components of coastal microbial communities (Table 1). NCDs are often found in nutrient-replete coastal waters and can reach high relative abundances, particularly in regions with terrestrial input, e.g. from riverine or estuarine sources (Moisander et al. 2008, Bombar et al. 2011, Kong et al. 2011, Hashimoto et al. 2012, Shiozaki et al. 2015, Rahav et al. 2016, Geisler et al. 2020, Selden et al. 2021, Hallstrøm et al. 2022b). It is important to note, however, that very little is known about the physiology or activity of NCDs in coastal areas; their presence suggests that coastal NCDs may be insensitive to dissolved inorganic N availability and/or ‘facultative’ diazotrophs capable of utilizing alternate N sources (discussed in ‘Diversity and ecophysiological features inferred from MAGs’).

NCDs reported in euphotic coastal waters often reflect the typical diversity of marine sediments, which suggests that sediment resuspension may be an important mechanism for introducing NCDs into the water column in coastal ecosystems. Periodic resuspension events are prevalent in shallow waters and areas with high wind or wave activity (Zouch et al. 2018). In particular, Cluster 3 *nifH* sequences have been widely reported in coastal regions, occasionally transcribing *nifH* (Short et al. 2004, Bentzon-Tilia et al. 2015b, Shiozaki et al. 2020, Hallstrøm et al. 2022b), suggesting NCD N_2 fixation in the oxygenated water column. In Western North Atlantic coastal waters, NCD *nifH* genes and transcripts dominate over those from cyanobacteria where salinity decreases due to freshwater input and turbidity increases through the resuspension of terrestrial and coastal sediments (Mulholland et al. 2019, Geisler et al. 2020, Selden et al. 2021, Hallstrøm et al. 2022b).

In the eutrophic Roskilde Fjord (Denmark), N_2 fixation rates were higher in sediment-amended seawater than in the surrounding seawater ($217 \text{ nmol N l}^{-1} \text{ d}^{-1}$ vs. $1.7 \text{ nmol N l}^{-1} \text{ d}^{-1}$, respectively), supporting the idea that NCDs resuspended from sediment fix N_2 in the water column (Pedersen et al. 2018). This same study demonstrated that NCDs were secondary colonizers of artificial particles in natural seawater, indicating their capability for motility and particle attachment. This successional attachment to particles may be indicative of a preference for sufficiently large particles that support low O_2 zones due to high activity of microbial respiration (see 'Marine particles').

Surface pelagic ocean waters

Open ocean gyres are typically N-limited and are known habitats for diazotrophs. Although the biogeography and environmental determinants are better understood for cyanobacterial diazotroph taxa, NCDs have been commonly recovered in *nifH* gene surveys dating back to early clone library-based studies in the North and South Pacific Oceans and the Mediterranean Sea (Zehr et al. 1998, 2003, Church et al. 2005b, Man-Aharonovich et al. 2007, Fong et al. 2008). Farnelid et al. (2011) were the first to apply *nifH* gene HTS in the global surface ocean, establishing that NCDs were broadly distributed and co-occur with cyanobacterial diazotrophs in tropical and subtropical waters. Farnelid et al. (2011) also revealed global taxa-specific NCD distribution patterns and confirmed that many NCDs were actively transcribing *nifH*, including those in small ($<10 \mu\text{m}$) size fractions. Since this foundational work, studies leveraging *nifH* gene HTS have significantly expanded the known habitats for NCDs in the ocean, most notably in pelagic ecosystems (Table 1).

Several studies have demonstrated that distinct NCD communities may be found in different regions of the global ocean (Farnelid et al. 2011, Shiozaki et al. 2017, Raes et al. 2020), but there are also several NCDs that appear to have more cosmopolitan distributions. This includes arguably the best-studied NCD Gamma A (Langlois et al. 2005), also referred to as AO15 (Zehr et al. 1998), UMB (Bird et al. 2005), and γ -24774A11 (Moisander et al. 2008).

Gamma A is broadly distributed throughout the tropical and subtropical North Atlantic and Pacific (reviewed by Langlois et al. 2015) and is theorized to be an oligotrophic specialist. Evidence of Gamma A as a potential contributor to N_2 fixation in oligotrophic gyres has been demonstrated in multiple surveys of *nifH* transcripts showing that Gamma A can account for a majority of the total transcript pool (Bird et al. 2005, Langlois et al. 2005, Church et al. 2005b, Bombar et al. 2011, Farnelid et al. 2011, Moisander et al. 2014, Messer et al. 2015, Shiozaki et al. 2017, Gradoville et al. 2020, Selden et al. 2021). Additionally, in the West-

ern South Pacific, Gamma A *nifH* genes and transcripts positively correlate with cyanobacterial diazotroph abundances, temperature and DOC, and negatively with depth, chlorophyll *a*, and nutrients, suggesting similar physiological constraints as *Trichodesmium* and *Crocospaera* (Moisander et al. 2014). The presence and *nifH* expression of Gamma A in well-lit surface waters suggests it benefits directly or indirectly from light. It has been speculated that Gamma A may have a photoheterotrophic lifestyle and utilize rhodopsin or bacteriochlorophyll-based energy generation or may rely on photosynthate from a photoautotroph (Langlois et al. 2015, Benavides et al. 2016b, Cornejo-Castillo and Zehr 2021). It is also found in association with larger size fractions ($>3 \mu\text{m}$), which has raised speculation of a symbiotic- or particle-bound lifestyle (Benavides et al. 2016b, Cornejo-Castillo and Zehr 2021; Fig. 2a).

Many other NCD taxa are routinely recovered in *nifH* gene surveys and, more recently, in metagenomic surveys (see 'Global ocean surveys using 'omics provide insights into NCD abundance and diversity') of the pelagic oceans. Several recent studies have demonstrated latitudinal shifts in NCD communities across the Pacific (Shiozaki et al. 2017, Gradoville et al. 2020, Raes et al. 2020). For example, cold, nutrient-rich sub-Antarctic waters support NCDs affiliated with α -, δ -, and ϵ -proteobacteria, along with Actinobacteria, while diverse γ -proteobacteria were seen in warmer, low latitude Pacific waters (Raes et al. 2020). Currently, the biological (e.g. nutrient limitation, grazing) or physical (e.g. advection on currents) mechanisms behind these observed latitudinal shifts in NCD community composition are not well-understood.

Aphotic waters and O_2 -deficient zones

Active N_2 fixation attributed to NCDs has been reported in aphotic waters in the Pacific (Fernandez et al. 2011, Hamersley et al. 2011, Bonnet et al. 2013, Löscher et al. 2014, Benavides et al. 2015, 2018c, Jayakumar et al. 2017, Selden et al. 2019), Mediterranean Sea (Rahav et al. 2013, Benavides et al. 2016a), Baltic Sea (Farnelid et al. 2013), and Black Sea (Kirkpatrick et al. 2018). Aphotic N_2 fixation rates are typically low ($< 1 \text{ nmol N l}^{-1} \text{ d}^{-1}$) making them difficult to measure with accuracy and precision, especially given methodological challenges associated with the low biomass often recovered and the different $^{15}\text{N}_2$ techniques (see 'Moving from genes to rates: are NCDs fixing N_2 in the pelagic oceans?'). However, considering the vastness of the marine aphotic zone, low but persistent rates of N_2 fixation could significantly affect the global marine N budget (Bonnet et al. 2013, Benavides et al. 2016a). Several early studies reported aphotic N_2 fixation rates in OMZs and suboxic waters (e.g. Fernandez et al. 2011, Hamersley et al. 2011, Farnelid et al. 2013), where NCDs are thought to be favored by O_2 -deplete and Fe-replete conditions where denitrification creates low N:P ratios (Deutsch et al. 2007, Löscher et al. 2014). However, in more recent studies, N_2 fixation was largely undetectable in the OMZs and suboxic waters in the Pacific and Indian Oceans (Chang et al. 2019, Selden et al. 2019, Löscher et al. 2020), implying spatial or temporal variability of aphotic N_2 fixation and/or changes to the calculation and reporting of detection limits (White et al. 2020).

