H$_2$ accumulation and N$_2$ fixation variation by Ni limitation in Cyanothece

Sing-how Tuo, Irene B. Rodriguez, Tung-Yuan Ho

Research Center for Environmental Changes, Academia Sinica, Taipei, Taiwan
Institute of Oceanography, National Taiwan University, Taipei, Taiwan

Abstract

Ni is an essential cofactor in NiFe-uptake hydrogenase, an enzyme regulating H$_2$ metabolism in diazotrophic cyanobacteria, the major H$_2$ producers in the surface ocean globally. Here, we investigated the effect of Ni supply on H$_2$ production and N$_2$ fixation by using a model marine cyanobacterial diazotroph, Cyanothece. By mediating total dissolved Ni concentrations from 100 to 0.03 nmol L$^{-1}$ in a trace metal-defined culture medium, we demonstrated that Ni deficiency results in H$_2$ accumulation, coupled with decreasing Ni quotas, growth rates, and occasionally relatively low N$_2$ fixation rates. These results indicate that Ni deficiency limits the growth of the Cyanothece to some extent, considerably decreases H$_2$ uptake by hydrogenase and leads to H$_2$ accumulation and N$_2$ fixation variation in the diazotroph. The findings show that Ni availability is a critical factor on controlling H$_2$ production and N$_2$ fixation in marine diazotrophic cyanobacteria. The information of Ni bioavailability for diazotrophic cyanobacteria is thus essential to evaluate the importance of Ni for H$_2$ cycling and N$_2$ fixation in oceanic surface waters.

Marine nitrogen-fixing cyanobacteria play important roles in regulating nitrogen and carbon cycling globally by transforming dinitrogen (N$_2$) to bioavailable nitrogen, ammonia (NH$_3$) (Karl et al. 2002; Bonnet et al. 2011). In addition to producing ammonia, hydrogen gas (H$_2$) is generated simultaneously as a byproduct through nitrogen fixation process as shown in Eq. 1:

$$\text{N}_2 + 8\text{H}^+ + 8\text{e}^- + 16\text{ATP} \rightarrow 2\text{NH}_3 + \text{H}_2 + 16\text{ADP} + 16\text{Pi}. \quad (1)$$

It has been widely observed that H$_2$ concentrations are supersaturated in the surface water of the tropical and subtropical oceans and the concentrations are closely related to cyanobacterial N$_2$ fixation activities. Early field studies observed that Trichodesmium, one of the major oceanic diazotrophic cyanobacteria, produces substantial amount of H$_2$ in the Caribbean Sea and the Atlantic Ocean (Scranton 1983, 1984; Scranton et al. 1987). In the North Pacific, Moore et al. (2009) also showed that net N$_2$ fixation rates exhibited a strong positive correlation with H$_2$ concentrations ($r = 0.96$), which ranged from 0.30 to 12.6 nmol L$^{-1}$ and were higher than its corresponding seawater equilibrium concentrations under H$_2$ atmospheric partial pressure. In the Atlantic Ocean, it was reported that H$_2$ concentrations were positively correlated with the abundances of Candidatus Atelocyanobacterium thalassa ($r = 0.77$), a dominant unicellular diazotrophic cyanobacterium (Moore et al. 2018). Laboratory culture studies also showed that prevalent marine diazotrophic cyanobacteria, including Trichodesmium, Crocosphaera, and Cyanothece, produced significant amount of H$_2$ during N$_2$ fixation process (Punshon and Moore 2008; Wilson et al. 2010a,b, 2012). In brief, both field and laboratory studies have demonstrated that cyanobacterial N$_2$ fixation is the major source of supersaturated H$_2$ in the upper ocean.

