Nitrogen-starvation-induced chlorosis in *Synechococcus* PCC 7942: adaptation to long-term survival

Margit Görl, Jörg Sauer, Tina Baier and Karl Forchhammer

When deprived of essential nutrients, the non-diazotrophic cyanobacterium *Synechococcus* sp. strain PCC 7942 undergoes a proteolytic degradation of the phycobiliproteins, its major light-harvesting pigments. This process is known as chlorosis. This paper presents evidence that the degradation of phycobiliproteins is part of an acclimation process in which growing cells differentiate into non-pigmented cells able to endure long periods of starvation. The time course of degradation processes differs for various photosynthetic pigments, for photosystem I and photosystem II activities and is strongly influenced by the illumination and by the experimental conditions of nutrient deprivation. Under standard experimental conditions of combined nitrogen deprivation, three phases of the differentiation process can be defined. The first phase corresponds to the well-known phycobiliprotein degradation, in phase 2 the cells lose chlorophyll a prior to entering phase 3, the fully differentiated state, in which the cells are still able to regenerate pigmentation after the addition of nitrate to the culture. An analysis of the protein synthesis patterns by two-dimensional gel electrophoresis during nitrogen starvation indicates extensive differential gene expression, suggesting the operation of tight regulatory mechanisms.

**Keywords:** cyanobacteria, dormant state, phycobiliproteins, photosynthesis, proteolysis

**INTRODUCTION**

Cyanobacteria are one of the most widespread groups of Gram-negative bacteria. They display considerable morphological diversity and unusual capacities for cellular differentiation, while their basic metabolism is fairly uniform. Cyanobacteria are capable of photoautotrophic growth, performing oxygenic photosynthesis, similar to that of eukaryotic algae and plants. Moreover, they employ highly efficient mechanisms to adapt to changes in ambient conditions (for a review see Tandeau de Marsac & Houmard, 1993). One of the essential requirements to survive in natural environments is the ability to withstand nutrient limitation or even deprivation. In the absence of combined nitrogen sources, diazotrophic cyanobacteria avoid nitrogen deficiency by fixing molecular nitrogen. Non-diazotrophic strains respond to nitrogen deprivation by degrading their photosynthetic pigments, resulting in a change in the colour of cultures from blue-green to yellow, a process known as chlorosis (Allen & Smith, 1969; Lau et al., 1977). Chlorosis also occurs upon starvation for other essential nutrients, although the chlorotic response shows subtle differences at the cellular level, depending on the limiting nutrient (Collier & Grossman, 1992; Wanner et al., 1986).

Of the different chlorotic reactions, that induced by nitrogen starvation has been most extensively studied. Most investigations have focused on the degradation of the major light-harvesting complexes, the phycobilisomes, which proceeds in an ordered manner (reviewed by Grossman et al., 1994). A small polypeptide (NblA) was identified that triggers the proteolytic degradation of phycobiliproteins; however, its precise function is still unknown (Collier & Grossman, 1994). In various studies, the time course and extent of phyocyanin (PC) degradation has varied from only 50% degradation of PC per volume of culture within 60 h (Lau et al., 1977) to 90% degradation within 9 h (Wanner et al., 1986). Wanner et al. (1986) showed that the decay of PC was...
accompanied by a dramatic degradation of chlorophyll a (chl a), whereas in other investigations the chl a content during chlorosis remained constant (Allen & Smith, 1969) or decreased only slightly (Collier & Grossman, 1992). However, these studies were performed with cells grown under different CO₂ concentrations and illumination intensities. Even different experimental procedures of induction of combined nitrogen deprivation (step-down) could have contributed to differences in the results. Nitrogen step-down was performed by one of two methods. Nitrate-grown cells were inoculated into a medium that contained limiting amounts of nitrate. The cells proliferated until the nitrogen source was exhausted and then started to bleach (Allen & Smith, 1969; Wanner et al., 1986). Alternatively, cells were harvested in the exponential growth phase by centrifugation, washed and re-suspended in a medium lacking a combined nitrogen source. This procedure allows a better definition of the onset of nitrogen starvation (Lau et al., 1977; Collier & Grossman, 1992). Here, the cells are able to divide once before growth ceases. Upon nitrogen starvation, cells accumulate products of photosynthesis that are stored as glycogen (Allen, 1984); however, a thermophilic *Synechococcus* strain was described that accumulates up to 27% poly-β-hydroxybutyrate per dry weight (Miyake et al., 1997). Within 1–3 d of the onset of nitrogen starvation, cyanobacteria are able to initiate growth upon a shift to nutrient-replete conditions (Allen, 1984, Wanner et al., 1986). A decline of the cellular chl a content following the initial chlorosis reaction has often been interpreted as an irreversible decay of cell viability (Tandeau de Marsac & Houmard, 1993). However, the capacity to survive longer starvation periods has not been investigated. Prolonged starvation of unicellular microorganisms often leads to cellular differentiation, which may result in the generation of dormant spores, as is found in many Gram-positive bacteria. In contrast, in various non-sporulating heterotrophic bacteria, a small subpopulation of non-dormant survivors differentiates, which may remain viable for years (Zambrano & Kolter, 1996; Watson et al., 1998). Here we demonstrate that the photoautotrophic organism *Synechococcus* PCC 7942 survives prolonged periods of nitrogen deprivation, by differentiating into non-pigmented cells that show characteristics of a dormant state.