NCDs present in aphotic waters are mostly Cluster 1 α -proteobacteria and γ -proteobacteria and Cluster 3 anaerobes affiliated with δ -proteobacteria and *Clostridia* (Fernandez et al. 2011, Hamersley et al. 2011, Bonnet et al. 2013, Farnelid et al. 2013, Löscher et al. 2014, Benavides et al. 2015). Clusters 2 and 4 *nifH* sequences have also been associated with OMZs (Löscher et al. 2014, Jayakumar et al. 2017). Although NCDs are well-established as the dominant diazotrophs in aphotic waters and low rates of N_2 fixation have been measured, little is known about which NCDs

are active and the factors controlling their activity (see discussion in 'Environmental drivers of NCD biogeography, activity, and presumed N₂ fixation').

Polar seas

Polar waters have long been assumed to be devoid of N₂ fixation. However, recent efforts to measure N₂ fixation in polar waters suggest it is an active process in both the Arctic and Antarctic Oceans, particularly in coastal and continental shelf regions (Blais et al. 2012, Sipler et al. 2017, Harding et al. 2018, Shiozaki et al. 2018b, 2020). While N₂ fixation by the UCYN-A/haptophyte symbiosis has been confirmed in polar waters (Harding et al. 2018), NCDs are also prevalent (Farnelid et al. 2011, Blais et al. 2012, Fernandez-Mendez et al. 2016, Shiozaki et al. 2018b) and are suspected to account for some portion of the N₂ fixation (Blais et al. 2012, Harding et al. 2018). In the Arctic, *nifH* gene surveys suggest that the NCD community in polar waters is primarily composed of δ -proteobacterial sequence types affiliating with Clusters 1A and 3 that are not closely related to sequence types recovered from other regions of the ocean (Blais et al. 2012, Fernandez-Mendez et al. 2016, Shiozaki et al. 2018b). Cluster 3 sequence types that appear endemic to the coastal Antarctic Ocean have also been described by Shiozaki et al. (2020). Collectively, the prevalence of putative anaerobes suggest that sediment resuspension plays an important role in shaping the pelagic diazotrophic community in these regions. However, several γ -proteobacterial NCDs have also been reported in the Antarctic with high *nifH* sequence similarity to oligotrophic taxa [including Gamma A, identified as SV009 in Shiozaki et al. (2020)].

Interestingly, a recent study reported the presence of 'ultra-small' (<0.22 μ m) NCDs (Pierella Karlusich et al. 2021) comprising up to 10% of the ultrasmall bacterioplankton in Arctic Ocean waters based on metagenome-derived abundances (described in 'Global ocean surveys using 'omics provide insights into NCD abundance and diversity'). An ϵ -proteobacterium, *Arcobacter nitrofigilis*, dominated bacterioplankton populations at the surface and deep chlorophyll maximum, while a γ -proteobacterium dominated in the mesopelagic. Both ultrasmall diazotrophs were also present in larger size fractions, suggesting they may have particle- or symbiont-associated lifestyles. At present, the ecological and biogeochemical importance of ultrasmall NCDs is unknown in the global oceans.

Environmental drivers of NCD biogeography, activity, and presumed N₂ fixation

Nutrient perturbation experiments

We are still in the early stages of understanding the environmental controls on marine NCD biogeography and activity. Since few marine NCDs are available in culture, environmental controls must be inferred from biogeographical surveys coupled to environmental data (discussed in 'Meta-analysis of *nifH* gene abundance data') and from *in situ* experiments involving nutrient and/or environmental manipulations. Such experiments have yielded important insights but can be challenging to interpret given the regular co-occurrence of cyanobacterial diazotrophs and NCDs, the complexity of microbial interactions among the broader community, and the lack of standardized approaches across experiments. For example, some studies employ *nifH* amplicon libraries, while others quantitatively target specific NCDs with qPCR/ddPCR. Additionally, some studies note enhanced N₂

fixation rates without observed changes in NCD abundances/*nifH* transcription, while other studies characterize the diazotroph community composition in the region, but not in the experiments themselves making interpretation challenging. Table 2 summarizes some environmental drivers of NCD abundance, *nifH* transcription or putative N₂ fixation based on representative studies using a variety of approaches.

Together these experiments indicate that NCD abundances and *nifH* transcription are sometimes limited by the availability of Fe, P (or both), and/or DOC in oligotrophic euphotic waters (Table 2). Gamma A abundances appear to be influenced (at least in part) by the availability of Fe in the Western South Pacific (Moisander et al. 2012). Regional differences are seen in the response of Gamma A to N availability; N additions resulted in decreased abundances of Gamma A in the Western South Pacific (Moisander et al. 2012), whereas in the North Atlantic, abundances increased in experiments with fixed N additions (Langlois et al. 2012). Aeolian dust input is an important source of Fe to surface oceans and has been shown to influence NCD abundances in the North Atlantic Ocean (Langlois et al. 2012) and the Mediterranean Sea (Ridame et al. 2022). The availability of labile DOC is also a control on N₂ fixation in regions where NCDs are thought to be the most prevalent diazotrophs, like in the Eastern Tropical South Pacific (Dekaezemacker et al. 2013, Turk-Kubo et al. 2014, Knapp et al. 2016).

DOC availability may also be a particularly important control on aphotic N₂ fixation (Table 2). For example, aphotic N₂ fixation was positively correlated with transparent exopolymeric particles (TEPs) in the oxygenated waters of the South Pacific Ocean and the Mediterranean Sea (Rahav et al. 2013, Benavides et al. 2015), suggesting that TEPs could provide C-rich and/or O₂-depleted microenvironments that favor NCD N₂ fixation. Moreover, additions of glucose and amino acids occasionally enhance aphotic N₂ fixation (Bonnet et al. 2013, Rahav et al. 2013, Löscher et al. 2014, Benavides et al. 2015, Gradoville et al. 2017a).

Although these studies illustrate that perturbations in nutrient availability (e.g. nutrient, trace metal, and labile organic C) or environmental conditions (e.g. availability of light) can lead to changes in the abundance and *nifH* transcription of NCDs, data from these experiments are relatively sparse and spatially heterogeneous. Fortunately, valuable insights into the environmental drivers behind diazotroph biogeography can also be inferred from biogeographical surveys coupled to environmental data ('Meta-analysis of *nifH* gene abundance data').

Meta-analysis of *nifH* gene abundance data

In the absence of direct cell counts, quantifying *nifH* gene abundances (via qPCR or ddPCR) is arguably the best method for enumerating NCDs and determining the biogeographical patterns and environmental drivers of specific taxa. Unfortunately, oceanographic sample collection, DNA extraction, and *nifH* gene quantification are laborious compared to more high-throughput methods (e.g. automated flow cytometry) and the coverage of qPCR/ddPCR *nifH* gene abundance datasets are often sparse in space, time, and NCD targets. Thus, it can be difficult to discern the global distribution patterns and environmental predictors of diverse NCD taxa from a single dataset.

A global *nifH* qPCR database has been compiled for marine cyanobacterial diazotrophs (Luo et al. 2012, Tang et al. 2019a), revealing taxon-specific biogeographical patterns. Linking these qPCR abundances to environmental data has revealed both common and distinct environmental drivers among taxa. For ex-

Table 2. Changes in NCD abundances, activity, and putative N_2 fixation in response to nutrient perturbations provide insights into their environmental drivers. NPSG—North Pacific Sub-tropical Gyre; WSP—Western South Pacific; ETSP—Eastern Tropical South Pacific; SCS—South China Sea; MedSea—Mediterranean Sea; RT-qPCR—reverse transcription qPCR; N—nitrogen; Fe—iron; P—phosphate; DOC—dissolved organic C; DON—dissolved organic N; DOP—dissolved organic P; DCMU—photosynthesis inhibitor (3-(3,4-dichlorophenyl)-1,1-dimethylurea); and GX—xanthan gum.