In terms of the stoichiometry in the nitrogen fixation reaction, the molar ratio of N$_2$ fixed to H$_2$ produced is 1:1 (Eq. 1). However, all of the measured ratios of net H$_2$ produced to N$_2$ fixed (hereafter the H$_2$:N$_2$ ratios) were less than 1 and varied significantly in both the field and laboratory studies. Field studies in the Caribbean Sea and the Atlantic Ocean (Scranton 1983, 1984; Scranton et al. 1987) reported that the H$_2$:N$_2$ ratio in Trichodesmium varied from 0.008 to 0.64. Laboratory studies showed that the H$_2$:N$_2$ ratios varied extensively for all of major diazotrophic cyanobacteria among different studies, such as 0.003–0.016 for Crocosphaera, 0.040–0.055 for Cyanothece, and 0.15–0.48 for Trichodesmium (Punshon and Moore 2008; Wilson et al. 2010a,b, 2012). The effort to apply H$_2$ production rates to estimate N$_2$ fixation rates, or vice versa, was hindered by the huge variations (Wilson et al. 2013; Moore et al. 2014). It has still remained unclear what environmental factors regulate the enormous variations and H$_2$ accumulation.

Since uptake hydrogenase oxidizes H$_2$ to proton and electron concurrently during N$_2$ fixation (Tamagnini et al. 2007), the oxidation would result in the decline of H$_2$ produced and
the molar ratio of H$_2$ produced to N$_2$ fixed to be less than 1. Indeed, hydrogen oxidation by uptake hydrogenase is beneficial to N$_2$ fixation since H$_2$ can inhibit nitrogenase activity (Guth and Burris 1983; Tamagnini et al. 2007). Moreover, uptake hydrogenase in unicellular diazotrophic cyanobacteria may also play a role in protecting nitrogenase from oxygen (O$_2$) toxicity (Zhang et al. 2014). In addition to uptake hydrogenase, diazotrophic cyanobacteria may contain a bidirectional hydrogenase, an enzyme generally existing in the nondiazotrophic cyanobacteria and being capable of regulating H$_2$ production and recycling with NAD(P)$^+$ and NAD(P)H reversibly (Tamagnini et al. 2007). However, the role of bidirectional hydrogenase on H$_2$ removal in diazotrophic cyanobacteria is not significant as uptake hydrogenase (Masukawa et al. 2002; Zhang et al. 2014). As cyanobacterial uptake hydrogenases are NiFe-bearing enzymes, Ni is expected to be a crucial factor for H$_2$ uptake in diazotrophic cyanobacteria. Indeed, previous freshwater culture studies showed that high Ni addition from 10 to 17,000 nmol L$^{-1}$ inhibited H$_2$ production for diazotrophic cyanobacteria Anabaena cylindrica and Oscillatoria sub brevevis (Daday and Smith 1983; Xiankong et al. 1984; Daday et al. 1985; Kumar and Polasa 1991). However, total dissolved Ni concentrations are generally around 2 nmol L$^{-1}$ in the surface waters of the tropical and subtropical open oceans (Bruland et al. 1979; Mackey et al. 2002; Wang et al. 2014; Schlitzer et al. 2018) so that the abnormally high Ni concentrations and nontrace metal buffered condition used in the freshwater study cannot reflect the impact of Ni deficiency on marine diazotrophic cyanobacteria. Although the Ni concentrations are around 2 nmol L$^{-1}$, the concentration level is one of the highest in comparison with other biologically essential trace metals in the surface water, such as Fe, Co, and Zn. It remains unclear whether 2 nmol L$^{-1}$ dissolved Ni level may limit the growth of some phytoplankton with Ni enzymes (Dupont et al. 2010; Ho 2013). Our previous laboratory culture experiments have demonstrated that Ni availability influences the growth and N$_2$ fixation of Trichodesmium (Ho 2013; Ho et al. 2013; Rodriguez and Ho 2014), which also show that a trace metal-defined culture medium with a low Ni background (Ho 2013) is essential to investigate the impact of Ni deficiency on marine phytoplankton.