**METHODS**

**Organism and culture conditions.** *Synechococcus* sp. strain PCC 7942, small plasmid cured (Kühlemeier et al., 1983), hereafter designated *Synechococcus* PCC 7942, was grown photoautotrophically in BG-11 medium (Rippka, 1988) that was modified as follows: ferric ammonium citrate was replaced by ferric citrate and the medium was buffered to pH 7.8 with 20 mM HEPES. Medium lacking combined nitrogen (BG-11) corresponded to BG-11 medium, but NaNO₃ was replaced by the same molarity of NaCl (17.6 mM). Stock cultures (40 ml) in 100 ml Erlenmeyer flasks were incubated at 30 °C without additional aeration and illuminated with a photosynthetic photon flux density (PFD) of approximately 10 μmol photons m⁻² s⁻¹ from fluorescent lamps. The photosynthetically active radiation was determined with a quantum sensor (LI-190SA, LI-COR). For starvation experiments, cultures were grown in 200 ml BG-11 in one litre Erlenmeyer flasks with magnetic stirring and aerated either with air, or with air supplemented with 2% CO₂; standard illumination conditions at a PFD of 50–60 μmol m⁻² s⁻¹ were provided from fluorescent lamps. For the determination of numbers of c.f.u., appropriate dilutions of liquid cultures were plated on BG-11 medium solidified by the addition of 0.9% (w/v) of Gel-Rite (Roth) and incubated at 30 °C with a PFD of 30 μmol m⁻² s⁻¹.

**Initiation of nitrogen deprivation.** A 100 ml volume of exponentially growing cells was harvested at OD₅₇₀ 0.5 by filtering the culture through a 0.45 μm PVDF membrane filter (Millipore, type HVLP) in a transparent filtration unit under constant illumination (55 μmol m⁻² s⁻¹). Care was taken not to dry out the cell layer on the filter. The cell layer was washed twice with 5 ml BG-11 medium and the cells were re-suspended in 200 ml BG-11 medium to give an OD₅₇₀ of 0.25; the incubation was continued in one litre Erlenmeyer flasks under the same growth conditions. The amount of evaporated water was determined daily by weighing and the corresponding volume of distilled sterile water was added to the culture.

**Regeneration experiments.** To assay the regeneration of photosynthetic pigments in individual cells, 5 ml samples were withdrawn from nitrogen-deprived cultures into sterile 25 ml flasks, supplemented with 17.6 mM sodium nitrate and 25 μg aztreonam ml⁻¹, and incubated at a PFD of 40 μmol photons m⁻² s⁻¹. Addition of aztreonam, an inhibitor of cell septation, was repeated after 3 and 5 d. Regeneration of pigments was observed by epifluorescence microscopy. To concomitantly visualize the non-regenerated cells, cells were stained with DAPI (4′,6′-diamino-2-phenylindole).

**Pigment and glycogen determination.** PC, chl a and glycogen were determined as previously described (Forchhammer & Tandeau de Marsac, 1995). The unit for PC determination (net A₅₇₀) results from the difference spectrum of cultures before and after heating to 75 °C for 8 min and is calculated according to the following equation: net A₅₇₀ = [A₅₇₀ (unheated)] - [A₅₇₀ (heated)] (Collier & Grossman, 1992).