NCD(e) ^a	Region (depth)	Type of analyses ^c	Environmental perturbation(s)	Synthesis of findings	Reference
Gamma A (AO15)	NPSG (0 – 25 m)	RT-qPCR; N_2 fixation	P	No significant stimulation of N_2 fixation or <i>nifH</i> transcripts	Zehr et al. (2007)
Cluster-3	NPSG (40 m)	qPCR	DOC	Cluster-3 increased in +DOC	Bombar et al. (2013)
Gamma A (γ24774A11)	WSP (5 m)	qPCR	N, P, Fe, DOC, Fe/P, N/P, N/Fe, N/P/Fe/DOC	Abundances increased in response to Fe and Fe/P in westernmost stations; decreased in +N	Moisaner et al. (2012)
Gamma A (γ24774A11)	WSP (3 m)	qPCR & RT-qPCR; N_2 fixation	DOC, DON, DOP, inhibition of photosynthesis using DCMU	No <i>nifH</i> expression across all treatment implying they were not actively fixing N_2 ; DCMU additions suppressed nearly all N_2 fixation implying that the most active N_2 -fixers were phototrophs	Benavides et al. (2018b)
P1, P4, P7	ETSP (95 m)	qPCR, N_2 fixation	Glucose, O ₂	N_2 fixation rates increased in +glucose, +O ₂ ; P7 abundances increased in +glucose, +O ₂	Löscher et al. (2014)
Unknown ^b	ETSP (15 m)	N_2 fixation rates	Fe, N or N/Fe, P, glucose	N_2 fixation rates increased in +Fe and +glucose and occasionally in +N/Fe	Dekaezemacker et al. (2013)
P2, P4, P6, P7	ETSP (0 m)	<i>nifH</i> transcript sequencing & RT-qPCR; N_2 fixation	Glucose	N_2 fixation stimulated by +glucose in eddy cores; <i>nifH</i> transcription from P2, P4, P6, P7 in eddy samples (not measured in experiments)	Löscher et al. (2016)
Unknown ^b	WNA (coastal)	N_2 fixation (ARA)	DOC, organic detritus, light and dark incubations	Water column N_2 fixation stimulated in +DOC, + organic detritus treatments	Pael and Prufert (1987)
Gamma A, gamma P, CIII-Church	ENA (1 – 3 m)	qPCR, N_2 fixation	N, P, Fe, dust	Gamma A abundances increased most in +NFe and +dust treatments; Gamma P and CIII were undetected or not quantifiable	Langlois et al. (2012)
Mainly 1G	MedSea (5 m)	<i>nifH</i> sequencing; N_2 fixation	Dust under contemporary and future temp. and pH scenarios	Increased N_2 fixation rates in response to dust additions in stations dominated by NCDs (cyanobacteria also present)	Ridame et al. (2022)
Diverse cluster I and III NCDs	MedSea (5 m)	<i>nifH</i> transcript sequencing; N_2 fixation	GX, N, P, DOC, NP, DOC/P, DOC/N, DOC/N/P; light and dark incubations	N_2 fixation stimulated by DOC in both light and dark incubations; increased relative abundances of NCD <i>nifH</i> transcripts in +DOC/N/P; increased N_2 fixation and NCD transcript relative abundances in +GX (cyanobacteria also present)	Rahav et al. (2016)
M6411A02, M6413A02, M6433A04	WSP (aphotic)	qPCR (environmental samples, not nutrient exp.); N_2 fixation	DOC, DON	N_2 fixation stimulated by amino acid (+DON) additions; assumed to be NCDs, but recently <i>Trichodesmium</i> has been shown to fix N_2 in mesopelagic waters (Benavides et al. 2022)	Benavides et al. (2015)
αETSP-2, cIII-ETSP	ETSP (aphotic)	<i>nifH</i> sequencing & qPCR, N_2 fixation	Amino acids, simple sugars	Aphotic N_2 fixation rates increased in +amino acids and +DOC treatments; identified NCDs did not change in abundance.	Bonnet et al. (2013)
Unknown (Gulf of Aqaba), 1 G (Med Sea)	MedSea, Gulf of Aqaba (aphotic)	<i>nifH</i> sequencing (environmental samples, not nutrient exp.); N_2 fixation	GX, amino acids	Aphotic N_2 fixation rates increased in +amino acids (Gulf of Aqaba) and +GX (Med Sea)	Rahav et al. (2013)

^aNCD *nifH* catalog name referenced when possible.

^bNCDs suspected to be dominant N_2 -fixers.

^c N_2 fixation measured on the whole community.

ample, abundances of UCYN-A, *Crocospaera*, *Trichodesmium*, and *Richelia* all correlate positively with temperature and negatively with depth, but UCYN-A is distributed across a larger temperature range, specifically being present in lower temperature waters (Tang et al. 2019a).

To our knowledge, previous compilations of global NCD *nifH* gene abundance data have only targeted Gamma A (Langlois et al. 2015, Shao and Luo 2022). To increase understanding of the distribution of NCDs beyond this phylotype, we compiled qPCR/ddPCR data from 59 published studies of the 55 targets included in our NCD *nifH* catalog (Fig. 1; Table S2, Supporting Information), yielding a total of 7385 water column observations (dataset doi: 10.5281/zenodo.6537451). This database shows that while NCD taxa have been quantified from many ocean regions, there has been a strong sampling bias, with the most samples collected from the North Pacific and few samples collected from the southern hemisphere (Figure S1, Supporting Information). Data is sparse for some NCD taxa (e.g. $n = 34$ observations of β -proteobacteria) but is particularly rich for γ -proteobacteria ($n = 4138$ observations). Here, we discuss the global distributions and environmental predictors of the three phylotypes for which the most data are available: Gamma A, Gamma 4, and a γ -proteobacterium of the order *Oceanospirillales* isolated by Ratten (2017) ('Gamma OcSpi'), as well as the most data-rich Cluster 3 phylotype ('ClusterIII-C'; Church et al. 2005a, Langlois et al. 2008). *nifH* gene transcripts of these groups have all been observed, except for Gamma OcSpi, for which no *nifH* RT-qPCR data (either presence or absence) have been reported (Figure S2, Supporting Information). There were eight additional NCD phylotypes with >100 *nifH* gene abundance observations; however, in most cases data for these phylotypes are only available from a single ocean region (Figure S3, Supporting Information).

The four most data-rich NCD phylotypes appear to have different global and depth distributions (Fig. 3). Abundances of the three γ -proteobacterial phylotypes are all highest at the surface and decrease with depth. Gamma A has the best sampling coverage of any NCD phylotype ($n = 2339$ samples), with *nifH* genes detected in 63% of samples. The highest Gamma A abundances ($>10^6$ *nifH* gene copies l^{-1}) have been observed in the North Pacific Ocean (Cheung et al. 2020), Indian Ocean (Wu et al. 2019), SCS (Liu et al. 2020), and New Caledonia Lagoon (Benavides et al. 2018b); Gamma A has not been detected in any samples collected from the Eastern South Pacific (Halm et al. 2012, Turk-Kubo et al. 2014, Shiozaki et al. 2018a; Fig. 3A). Gamma 4 has only been quantified in the Pacific Ocean, where its *nifH* gene sequence has been detected in 81% of samples ($n = 943$). The highest abundances ($>10^6$ *nifH* gene copies l^{-1}) have been observed in the North Pacific and Eastern Tropical South Pacific. Gamma 4 values below detection limits have been observed in the North Pacific, Central South Pacific, Eastern Tropical South Pacific, and SCS (Halm et al. 2012, Löscher et al. 2014, Chen et al. 2019, Cheung et al. 2021). Gamma OcSpi has only been explored in the upper 300 m of the North Atlantic, where *nifH* genes were detected in 98% of samples ($n = 440$; Ratten 2017). Maximum abundances of Gamma OcSpi were generally lower ($<10^5$ *nifH* gene copies l^{-1}) compared to Gamma A and Gamma 4 and the only observations of Gamma OcSpi below detection limits occurred in the western North Atlantic. Finally, ClusterIII-C *nifH* genes have been detected in 51% of samples ($n = 147$) and though a smaller number of samples have been collected than for the γ -proteobacterial phylotypes, coverage has been distributed over many ocean basins (Fig. 3). The highest ClusterIII-C abundances ($>10^5$ *nifH* gene copies l^{-1}) have been observed in the North Pacific Subtropical Gyre (Church et al. 2005a),

while relatively high abundances ($>10^4$ *nifH* gene copies l^{-1}) have been detected in the North Atlantic and the Bay of Bengal (Langlois et al. 2008, Löscher et al. 2020). Measurements of ClusterIII-C below detection limits have been reported in the Atlantic (Langlois et al. 2008). Notably, abundances of ClusterIII-C do not decrease with depth (Fig. 3), unlike the γ -proteobacterial phylotypes, and instead have a positive depth trend.