Among major marine unicellular diazotrophic cyanobacteria (Luo et al. 2012; Cheung et al. 2017), Cyanothece strain ATCC 51142, hereafter Cyanothece, has been widely used as a model diazotrophic cyanobacterium for H$_2$ production studies (Bandyopadhyay et al. 2010, 2011; Min and Sherman 2010; Wilson et al. 2010a; Aryal et al. 2013). Its decoded genome also indicates that it possesses NiFe-uptake hydrogenases (Welsh et al. 2008; Bandyopadhyay et al. 2010). Most importantly, Cyanothece does not possess urase and Ni-superoxide dismutase (NiSOD) (Welsh et al. 2008) so that the effect of Ni deficiency is most likely manifest through effects on uptake hydrogenase. The major objective of this study is to investigate the effects of Ni deficiency on H$_2$ accumulation and N$_2$ fixation in Cyanothece and the implication on understanding the environmental control on H$_2$ cycling and N$_2$ fixation rates in the ocean.

**Methods**

*Cyanothece* sp. ATCC 51142 was obtained from American Type Culture Collection. Batch cultures of the strain were grown in 1-liter trace metal-clean polycarbonate Erlenmeyer flasks with 500 mL of trace metal-defined medium (Supporting Information Table S1) modified from YBCII recipe (Chen et al. 1996; Ho 2013). To prepare the trace metal-defined YBCII medium, we removed trace metal impurities from the artificial seawater base and working stocks of phosphate by passing the solutions through quartz column filling with chelating resins (Chelex$^\text{®}$ 100, Bio-Rad) in a clean room (Ho 2013). Sterilization and preparation for the medium followed the procedures for preparing trace metal-defined culture media (Price et al. 1989; Ho et al. 2003). All cultures were kept in an incubator with orbital shaking speed at 45 revolutions min$^{-1}$, temperature controlled at 26°C, and illumination at photon flux density of 450 μmol quanta m$^{-2}$ s$^{-1}$ operated at a 12:12 h light:dark square wave cycle.

To examine the effects of Ni under different Fe supplies, we conducted a 3 × 2 factorial design for the experiments. Three levels of total dissolved Ni supply, 0.03 nmol L$^{-1}$ (background concentration of the culture medium without Ni addition, low Ni treatment), 10 nmol L$^{-1}$ (medium Ni treatment), and 100 nmol L$^{-1}$ (high Ni treatment) were set under two levels of Fe addition, with total dissolved Fe concentrations to be 100 and 400 nmol L$^{-1}$. Trace metal availability was buffered by adding 20 μmol L$^{-1}$ ethylenediaminetetraacetic acid (EDTA). Each Ni-Fe treated culture was acclimated for around 20 generations (about 2 months) before performing the formal experiments (Supporting Information Table S2). Each treatment of batch culture had three replicate polycarbonate flasks for the formal experiments. In addition to the formal experiments, the samples obtained from the preliminary experiments during the acclimation process were also measured (Supporting Information Table S2).

We monitored cell density (cells mL$^{-1}$) and cell volume (μm$^3$ cell$^{-1}$) almost every day using a Multisizer 3 particle counter (Beckman Coulter). Specific growth rate (d$^{-1}$) was calculated using cell density data acquired during the late exponential growth phase, generally ranging from 10$^3$ to 2 × 10$^6$ cells mL$^{-1}$ (Fig. 1). Culture samples were concurrently processed during the late exponential phase for the measurement of the following parameters, including H$_2$ accumulation and N$_2$ fixation rates, elemental quotas, and chlorophyll $a$ (hereafter Chl $a$) content. The sampling time for each treatment was labeled with asterisk at the lower right position of the data point (Fig. 1). It should be noticed that the cells grown in the treatments with medium Ni-high Fe (all the replicates) and high Ni-low Fe (one of the replicates) concentrations had entered the
early stationary phase when collecting samples for the measurements (Fig. 1).