**Measurement of photosynthetic activities.** Photosynthetic oxygen evolution was measured using a Clark-type oxygen electrode (Hansatech DW1) as described by Walker (1987). Light was provided from a high-intensity white-light source (Hansatech LS2) and light intensity was adjusted using neutral density filters. Oxygen consumption or evolution in 2 ml samples was measured at the following PFDs: 0, 7, 12, 24, 82 and 3000 μmol m⁻² s⁻¹. PS II activity was determined in intact cells by measuring oxygen evolution in a Hill reaction after addition of 0.5 mM ferricyanide and 0.1 mM benzoquinone using saturating actinic light of 3000 μmol m⁻² s⁻¹ (Shen et al., 1993). PS I activity was estimated by measuring the light-induced oxygen uptake (Mechler reaction) in permeabilized cells using the protocol of Haukka (1980), adapted for use with cyanobacteria. To 4 ml cell suspension (with an OD₅₇₀ of 0.25–0.5), 20 μl 1 M HEPES pH 8.0, 20 μl 5 M NaCl, 100 μ1 0.1 M MgCl₂, 80 μl 1 mM DCMU [3-(3,4-dichloro-phenyl)-1,1-dimethylurea] in ethanol, 4 μl 1 M DTT, 133 μl 1 M Na₂CO₃, 10 μl 1 M t-arabonic acid and 133 μl 0.75% (w/v) CTAB (cetyltrimethylammonium bromide) were added and mixed thoroughly for 30 s. After a 5 min incubation at 28 °C under constant shaking, 40 μl 10 mM benzyl viologen and 10 μl N,N,N′,N′-tetramethyl-p-phenylenediamine were
added and incubated for another 10 min; 2 ml of the suspension was transferred to the chamber of the oxygen electrode and oxygen consumption was recorded in the dark. Then, saturating actinic light (3000 μmol m⁻² s⁻¹) was supplied and the light-stimulated oxygen consumption was determined.

**Epifluorescence microscopy.** Epifluorescence microscopy was performed with a Zeiss Axiosplan epifluorescence microscope equipped with a camera (MC100, Zeiss). Cells were stained with the LIVE/DEAD BacLight bacterial viability kit (Molecular Probes) according to the manufacturer’s instructions. Briefly, 1 μl of each reagent A and B were diluted in 660 μl PBS (137 mM NaCl, 2.7 mM KCl, 10.1 mM Na₂HPO₄ and 1.8 mM KH₂PO₄). Cells were harvested by centrifugation and 8 × 10⁵ cells resuspended in 50 μl of the dye solution. After 10 min incubation in the dark, 15 μl cell suspension was spread onto a microscope slide covered with a dried layer of 2% agarose that immobilized the cells. For simultaneous visualization of live and dead cells, the Zeiss 487909 filter combination (365 nm excitation, 490 nm dichroic and >520 nm emission filter) was used. DAPI staining of cells was performed according to Coleman (1980) and the stained cells were examined using the Zeiss 487901 filter combination (365 nm excitation, 395 nm dichroic and >397 nm emission filter), which allowed the simultaneous detection of the blue fluorescence from DAPI-stained chlorotic (non-autofluorescent) cells and the red autofluorescence of fully pigmented cells.

**In vivo [³⁵S]methionine labelling and two-dimensional gel electrophoresis.** In vivo labelling of proteins in nitrogen-depleted cells was performed by removing 8 ml samples from nitrogen-deprived cultures into sterile 50 ml Erlenmeyer flasks, the samples were supplied with 25 μCi [³⁵S]methionine (43.5 TBq) and incubated as before for the times indicated (see Fig. 6 legend). The cells were then chilled on ice, harvested by centrifugation, washed twice with ice-cold 80% acetone and resuspended in 120 μl lysis buffer [8 M urea; 0.2%, v/v, DTT; 0.02%, w/v, Pefablock (Boehringer Mannheim)]. Incorporation of [³⁵S]methionine into proteins was quantified by determining the radioactivity from a 10 μl aliquot after precipitating proteins with 10% TCA. For each electrophoretic analysis, a sample volume corresponding to 0.6 × 10⁵ c.p.m. was loaded, unless otherwise indicated. Gel electrophoresis in two dimensions was carried out according to Görg et al. (1995). The first dimension was separated on 14 cm immobilized pH gradients (IPG dry strips, pH 4–7), using the multiphor flatbed electrophoresis system together with the immobiline dry strip kit (Pharmacia). The second-dimension gels were all run together on 16 × 20 cm Tricine SDS 10% acrylamide gels (Schägger & van Jagow, 1987). Gels were silver stained (Blum et al., 1987), dried, exposed to phosphor screens (Molecular Dynamics) for 2–4 d and scanned with the PhosphorImager system (Molecular Dynamics) at 100 μm resolution.