Comparing the distribution of these four NCD phylotypes is complicated by differences in sampling efforts and coverage (Fig. 3). For instance, samples for Gamma 4 and Gamma OcSpi have only been collected in the Pacific and Atlantic Oceans, respectively, and the three γ -proteobacterial phylotypes are heavily biased toward surface samples. Sampling strategies designed to target NCDs in undersampled ocean regions may be particularly useful for helping to constrain NCD biogeography. Reporting non-detects is also useful for efforts to model diazotroph distributions (Meiler et al. 2022) and we encourage researchers to do so in future studies.

To investigate the potential environmental determinants of NCD phylotypes, we colocalized qPCR abundance data with available World Ocean Atlas climatology, satellite observations, Argo float data, and PICES model products using the Simons Collaborative Marine Atlas Project (Simons C-MAP, <https://simonscmap.com>, Ashkezari et al. 2021). Details on variables and space/time/depth tolerances used for colocalization are provided in Table S4 (Supporting Information). All sampling depths were included in this analysis, and while many environmental variables covaried with depth, their relationships with *nifH* gene abundances were generally consistent when restricting analyses to surface samples (Figure S4, Supporting Information).

Environmental drivers of NCDs appear to differ among the four phylotypes investigated and diverge from those described for cyanobacterial diazotrophs (Fig. 4). Abundances of the three γ -proteobacteria correlated positively with temperature and negatively with NO_3^- and phosphate (PO_4^{3-}) concentrations; these same trends have been observed for cyanobacterial diazotrophs (Tang et al. 2019a). In contrast, abundances of ClusterIII-C correlated negatively with temperature and positively with NO_3^- and PO_4^{3-} concentrations (Fig. 4), reflecting the positive depth trend for this phylotype (Figure S4, Supporting Information). Gamma A and Gamma 4 abundances correlated negatively with the $NO_3^-:PO_4^{3-}$ (N:P) ratio, a trend also observed for most cyanobacterial diazotrophs (Tang et al. 2019a), while Gamma OcSpi and ClusterIII-C had weakly positive but nonsignificant correlations to N:P. The highest abundances of the γ -proteobacterial phylotypes were associated with moderate O_2 concentrations (~ 4.5 – 6.5 ml l^{-1}), while their abundances were lower at low O_2 concentrations (deep samples) and at high O_2 concentrations (supersaturated and/or cold surface ocean waters). Cluster III-C abundances appear negatively related to O_2 (although this trend was not statistically significant), with some of the highest abundances observed in deep, low- O_2 waters (<1 ml l^{-1}). This is consistent with a potentially anaerobic or facultative anaerobic lifestyle, i.e. a characteristic of many Cluster 3 diazotrophs. Relationships between NCD abundance and additional environmental variables are presented in Figure S4 (Supporting Information).

Overall, our analysis suggests greater variability in the biogeography and environmental predictors among NCD taxa than among cyanobacterial diazotroph taxa. This likely reflects the taxonomic and metabolic diversity of NCDs, which are distributed over broad bacterial and archaeal lineages (Fig. 1) and have diverse energy generation and nutrient transport mechanisms (see 'Diversity and ecophysiological features inferred from MAGs'). While

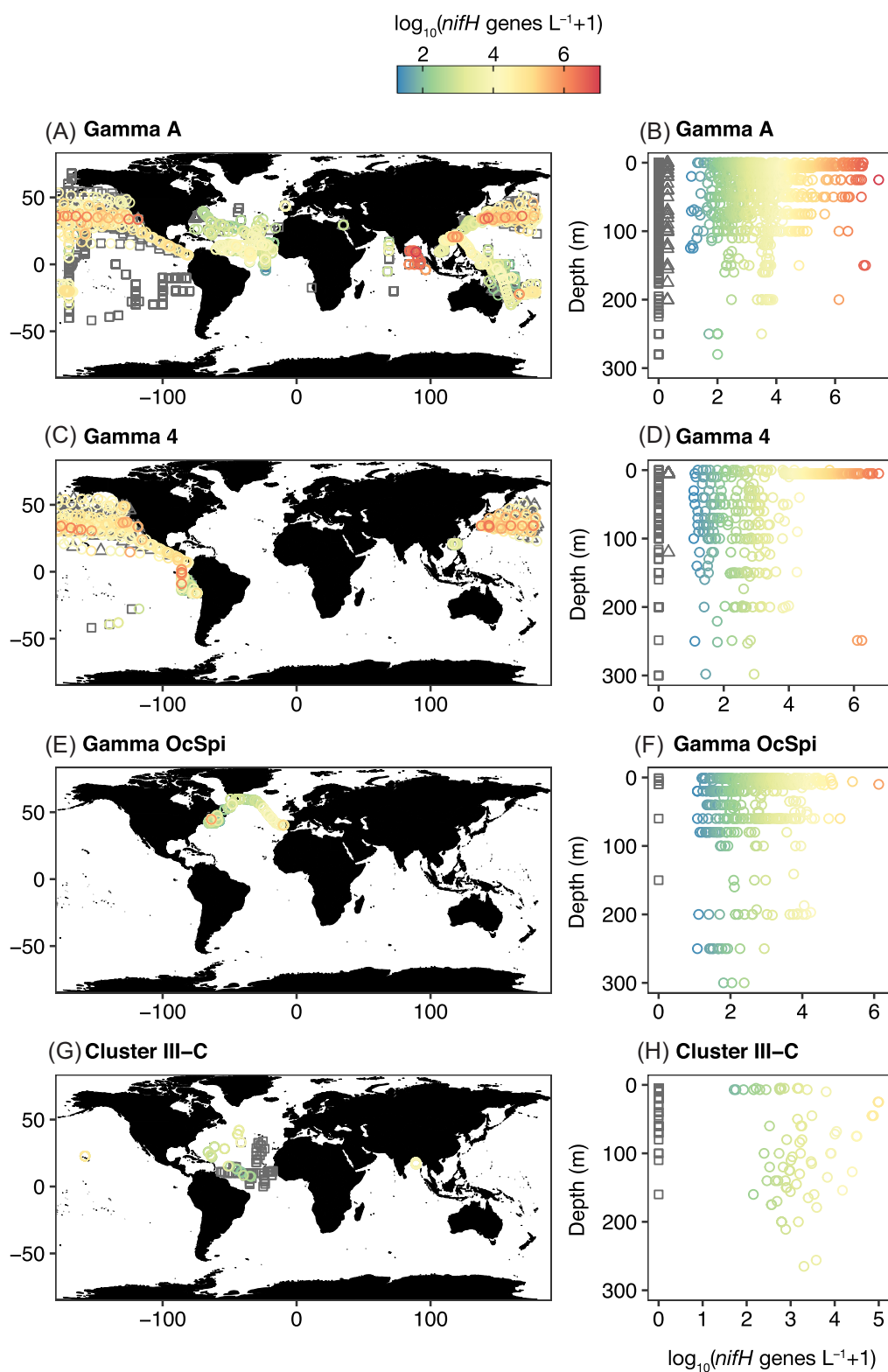


Figure 3. NCD taxa have distinct global and depth distribution patterns. Maps (A), (C), (E), and (G) show the global *nifH* gene abundances of four NCD taxa at all sampling depths (0–4000 m) while depth plots (B), (D), (F), and (H) show the upper 300 m only. Observations of no detect are represented by gray squares; observations of detect but not quantifiable (DNQ) were given a nominal value of 1 *nifH* gene L^{-1} in the database and are represented by gray triangles. Dataset doi: 10.5281/zenodo.6537451.