We measured the rates of H$_2$ accumulation and N$_2$ fixation by mercury oxide (HgO) reduction technique (Herr et al. 1981) and acetylene (C$_2$H$_2$) reduction (Capone and Montoya 2001), respectively. Two hours before the dark period started, each 10 mL aliquot of culture grown from each replicate flask under each treatment was transferred to a 20 mL vial and was crimp-sealed using Teflon-coated cap in a clean room. For H$_2$ measurements, the vials were incubated in the dark for 12 h. After incubation, H$_2$ concentrations were measured by injecting the gaseous samples into a 0.1 mL sample loop of a Peak Performer 1 Reducing Compound Photometer (Peak Laboratories) equipped with a heated HgO bed and UV absorption detector. For N$_2$ fixation assay, 2 mL of the headspace gas in the vials was sucked out and replaced with 2 mL of commercially available C$_2$H$_2$ by needles, then the vials were incubated in the dark for 12 h. After incubation, H$_2$ concentrations were measured by injecting the gaseous samples into a 0.1 mL sample loop of a Peak Performer 1 Reducing Compound Photometer (Peak Laboratories) equipped with a heated HgO bed and UV absorption detector. For N$_2$ fixation assay, 2 mL of the headspace gas in the vials was sucked out and replaced with 2 mL of commercially available C$_2$H$_2$ by needles, then the vials were incubated in the dark for 12 h. After incubation, 1 mL of the headspace gas from each vial was drawn for the measurement every 2 h during the incubation period. N$_2$ fixation was estimated by analyzing ethylene (C$_2$H$_4$) production using an Agilent 7890A gas chromatograph with a Poropak N column (Agilent, HP-PLOT Al$_2$O$_3$ S) and a flame ionization detector. The N$_2$ fixation rate was calculated using a reduction ratio (C$_2$H$_2$:N$_2$) of 4:1 for the conversion (Capone and Montoya 2001; Punshon and Moore 2008; Wilson et al. 2010a). Both the rates were normalized to cell volume subjected to the respective assays. We have used the data point of 12-h yield in the dark to calculate the daily rate (amol $\mu$m$^{-3}$ d$^{-1}$). The time-series hourly rates (amol $\mu$m$^{-3}$ h$^{-1}$) were also calculated by subtracting accumulated H$_2$/C$_2$H$_4$ concentrations observed at the preceding point then divided by the duration.

The elemental quotas were determined by collecting cells with 2 $\mu$m acid-washed polycarbonate membrane filters for P and trace metal analysis and precombusted GF/F filters for C and N analysis at the middle of the dark period. For phosphorus and trace metal analysis, the harvested cells were washed with ultrapure water to remove the culture medium residue in a class 100 trace metal clean laboratory. The filters with cells were digested using super pure concentrated nitric acid (ULTREX II, JT-Baker) in trace metal clean Teflon vials prior to elemental analyses using high-resolution inductively coupled plasma mass spectrometer (Element XR, Thermo Scientific). For C and N analysis, the harvested cells were fumed for 24 h with concentrated hydrochloric acid to remove particulate inorganic carbon prior to measurement by an elemental analyzer (2400 CHNS/O Series II, PerkinElmer). Filter blanks were subjected to the same digestion, dilution, and analysis. The blank values were subtracted from sample measurements. The details of the analytical precision, accuracy, and detection limits of the pretreatment and inductively coupled plasma mass spectrometry method for elemental measurement were described in our previous studies (Ho et al. 2003; Ho 2013). Cells for Chl a concentration measurements were harvested by filtration onto GF/F filters at the middle of the dark period. The filters were preserved in tissue embedding cassettes and were kept frozen at $-80^\circ$C until further processing. Water content from the filters was removed by freeze-drying the filters for 24 h prior to Chl a extraction in dark by using 90% acetone with sonication in an ice slurry for 1 h. Extracted Chl a samples were then filtered through a 0.2-$\mu$m filter to remove particulate debris and subsequently analyzed by high-performance liquid chromatography (LC-10A, Shimadzu). The Chl a concentrations measured include both mono-Chl a and

![Fig. 1. Growth curves of *Cyanothece* under various Ni and Fe treatments. The number next to elemental symbol (Ni) stands for total dissolved Ni concentration, which was either 0.03, 10, or 100 nmol L$^{-1}$. The Fe concentrations were either 100 or 400 nmol L$^{-1}$. Error bars stand for 1 SD from the average. For Ni100-Fe100 treatment, one replicate (R1) is not combined with the other two (R2 and R3) due to their unsynchronized growth paces. Asterisk below a symbol indicates the time point for the measurements of cell volume, Chl a contents, cellular elemental quotas, and rates of H$_2$ accumulation and N$_2$ fixation. [Color figure can be viewed at wileyonlinelibrary.com](image-url)
divinyl-Chl a. The details of the method were described in our previous study (Ho et al. 2015).