**RESULTS**

**Nitrogen-starvation-induced pigment degradation under different experimental conditions**

In previous reports, the time course and extent of PC and chl a loss that was observed upon depriving *Synechococcus* PCC 7942 cells of combined nitrogen varied considerably. Therefore, we examined the influence of different experimental conditions on pigment degradation. To precisely determine the onset of nitrogen deprivation, we intended to use cells in the exponential growth phase and resuspend them in nitrogen-deficient medium. We had previously observed that centrifugation caused a global distortion of cellular physiology, of protein synthesis patterns and of photosynthetic activities (J. Sauer, M. Gör & K. Forchhammer, unpublished). For a minimal impact on cell physiology, cells were harvested by filtration through a 0.45 μm PVDF filter in a transparent filtration unit, which allowed continued illumination during the procedure. Under these conditions, degradation of PC was nearly complete 24 h after nitrogen depletion (Fig. 1c, d), whereas cultures that were shifted by centrifugation still retained 40 to 50% PC per culture volume (Fig. 1a, b). However, no significant influence of the shift procedure could be observed with respect to the chl a content of the culture (Fig. 1a, c). With both procedures, the CO₂ supply was found to have a major influence on chl a degradation; in the presence of 2% CO₂, chl a degradation was significantly retarded compared to cultures supplied with air (0.03% CO₂). The time course of the chlorotic response depended further on the PFD: PC degradation increased in parallel with the increase in light intensity and the chl a content decreased even more rapidly at higher light intensities (data not shown).

**Long-term nitrogen-deprivation and survival of *Synechococcus* PCC 7942**

*Synechococcus* PCC 7942 cells were shifted to medium lacking combined nitrogen and were incubated under standard conditions (2% CO₂, 50 μmol photons s⁻¹ m⁻²) for up to 30 d. At various times, aliquots were removed and analysed for PC, chl a and glycogen. The OD₅₇₀ was measured and numbers of c.f.u. after plating dilutions on nitrate-supplemented solid BG11 medium were also determined (Table 1). Based on the results of these experiments, three phases of the nitrogen-deprivation response could be defined. Phase 1 corresponded to the chlorotic response, which is characterized by a rapid decline of PC while the chl a content of the culture only slightly decreased. The cells divided once and glycogen accumulated. Under our experimental conditions, this phase occupied the first 2 d of nitrogen deprivation. Phase 2 started when the PC degradation was complete. During phase 2, the chl a content of the culture decreased progressively, reaching nearly undetectable values; the carotenoid content also decreased during this phase (data not shown). The colour of the culture turned from yellowish-green to pale white during this phase, which lasted about 8–10 d. The decay of the membrane-bound photosynthetic pigments was accompanied by a moderate reduction of the OD₅₇₀ of the culture and by a decline in the numbers of c.f.u., whereas the glycogen content remained at a high level. Finally, the cells entered phase 3, where no further changes in the appearance of the culture were observed. In phase 3, pigments were almost undetectable and the glycogen content remained at a high level. Surprisingly, the number of c.f.u. on nitrate-supplemented medium remained at a relatively low, but constant, level. However, a microscopic examination of the nitrogen-starved cells showed that cells tended to form aggregates which could...
distort the determination of c.f.u. numbers. To more precisely examine the viability of long-term nitrogen-deprived cultures, the cells were stained with the ‘LIVE/DEAD BacLight’ viability kit, which allows differentiation between living and dead cells on the basis of their membrane potential. By epifluorescence microscopy, dead cells show a red fluorescence whereas living cells fluoresce green. Cells from a culture starved for 1 d had already lost the typical cyanobacterial red autofluorescence and appeared green upon staining with ‘LIVE/DEAD BacLight’. As a control for dead cells, an aliquot was boiled for 5 min; these cells showed an orange-red fluorescence (Fig. 2a, b). The intensity of the green fluorescence faded in phase 2 cells and a small portion of red fluorescent cells became visible. Surprisingly, in samples of phase 3 cultures, almost all cells retained a green fluorescence (Fig. 2c), indicating that the cells still exhibited membrane potential. This result suggested that the majority of cells remained viable. Therefore, we developed a new method to analyse the regeneration potential of the nitrogen-deprived cells. Nitrogen-replete cyanobacteria display a red autofluorescence, which originates from PS II and phyco-biliproteins. As a consequence of the degradation of photosynthetic pigments and of quenching effects (Collier et al., 1994), nitrogen-deprived cells rapidly lose their red autofluorescence. The reappearance of the autofluorescence after the addition of nitrate indicates the regeneration of their pigments. Since a small portion of viable cells could overgrow the culture during regeneration experiments, the antibiotic aztreonam, which specifically inhibits septum formation (Zambrano & Kolter, 1996), was added to avoid this. At different times after the addition of nitrate and aztreonam to chlorotic cultures, aliquots were analysed by fluorescence microscopy. To allow the simultaneous visual-