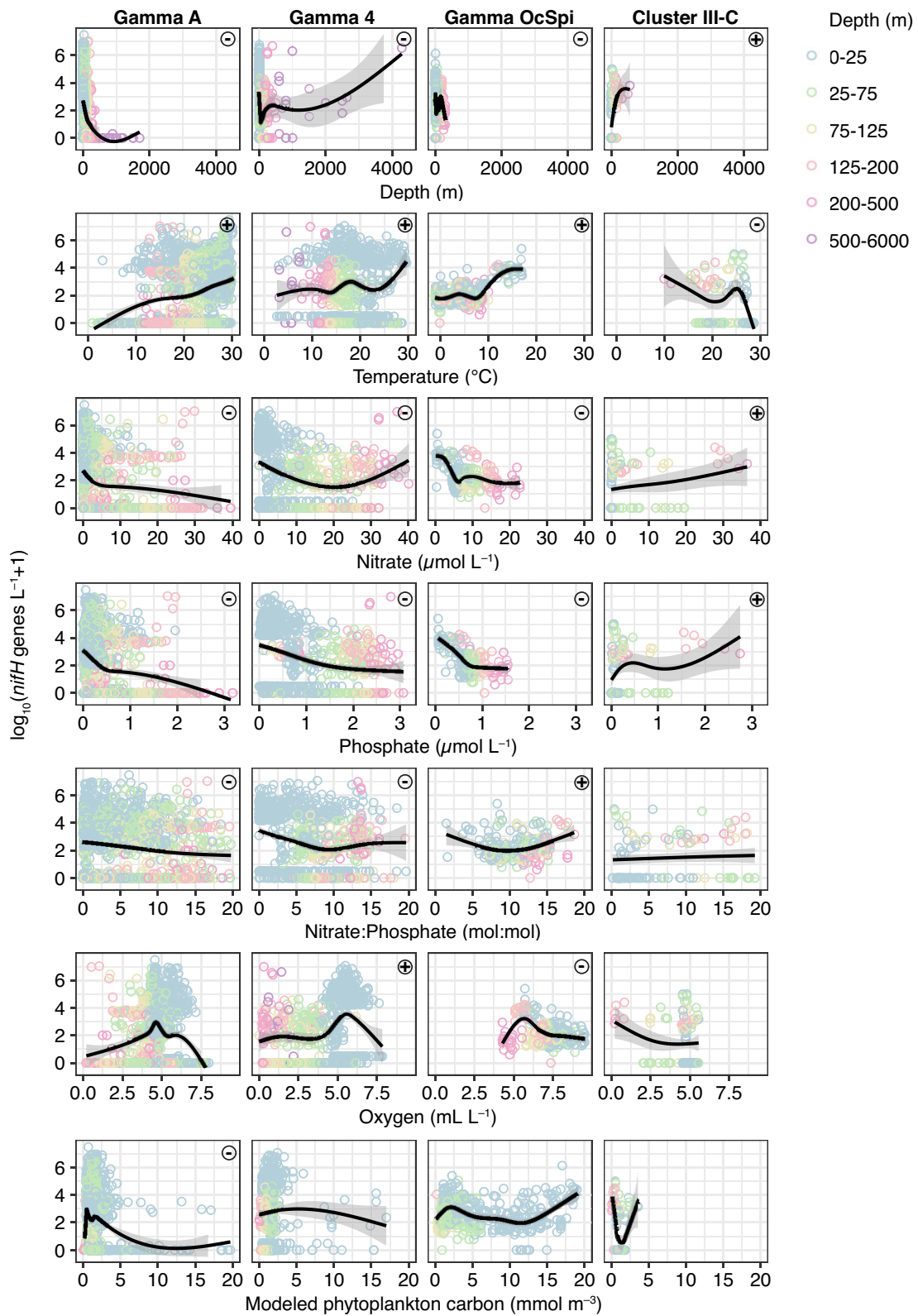


Figure 4. Environmental predictors vary among NCD taxa. NCD *nifH* gene abundances are plotted against environmental metadata from World Ocean Atlas monthly climatology (temperature, nitrate, phosphate, N:P, and O_2 concentration) and Pisces model output of phytoplankton biomass in units of carbon (modeled phytoplankton C). Tolerances for colocalization are presented in Table S4 (Supporting Information). Each point represents an individual *nifH* gene abundance sample, with sample depth shown in color. Note that a small fraction of data with outlying x-axis values were excluded from plots. Black lines and gray shading represent the smoothed conditional mean and 95% confidence intervals. Symbols (+/-) reflect positive and negative correlations (Spearman rank with Bonferroni correction for 28 comparisons).

genome comparisons among the four NCD taxa examined here are not yet possible, we presume that they have divergent ecophysiological features that may explain their different environmental drivers and distributions.

NCD biogeography and ecophysiology from metagenomes and metatranscriptomes

Global ocean surveys using 'omics provide insights into NCD abundance and diversity

Investigations of the global diversity and distribution of NCDs from an 'omics perspective are now possible due to recent improvements in HTS techniques, decreases in sequencing costs, and global ocean surveys focused on primer independent sequencing including Tara Oceans (Karsenti et al. 2011) and the Malaspina (Duarte 2015) expeditions. The Tara Oceans expedition has provided the most samples and deepest metagenomic sequencing effort to date for the open ocean (Sunagawa et al. 2015, Carradec et al. 2018, Salazar et al. 2019), with sampling coverage including all major ocean regions from the euphotic layer to the mesopelagic sea. The first version of the Ocean Microbiome Reference Gene Catalog (OM-RGC; Sunagawa et al. 2015) focused on the free-living size fraction (<3 μm) and provided a total of 28 *nifH* gene sequence variants, only two of which belonged to cyanobacteria (Cornejo-Castillo 2017). In recent years, increased metagenomic data available from undersampled environments, including polar regions and the deep ocean, has revealed new NCD *nifH* sequence variants, altogether showing that NCDs are globally distributed in surface and mesopelagic layers (Cornejo-Castillo 2017, Salazar et al. 2019, Acinas et al. 2021, Pierella Karlusich et al. 2021, Delmont et al. 2022). Some NCD *nifH* gene sequences are different from sequences identified with primer-based approaches, suggesting they may represent novel diazotrophic diversity not found in previous studies (Fig. 1; Cornejo-Castillo 2017, Delmont et al. 2018, 2022). In addition, the reconstruction of MAGs has provided vital insight into the genomic content of some marine NCDs, which is crucial to gain a better understanding of their physiology and ecology (see 'Diversity and ecophysiological features inferred from MAGs').

One advantage of metagenomic approaches is that normalization of nitrogenase genes to other genetic markers can provide an estimate of the relative abundance of NCDs to the total bacterioplankton community. Using this method, NCDs have been estimated to be more abundant than some previous reports ($\sim 10^6$ cells l^{-1} ; Delmont et al. 2018); NCDs were also among the top contributors to the *nifH* transcript pool in Tara Ocean samples (Salazar et al. 2019). The relative contributions of NCDs to total bacterioplankton were higher in large (5–20 μm) size fractions than in small (0.2–3 μm) size fractions (Pierella Karlusich et al. 2021) and relative abundances were significantly higher in the mesopelagic than in surface waters (Cornejo-Castillo 2017, Salazar et al. 2019, Pierella Karlusich et al. 2021). However, these estimations should be interpreted with caution since they have yet to be corroborated using other quantitative approaches.