Data acquired at the early stationary phase, including one replicate of the treatment with high Ni-low Fe as well as all three replicates of the treatment with medium Ni-high Fe (the treatment as a missing cell) (Supporting Information Tables S3–S6), were not included into the statistical data sets for comparisons among different Ni and Fe treatments. Responses among different Ni and Fe treatments were compared using two-way ANOVA and post hoc Tukey’s studentized range test (only for the balanced data of specific growth rate) or Tukey–Kramer test. For such unbalanced data sets with the missing cell, a Type IV sum-of-square method was applied for ANOVA. If the interaction effect between Ni and Fe was significant ($p < 0.05$), then the post hoc test would be performed to investigate the difference among the Ni-Fe treatments. Relationships between the rates of $H_2$ accumulation and $N_2$ fixation as well as between the $H_2$:$N_2$ ratio and total dissolved Ni concentrations were tested with Pearson’s correlation coefficient ($r$). The data used for correlation test were log$_{10}$-transformed. Student’s $t$-tests were applied to compare the differences on elemental quotas between cells harvested at the early stationary and late exponential phases. All the data were presented as mean ± 1 standard deviation (SD). All statistical procedures were carried out by SAS program (SAS Institute).

**Results**

Cyanobacteria showed relatively low growth rates in the treatments without Ni addition, in which the growth rate (0.63 d$^{-1}$) was corresponding to 84% of maximal growth rates (0.75 d$^{-1}$) obtained from the medium Ni-high Fe treatment (Fig. 2a; Supporting Information Tables S3, S7). In the low Ni treatments, cell volume was larger than the medium and high Ni treatments (Fig. 2b; Supporting Information Tables S3, S7). We found that the effects of low Ni treatments on growth rates and cellular volume were repeatedly observed in the low Ni treatments in both formal experiments (Fig. 2a, b) and preliminary experiments (Supporting Information Fig. S1). No significant difference on Chl $a$ content was found among the treatments (Fig. 2c; Supporting Information Tables S3, S7).

Cellular elemental compositions are shown in Figs. 3, 4. Due to the significant variations of cell volume among different Ni treatments, elemental quota and rate parameters were presented by being normalized to cell volume. Except the treatment harvested at the early stationary phase, the averaged quotas of C, N, and P for all other treatments ranged from 8.4 to 13, from 2.4 to 2.7, and from 0.036 to 0.048, respectively (Fig. 3a–c; Supporting Information Tables S4, S7). No significant differences on the quotas of N and P were found (Fig. 3b,c; Supporting Information Tables S4, S7). In terms of trace metal quota, cellular Ni quota positively responded to Ni availability, possessing quotas ranging from 0.6 to 1.4 zmol $\mu m^{-3}$ for the low Ni treatments and from 29 to 34 zmol $\mu m^{-3}$ for the high Ni treatments and from 20.8 to 26.8 amol $H_2$ $\mu m^{-3}$ d$^{-1}$ were fivefold to twentyfold higher than those obtained in the higher Ni treatments, ranging from 1.1 to 4.0 amol $H_2$ $\mu m^{-3}$ d$^{-1}$ (Fig. 5a; Supporting Information Table S6). These trends on $H_2$ accumulation and $N_2$ fixation were also repeatedly observed in the preliminary experiments (Supporting Information Fig. S2). Although the correlation between the rates of $H_2$ accumulation and $N_2$ fixation was not significant for the formal experiment alone ($r = −0.30$, $n = 14$, $p = 0.29$), combining the data of preliminary and formal experiments all
together, the H$_2$ accumulation rate was negatively correlated with N$_2$ fixation rate ($r = -0.48$, $n = 24$, $p < 0.05$) (Fig. 6).