**Fig. 1.** Effect of experimental procedures initiating nitrogen deprivation and of the CO₂ supply on the level of PC and chl a per culture volume and OD₅₇₀ of Synechococcus PCC 7942. The 100% values correspond to the values before the shift. Representative results are shown. ▼, OD₅₇₀; ○, relative PC level ml⁻¹; ●, relative chl a level ml⁻¹. Cells were shifted to nitrogen-deprived medium by centrifugation (a, b) or by filtration (c, d). Cultures were supplied with either 2% CO₂ in air (a, c) or with air alone (b, d). The absolute values corresponding to 100% were: PC, 0.1 net A₅₇₀; chl a, 1.8 µg ml⁻¹.
Table 1. Change in the levels of PC, chl a and glycogen, in OD750 and in c.f.u. per culture volume during the course of nitrogen deprivation

<table>
<thead>
<tr>
<th>N-deprivation (d)</th>
<th>Percentage PC*</th>
<th>Percentage chl a*</th>
<th>Percentage glycogen*</th>
<th>Percentage OD750*</th>
<th>Percentage c.f.u. (±SE)</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>100</td>
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<tr>
<td>1</td>
<td>5</td>
<td>87</td>
<td>ND</td>
<td>198</td>
<td>ND</td>
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<tr>
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<td>220</td>
<td>176 ± 15</td>
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<tr>
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<td>0</td>
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<td>370</td>
<td>198</td>
<td>154 ± 15</td>
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<tr>
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<td>140</td>
<td>96 ± 12</td>
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<td>110</td>
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<tr>
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<td>420</td>
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<td>8 ± 4</td>
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<td>&lt;1</td>
<td>410</td>
<td>73</td>
<td>7 ± 4</td>
</tr>
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* The SE for the determination of PC, chl a, glycogen and OD750 in a single experiment was below 5%.

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The results are from one of three independent experiments and are expressed as percentages of the value before nitrogen step-down. The time course of the degradative processes in the different experiments varied by about 10% but proceeded in a similar manner. ND, Not done. The absolute values corresponding to 100% were: PC, 0.102 net A420; chl a, 18 µg ml⁻¹; glycogen, 25 µg ml⁻¹; OD750, 0.25; c.f.u, 5.2 x 10⁶ ml⁻¹.

The decomposition of the non-regenerated cells, the samples were counter-stained with the blue fluorescent dye DAPI. Fig. 3 shows a typical experiment with cells that had been nitrogen-depleted for 20 d. In different experiments, about 5–10% of the cells recovered the red autofluorescence within the first 2 d. Following a lag phase of another 2–3 d with no significant increase of regeneration, nearly all cells regained red autofluorescence after a total regeneration time of 4–5 d.

Photosynthetic activities during nitrogen deprivation

To define any relationship between nitrogen deprivation and photosynthesis, we assayed photosynthesis-related activities during chlorosis. At various times following nitrogen depletion, light-dependent oxygen exchange was assayed in the dark and at several PFDs to estimate dark respiration, quantum efficiency and Vmax of complete photosynthesis. In cells that were shifted to nitrate-deprived medium, a rapid decline in photosynthetic capacity was observed (Fig. 4a). Six hours after the shift, a 50% decline in Vmax became evident and after 24 h, the quantum yield decayed strongly as evidenced by the decreased slope of the light-dependent region. Photosynthetic oxygen evolution vanished almost completely after 72 h. PS II activity was determined at saturating light (3000 µmol s⁻¹ m⁻²) using the Hill reaction (Fig. 4b). PS II activity followed the decline of Vmax with some delay (Fig. 4b). PS I activity was measured as light-induced oxygen uptake (Mehler reaction) in cells in which PS II activity was inhibited by DCMU (Hauska, 1980) (Fig. 5). Although this method only allows a rough estimate of PS I activity, it is evident that PS I activity decays much more slowly than light-dependent oxygen evolution or PS II activity. Only after 4 d of nitrogen starvation was a decay of PS I activity apparent, which paralleled the degradation of chl a. PS I activity was undetectable after 16 d of nitrogen starvation.