Diversity and ecophysiological features inferred from MAGs

Acquiring NCD genomes from cultivated isolates and the assembly and binning of MAGs from the metagenomic datasets described above enables the prediction of metabolic pathways, and hence ecophysiological features, of marine NCDs from the surface ocean, estuaries, OMZs, and the bathypelagic (Bentzon-Tilia et al. 2015b, Martinez-Perez et al. 2018, Acinas et al. 2021, Cheung et

al. 2021, Poff et al. 2021, Delmont et al. 2022). Recently, 40 putative NCD MAGs with a high percent of completion were reconstructed from the Tara Oceans expedition (termed Heterotrophic Bacterial Diazotrophs, see Box 1) providing the largest dataset of the genomic potential of marine NCDs to date (Delmont et al. 2022). These MAGs represent diverse NCDs affiliated to Proteobacteria, Planctomycetes, and Verrucomicrobia (*nifH* Clusters 1 and 3). NCD MAGs contributed nearly half of the total *nifH* gene sequences detected in the Tara Oceans metagenomic dataset, although their *nifH* gene transcripts were greatly outnumbered by those of cyanobacterial diazotrophs (Figure S5A, Supporting Information). Among the NCD MAGs, γ -proteobacteria contributed the majority of the *nifH* sequences in both the metagenomic (55%) and metatranscriptomic (78%) datasets, followed by δ -proteobacteria ($\sim 15\%$ of both *nifH* genes and transcripts). In contrast, Planctomycetes and α -proteobacteria contributed 2%–3% of the *nifH* transcripts, while their relative abundances at DNA level were similar to δ -proteobacteria. The remaining NCD MAGs were affiliated with β -proteobacteria, ε -proteobacteria, and Verrucomicrobiota, which contributed <3% to the total *nifH* genes and transcripts (Figure S5B, Supporting Information). These findings suggest that γ -proteobacteria and δ -proteobacteria may be the dominant NCDs in the pelagic oceans.

We examined the ecophysiological features of NCD MAGs, focusing specifically on pathways involved in energy generation other than aerobic respiration and in the transport of fixed N, which can inhibit cyanobacterial N_2 fixation (Knapp 2012). In addition, we inferred potential regulatory mechanisms of N_2 fixation based on our understanding of similar pathways from terrestrial model NCDs, described in detail below. All MAGs with >90% completeness (as estimated by Anvi'o; Eren et al. 2015) containing all the nitrogenase structural genes (*nifHDK*) were included in this analysis (30 MAGs) and potential pathways in these MAGs were reconstructed using KEGGMAPPER (Kanehisa et al. 2012).

Although there have been reports of photoheterotrophic and chemolithoautotrophic NCDs in estuarine and bathypelagic waters (Bentzon-Tilia et al. 2015a, Acinas et al. 2021), all Tara Ocean NCD MAGs were originally identified as chemoheterotrophs (Delmont et al. 2022). Nevertheless, several energy generating pathways other than aerobic respiration were detected in the NCD MAGs, including e.g. anoxygenic photosystem II (APII), dissimilatory sulfate reduction (DSR), the SOX pathway (i.e. sulfur oxidation), dissimilatory NO_3^- reduction to NH_4^+ (DNRA), and denitrification (Fig. 5A; Hu et al. 2002, Kraft et al. 2011, Santos et al. 2015, Grabarczyk and Berks 2017). APII was only detected in an α -proteobacterial MAG and DSR was constrained to δ -proteobacteria. In contrast, the SOX pathway and DNRA were detected across MAGs of various groups. Denitrification was found in an α -proteobacterial MAG and two γ -proteobacterial MAGs. Approximately, half of the NCD MAGs contained at least one of these energy generating pathways, and one-third of them contained multiple pathways.

The NCD MAGs contained the genes for ABC transporters of various chemical forms of N, P, and Fe (Fig. 5B; Figure S6, Supporting Information). The genes encoding NH_4^+ transporters were detected in all NCD MAGs but only in some cyanobacterial diazotrophic MAGs. Additionally, the γ -proteobacterial and α -proteobacterial NCDs were genetically capable of transporting NO_3^- , NO_2^- , cyanate, urea, and amino acids, while the other NCDs and cyanobacterial diazotrophs generally lacked some or all genes in these pathways. The genetic capacity for uptake/transport of more forms of fixed N in NCDs than in cyanobacterial diazotrophs suggests that some NCDs may be facultative diazotrophs and

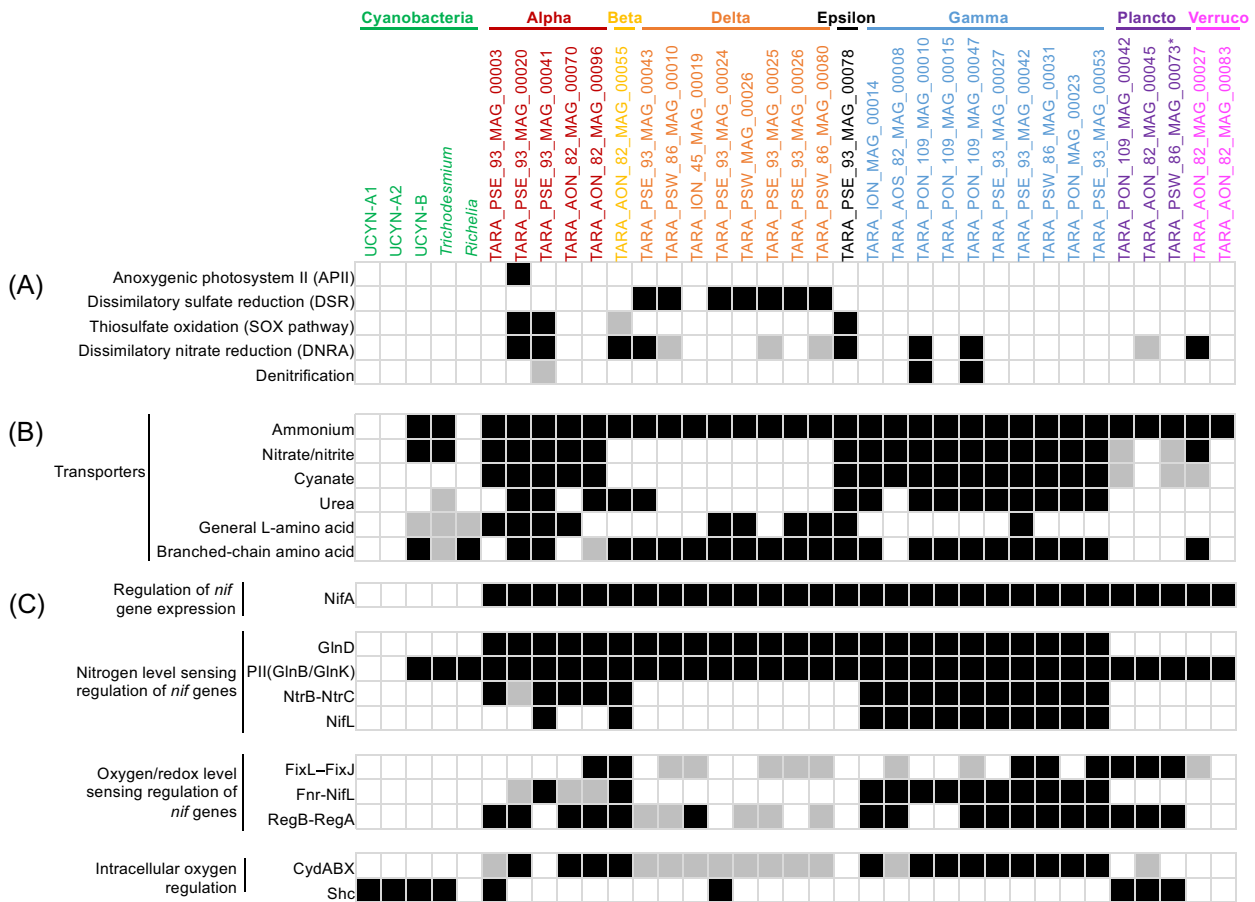


Figure 5. Ecophysiological features of NCDs include pathways involved in energy generation and transport of fixed N not present in cyanobacterial diazotrophs. Color represents presence/absence of metabolic pathways (black: all genes; gray: one missing gene; and white: no genes) related to energy acquisition (A), N uptake (B), and regulation of nitrogenase synthesis and activity (C) of NCDs and cyanobacterial diazotrophs as inferred from reconstructed Tara Oceans MAGs (Delmont et al. 2022). Alpha = α -proteobacteria; Beta = β -proteobacteria; Delta = δ -proteobacteria; Epsilon = ϵ -proteobacteria; Gamma = γ -proteobacteria; Plancto = Planctomycetes; and Verruco = Verrucomicrobiota. See Fig. 6 for more details about these regulatory pathways and the proteins involved.

switch to alternative fixed N forms when available under certain conditions. Most NCD MAGs also contained diverse P and Fe uptake pathways (Figure S6, Supporting Information).