**Discussion**

The results of Fig. 2a demonstrate that Ni deficiency can limit the growth of *Cyanothece* to some extent, with growth rates decreasing 13–24% from the maximal observed growth rates. The condition of Ni deficiency was validated by the corresponding decreasing Ni quotas among the treatments (Fig. 4a). In addition to the impact on growth rate, our study revealed that Ni limitation in *Cyanothece* causes significant H$_2$ accumulation, with one order of magnitude higher accumulation rates in the low Ni treatments than medium and high Ni treatments (Fig. 5a). The results suggest that the production of NiFe-uptake hydrogenases was hindered owing to Ni deficiency.

The results of Figs. 5, 6 and Supporting Information Fig. S2 clearly show that Ni availability regulates H$_2$ accumulation and N$_2$ fixation in *Cyanothece*. As H$_2$ is a competitive inhibitor for nitrogenase (Guth and Burris 1983; Tamagnini et al. 2007), elevated H$_2$ accumulation caused by deficient Ni supply would concurrently hinder N$_2$ fixation. Under the condition of sufficient Ni supply, the ample production of NiFe-uptake hydrogenase would reduce H$_2$ level (Daday and Smith 1983; Xiankong et al. 1984; Daday et al. 1985; Kumar and Polasa 1991) and maintain N$_2$ fixation activities. The pooled daily based data showed a significant negative relationship between the rates of H$_2$ accumulation and N$_2$ fixation (Fig. 6). Similarly, we also observed that the variations of N$_2$ fixation rates and H$_2$ accumulation rates seem to be inverse during the dark period (Supporting Information Figs. S3, S4). Both the daily and hourly rates suggest that H$_2$ removal by uptake hydrogenase enhances nitrogenase activities.
and results in elevated N₂ fixation rates overall. Indeed, previous study clearly demonstrated that nitrogenase expression was hindered in \textit{Cyanothece} mutant with an uptake hydrogenase subunit knockout (Zhang et al. 2014), showing the essential role of uptake hydrogenase on cyanobacterial N₂ fixation. For the treatments with 400 nmol L⁻¹ Fe supply, N₂ fixation rates were all relatively low for all three Ni groups (Fig. 5b). It is unclear to us why the N₂ fixation rates were relatively low for the high Ni-high Fe treatment. In addition, it should be noticed that the daily N₂ fixation rates estimated from 12-h short-term integration may not be representative to calculate the overall biomass N due to various uncertainties involved (Supporting Information Text S1). In brief, under Ni deficiency condition, our experiments suggest that increasing H₂ accumulation rates would cause the decline of N₂ fixation rates.

Figure 7 shows that bioavailable Ni concentration is an important factor controlling the variations of the H₂:N₂ ratios in \textit{Cyanothece}. The highly varied ratios were negatively correlated with Ni concentrations (r = −0.71, n = 14, p < 0.01). A previous laboratory study showed significant differences on the ratio among various diazotrophic cyanobacteria and the authors argued that the two orders of magnitude differences reflected the ability of H₂ reutilization among the species (Wilson et al. 2010a). Supposedly, while expressing urease or...
NiSOD (Ho 2013), these two diazotrophs are expected to require more Ni than *Cyanothece* and exhibit higher H2:N2 ratios than *Cyanothece* if they all grow under the same level of low Ni supply. It should be noticed that the culture media used in the study (Wilson et al. 2010), either natural seawater-based SO (Waterbury and Willey 1988) or artificial seawater-based YBCII (Chen et al. 1996), were not added with Ni and were not trace metal defined so that the conditions of bioavailable Ni concentrations were possibly inconsistent in the experiments. It is thus essential to use a trace metal-defined culture medium to study N2 fixation rates or the H2:N2 ratios of diazotrophic cyanobacteria by laboratory culture experiments. The results of Fig. 7 imply that the H2:N2 ratios of some diazotrophic cyanobacterial species may also vary significantly in the field with different Ni availability in ambient seawater. The condition of Ni availability can be a potential cause for the huge variations of the H2:N2 ratios (0.008–0.64) observed in *Trichodesmium* colonies collected in the surface seawaters of the Caribbean Sea and the Atlantic Ocean (Scranton 1983, 1984; Scranton et al. 1987).