N_2 fixation is a highly regulated process, due both to the energetic demands (high ATP and reductant requirements) and O_2 sensitivity of nitrogenase. As such, there are multiple complex pathways for the sensing of alternative N sources and intracellular O_2 conditions. Many potential N/ O_2 sensing regulatory mechanisms of N_2 fixation were found in the NCD MAGs (Figs 5C and 6). All NCD MAGs contained the *nifA* gene, which is the major regulator that interacts with the upstream environmental sensing pathways and controls the transcription of the *nif* operon in terrestrial NCDs (Dixon and Kahn 2004). In terms of known N-sensing regulatory mechanisms, only the proteobacterial MAGs contained the genes encoding the N sensing protein (GlnD) and PII proteins (GlnB or GlnK). Additionally, within that group, nearly all the α -proteobacterial and γ -proteobacterial MAGs contained the genes for the two-component system (NtrB-NtrC) that regulates the expression of *nifA* gene. Briefly, when alternative N forms are in excess, activation of *nifA* transcription is inhibited by the removal of uridylyl groups from PII by GlnD, which then interacts with NtrB and inhibits NtrC (Bueno Batista and Dixon 2019). When N is limiting, the PII proteins are uridylylated by GlnD, which cannot interact with NtrB-NtrC; uridylylated PII proteins can also directly

activate the NifA protein, and thus allow the expression of the *nif* operon (Dixon and Kahn 2004). Another regulatory protein (NifL), known to interact with non-uridylylated PII and inhibit NifA under excess N conditions in the terrestrial γ -proteobacterial diazotroph *A. vinelandii* (Little et al. 2000) was also detected, but mostly in γ -proteobacterial MAGs (Fig 5C).

Some NCD MAGs also contained genes for O_2 (FixL-FixJ) and redox (RegB-RegA) sensing two-component regulatory systems (Figs 5C and 6). When the intracellular conditions favor the activity of the O_2 -sensitive nitrogenases, these systems can activate the transcription of the *nifA* gene and allow the synthesis of nitrogenase (Dixon and Kahn 2004). The Fnr protein, which was mostly detected in γ -proteobacterial MAGs, can also sense O_2 levels and is important in maintaining the non-inhibitory form of NifL and the functioning of NifA in the terrestrial γ -proteobacterial diazotroph *Klebsiella pneumoniae* (Grabbe et al. 2001). Moreover, the NifA in some NifL-lacking terrestrial NCDs can also directly sense O_2 level (Dixon and Kahn 2004), which could theoretically be the case for some of the marine NCDs that appear to lack known O_2 sensing systems.

In addition to these regulatory processes, genes coding for cytochrome bd (CydABX) were found in most γ -proteobacterial and α -proteobacterial MAGs (Figs 5C and 6). The cytochrome bd consumes large amounts of O_2 during ATP generation, which was

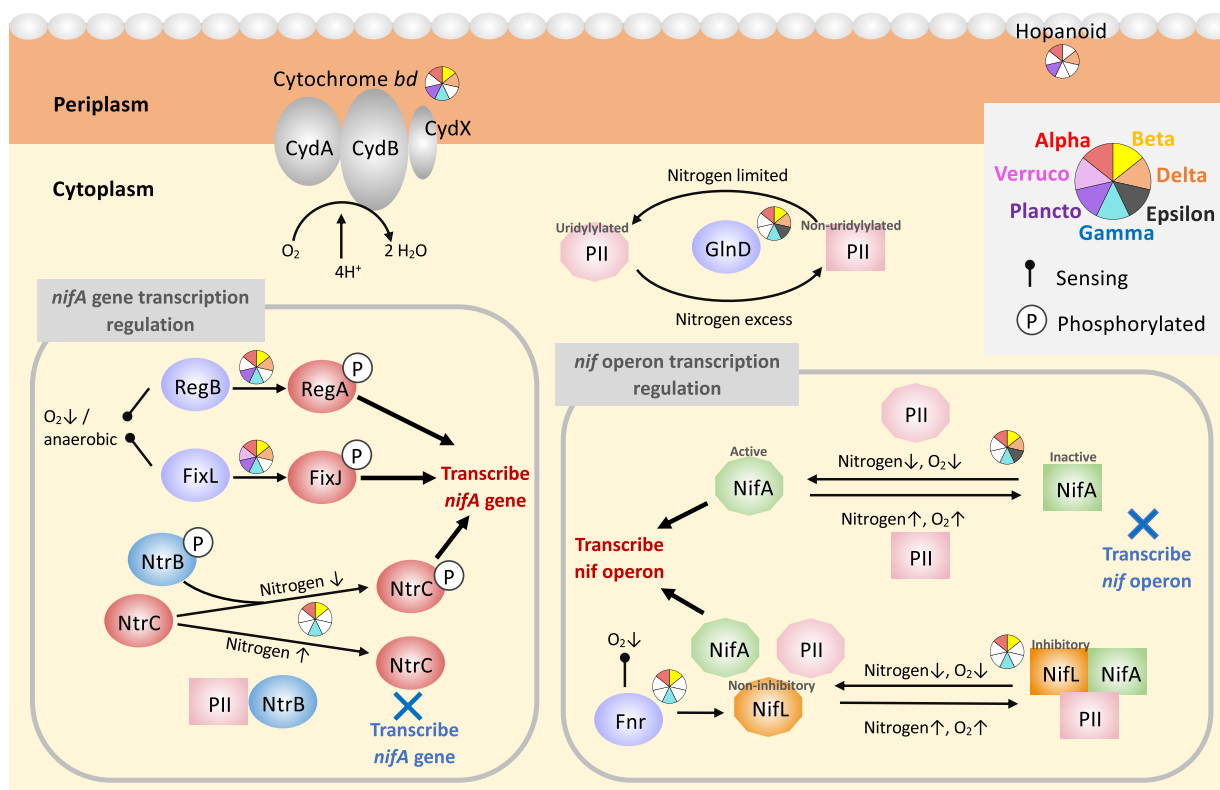


Figure 6. NCDs use diverse regulatory pathways for N_2 fixation. Circular diagrams next to each regulatory pathway represent the presence or absence of the corresponding genes in at least one MAG from each NCD group (Alpha = α -proteobacteria; Beta = β -proteobacteria; Delta = δ -proteobacteria; Epsilon = ϵ -proteobacteria; Gamma = γ -proteobacteria; Plancto = Planctomycetes; and Verruco = Verrucomicrobiota) as indicated by color (presence) or white (absence).

found to be important in reducing intracellular O_2 levels through respiration for aerobic N_2 fixation in *A. vinelandii* (Poole and Hill 1997). Additionally, of relevance to intracellular O_2 regulation, a recent study has proposed that hopanoids can act as an O_2 -diffusion barrier that protects the O_2 -sensitive nitrogenase in marine cyanobacteria (Cornejo-Castillo and Zehr 2019); however, the gene for hopanoid synthesis [squalene-hopene cyclase (*shc*)] was absent in most NCD MAGs, except Planctomycetes (Fig. 5C). Interestingly, many of these regulatory pathways were not detected in the MAGs of cyanobacterial diazotrophs (Fig. 5).

Acquisition and analysis of these MAGs provides unprecedented insight into the genomic potential of marine NCDs. The emerging pattern suggests that collectively this group may rely on alternative N sources in place of N_2 fixation under oxygenated conditions, as evidenced by the diverse N uptake proteins detected, along with the presence of N/ O_2 -sensing regulatory pathways. Future studies leveraging *in situ* whole genome transcription from NCDs are needed to link metabolic status (e.g. fixing N_2 vs. relying on N-uptake) with environmental conditions.

Moving from genes to rates: are NCDs fixing N_2 in the pelagic oceans?

The high diversity and wide distribution of NCDs and the occasional detection of *nifH* transcripts in PCR-based and metatranscriptomic datasets are often referenced as indicators of the significance of NCDs to marine N_2 fixation. Numerous attempts have been made to link NCD presence and/or active transcription of *nifH* to whole community N_2 fixation. This is challenging in the well-lit surface ocean given the generally low abundance of NCDs

relative to co-occurring cyanobacterial diazotrophs (Moisander et al. 2017). However, there are some reports of N_2 fixation rates where NCDs are abundant and cyanobacterial diazotrophs are absent (Yogev et al. 2011, Großkopf et al. 2012, Halm et al. 2012, Löscher et al. 2014, Shiozaki et al. 2014, 2017, Bentzon-Tilia et al. 2015b, Rahav et al. 2016), occasionally at high rates (e.g. $24.8 \pm 8.4 \text{ nmol N l}^{-1} \text{ d}^{-1}$; Löscher et al. 2014). But even when such direct comparisons are possible (e.g. NCD cell abundances and N_2 fixation rates are both reported), discrepancies often exist between NCD abundances and potential N_2 fixation rates (Turk-Kubo et al. 2014), suggesting underestimates of NCD abundances and/or cell-specific rates or overestimates of whole community N_2 fixation rates.

Proving that NCDs fix N_2 in aphotic waters is also challenging. The low rates reported for many aphotic samples ($<1 \text{ nmol N l}^{-1} \text{ d}^{-1}$) are near detection limits for the $^{15}N_2$ uptake assay, which have not historically been reported throughout the literature (Gradoville et al. 2017a). Reports of commercial $^{15}N_2$ gas stocks contaminated with $^{15}NH_4^+$ and/or $^{15}NO_3^-/^{15}N$ -nitrite (NO_2^-) raise a concern that some apparent rates could be artifacts driven by fixed N uptake (Dabundo et al. 2014). Additionally, measuring accurate N_2 fixation rates from the mesopelagic, where particulate N concentrations are low, can be logistically challenging due to the large filtration volumes needed to produce sufficient N content for mass spectrometry (White et al. 2020). Fortunately, the scientific community is now coming to a consensus regarding best practices for measuring and reporting N_2 fixation rates in the face of these challenges (White et al. 2020). However, more data validating active N_2 fixation by NCDs in aphotic waters is needed and linking rates to a particular NCD taxon remains a challenge.




	 free-living	 particle-/aggregate-associated	 symbiotic association
diversity	high	high	?
energy	C_{org} , light? inorganic compounds?	C_{org} , light? inorganic compounds?	C_{org} (host supplied), light?
carbon	C_{org} , C_{inorg} ?	C_{org} , C_{inorg} ?	C_{org} (host supplied)
N uptake	versatile N-acquisition	versatile N-acquisition	?
O₂ mitigation	↑ respiration ↓ EPS ↓ size?	↑ respiration ↓ EPS low O ₂ microenvironments	host controlled?
N₂ fixation	?	?	? host controlled?

Figure 7. Predicated features of euphotic marine NCD communities are likely influenced by habitats. Potential strategies used by NCDs to acquire energy and C needed to support N₂ fixation with energetic constraints and O₂ inactivation of nitrogenase. Question marks emphasize where many open questions remain. Notably, there is recent evidence that NCDs may be fixing N₂ on particles, as discussed in this review (Geisler et al. 2019). EPS—extracellular polymeric substances. Created with BioRender.com.

Measurements of taxonomically-resolved single-cell NCD N₂ fixation rates in marine waters are extremely sparse. Farnelid et al. (2014) isolated more than 60 strains of putative NCDs from Baltic Sea surface waters, more than half of which fixed N₂ under culture conditions. Single-cell N₂ fixation rates (up to 0.17 fmol N cell⁻¹ h⁻¹) of Baltic Sea isolates indicated that NCDs could be a locally-important N source if they fix N₂ at these rates in the environment given *in situ* abundances (10⁴ cells L⁻¹; Bentzon-Tilia et al. 2015a). Additionally, Martinez-Perez et al. (2018) isolated an α -proteobacterial NCD, *Sagittula castanea* (targeted by the qPCR assay CE1_45m_12_a, Zhang et al. 2011), from sulfidic waters in the upwelling region off the coast of Peru and successfully measured single-cell rates of 0.0008–0.060 fmol N cell⁻¹ d⁻¹ under culture conditions. N₂ fixation was only detected under anoxic conditions (despite being isolated from oxic waters) and was stimulated by NO₂⁻ and inhibited by NH₄⁺. Importantly, *in situ* single-cell N₂ fixation rates by *S. castanea* were undetectable, despite measurable bulk rates during the time of isolation, suggesting that *in situ* N₂ fixation by *S. castanea* may be transient and dependent on the development of anoxia.

Single-cell N₂ fixation rates from marine NCDs, both *in situ* and from nutrient perturbation experiments, are critically needed to validate their role in marine N₂ fixation and better understand environmental controls on their activity. Advances in single cell visualization techniques including CARD-gene-FISH (Moraru et al. 2010), mRNA-FISH (Pilhofer et al. 2009, McInnes et al. 2014), FISH-TAMB (Harris et al. 2021), and nitrogenase immunolabeling (Geisler et al. 2019, 2020), hold promise for measuring targeted NCD single-cell N₂ fixation rates if they could be successfully coupled to ¹⁵N₂ incubations and nanoscale secondary ion mass spectrometry (nanoSIMS). However, technical challenges arising from low cell concentrations in complex environmental samples and sample preparation steps that dilute the ¹⁵N signal (Meyer et al. 2021) require innovative solutions. Furthermore, given that NCD N₂ fixation may be a transient process, especially in particle microenvironments (Riemann et al. 2022), capturing active NCD N₂ fixation may currently be hampered by insufficient temporal and spatial sampling resolution (Benavides and Robidart 2020).

Future perspectives

The past two decades of research have greatly increased our understanding of marine NCDs and a framework for understand-

ing these organisms is now emerging (Fig. 7). Diverse NCD sequences have been detected from many ocean environments and habitats using both primer-based and primer-free approaches, underscoring their prevalence in the marine system. Insights from cultivation-based studies and marine NCD MAGs have revealed potential physiological strategies that marine NCDs may use to counter the inhibition of nitrogenase by O₂ and acquire enough energy to fuel N₂ fixation, such as forming self-aggregates in microaerophilic conditions and the ability to use alternative energy sources (Bentzon-Tilia et al. 2015a, Martinez-Perez et al. 2018, Acinas et al. 2021; Fig. 5A). Furthermore, reconstructing the metabolic potential of NCDs from MAGs shows that the phylogenetic diversity of this group is mirrored by diverse ecophysiological strategies. Marine NCDs have complex genetic systems to sense O₂ and N-availability and are able to use many alternative N sources, suggesting that some NCDs may be functioning as facultative diazotrophs. Collectively, these findings help explain why NCDs appear to inhabit such disparate habitats, from sunlit surface waters to the bathypelagic and to diverse benthic systems.

Despite these advances, many open questions remain (emphasized in Fig. 7). In addition to the critical need to demonstrate active N₂ fixation in marine NCDs (discussed in ‘Moving from genes to rates: are NCDs fixing N₂ in the pelagic oceans?’), better characterizing habitats and lifestyles of NCDs will be key to understanding the factors promoting active N₂ fixation. This includes determining whether NCD N₂ fixation is restricted only to low O₂ microhabitats (e.g. particle- or aggregate-associated), or if NCDs can fix N₂ aerobically (like some terrestrial counterparts) and if so, how they mitigate O₂ inactivation of nitrogenase. Another interesting question to resolve is whether some NCD taxa have formed obligate symbioses with other planktonic species, an evolutionary strategy that has led to many symbioses between cyanobacterial diazotrophs and algae (Villareal 1992, Thompson et al. 2012, Schvarcz et al. 2022).

At this time, the substantial knowledge gaps discussed throughout this review impede obtaining reasonable estimates of NCD contributions to the global marine N cycle, and the biogeochemical significance of NCDs remains uncertain. Ultimately, determining the importance of marine NCDs will require measurements of single cell N₂ fixation rates as well as advances in techniques to collect, enrich, characterize, and manipulate aggregates and particles, cultivate rare marine microbes, and increase tem-

poral and spatial resolution of N₂ fixation rates coupled to PCR-based and PCR-free surveys.

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Supplementary data

Supplementary data are available at [FEMSRE](#) online.

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