The results of Fig. 5a suggest that Ni availability can be an important factor causing the huge variations of H2 concentrations observed in oceanic surface water. Although both H2 concentrations and N2 fixation rates varied considerably in the tropical and subtropical oceans (Moore et al. 2009, 2018; Chen et al. 2014), field data show that total dissolved Ni concentrations are relatively comparable in the surface water (Bruland et al. 1979; Mackey et al. 2002; Wang et al. 2014; Schlitzer et al. 2018), around 2 nmol L\(^{-1}\). Unlike Fe and Cu which are strongly chelated by organic ligands in the surface water, studies showed that a significant portion (40–80%) of dissolved Ni in the surface water is labile (Achterberg and van den Berg 1997; Saito et al. 2004; Boiteau et al. 2016), which was operationally determined by the exchangeable dissolved
Ni with 12-h 20-μmol L\(^{-1}\) dimethylglyoxime treatment. It remains unclear whether this relatively weakly bound labile Ni are bioavailable to diazotrophic cyanobacteria or not (Ho 2013). Although the study of Boiteau et al. (2016) provided preliminary information about the elemental composition of Ni organic ligand extracted, the extracted fraction only accounted for 1–2% of total Ni concentrations. Further studies should focus on developing methods to investigate the structure and bioavailability of the majority of total dissolved Ni in the surface water. Then we may be able to assess whether Ni supply can serve as a critical variable for the H\(_2\) cycling and N\(_2\) fixation of diazotrophic cyanobacteria in the ocean.

In addition to Ni availability in seawater, biological Ni demand may vary significantly among different diazotrophic cyanobacterial groups, attributed to the difference of their intracellular Ni-containing enzymes. As mentioned previously, both *Trichodesmium* and *Crocospheara* contain NiFe hydrogenase, NiSOD, and urease (Dupont et al. 2008). It deserves further study to investigate their Ni requirement, the expressions of Ni-containing enzymes, and the impacts on N\(_2\) fixation and H\(_2\) production under various Ni supply in different oceanic regions. Enzymatic antibody approach and in situ Ni enrichment experiments may be applied to validate the importance of individual enzyme among different diazotrophic cyanobacteria. How Ni supply controls the dynamics of H\(_2\) concentration and N\(_2\) fixation in the ocean deserves further investigation.

**Conclusion**

Ni limitation in *Cyanothece* resulted to elevated H\(_2\) accumulation and caused the variations on N\(_2\) fixation rates. These results indicate that Ni deficiency decreases H\(_2\) uptake and occasionally N\(_2\) fixation, leading to a decline in growth rate while cellular N quota keeps constant. Our findings suggest that H\(_2\) supersaturation in the upper ocean may thus be associated with Ni bioavailability for diazotrophic cyanobacteria. Ni bioavailability in oceanic surface water would be essential information to evaluate the role of Ni on H\(_2\) cycling and N\(_2\) fixation for marine diazotrophic cyanobacteria. Moreover, in situ Ni enrichment experiments can be carried out to demonstrate the importance of Ni on cyanobacterial H\(_2\) cycling and N\(_2\) fixation in the ocean, which can be elucidated via examining the expressions of uptake hydrogenase and nitrogenase.

**References**


Tamagnini, P., E. Leitão, P. Oliveira, D. Ferreira, F. Pinto, D. J. Harris, T. Heidorn, and P. Lindblad. 2007. Cyanobacterial hydrogenases: diversity, regulation and applications. FEMS


Acknowledgments
We thank Wan-Yen Cheng, Jie-Cheng Chang, Chih-Chiang Hsieh, and Wan-Chen Tu for providing technical support. This study was financially supported by grant MOST 105-2119-M-001-039-MY3 by Taiwan Ministry of Science and Technology and Career Development Award CDA-104-M11 by Academia Sinica.

Conflict of Interest
None declared.

Submitted 12 February 2019
Revised 12 June 2019
Accepted 25 July 2019

Associate editor: James Moffett