Protein synthesis in nitrogen-deprived cells

To obtain a preliminary insight into global changes of gene expression associated with nitrogen deprivation, we investigated the protein composition and new protein synthesis during the different phases of nitrogen starvation using two-dimensional gel electrophoresis. At various times following nitrogen deprivation, aliquots were removed from the chlorotic cultures and labelled with [35S]methionine. Subsequently, cell extracts were analysed by electrophoresis (Fig. 6). About 5 h after the shift to nitrogen-deprived medium, the synthesis of a multitude of proteins appeared to be affected, when compared to the synthesis pattern of nitrogen-replete cells. Computer-assisted evaluation of the autoradiograms identified at least 70 proteins that were strongly induced and about 50 which decreased in intensity, following nitrogen deprivation (data not shown). The protein synthesis pattern of cells in phase 2 showed a significant reduction of the number of spots. Interestingly, several spots appeared only after long-term starvation (some representatives are indicated by rectangles in Fig. 6). In phase 3, the labelling efficiency decreased dramatically. Only very few proteins could be identified which were synthesized under these conditions. Similar to the labelling patterns, the silver-stained gels showed a strong reduction of the protein inventory during phase 2 of chlorosis, reaching a minimal protein composition in the phase 3 cells (data not shown).
Fig. 2. Epifluorescence microscopy of *Synechococcus* PCC 7942 cells after staining with the ‘LIVE/DEAD BacLight’ viability kit. (a) Cells after 1 d nitrogen deprivation; (b) the same cells after boiling for 5 min; (c) cells after 28 d nitrogen deprivation. Bar, 10 μm.

Fig. 3. Regeneration of nitrogen-deprived *Synechococcus* PCC 7942 cells visualized by epifluorescence microscopy. Cells that had been starved for nitrogen for 20 d were regenerated by the addition of nitrate in the presence of aztreonam; aliquots were removed at various times, stained with DAPI and analysed by epifluorescence microscopy. Regenerated cells display a bright-red autofluorescence, whereas chlorotic cells show the blue DAPI fluorescence. (a) Before the addition of nitrate, (b) 2 d, (c) 3 d, (d) 4 d after the addition of nitrate. Bar, 10 μm.
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**Fig. 4.** (a) Photosynthetic oxygen evolution of *Synechococcus* cells at increasing PFDs after 0, 0.2, 2, 6, 24, 48, 72 and 120 h of nitrogen starvation. (b) PS II activity was determined at saturating light intensity by the use of the Hill reaction from the same cultures as those in (a).

**Fig. 5.** PS I activity measured as light-induced oxygen uptake (Mehler reaction) in permeabilized cells after nitrogen deprivation.

**DISCUSSION**

We have presented evidence that depletion of the non-diazotrophic cyanobacterium *Synechococcus* PCC 7942 for combined nitrogen causes a differentiation process that enables the cells to survive long periods of starvation. Based on the temporal sequence of adaptive responses, this differentiation process can be divided into three phases, reminiscent of three stages of the starvation-survival process described in heterotrophic bacteria (Morita, 1993). In the nitrogen-starvation response of *Synechococcus* PCC 7942, these phases are characterized by the following events. In phase 1, the cells divide once, while the phycobiliproteins are degraded to near completion. Photosynthetic oxygen evolution decayed after 1 d of nitrogen deprivation, as previously reported (Allen et al., 1990); PS I activity remained unaffected during this time (Collier et al., 1994). The decay of the $V_{\text{max}}$ of photosynthesis preceded the decay of PS II activity. This result implies that the reduction of photosynthetic electron transport is caused by an impairment in the metabolic reactions linked to photosynthesis. Since no nitrogen is available to the cells for synthesis of nitrogen-containing metabolites, the flow of photosynthate is channelled into glycogen synthesis. Indeed, Coronil et al. (1993) showed that immediately after nitrogen depletion, *Synechococcus* cells reduce the flow of newly fixed CO$_2$ into the amino acid pool. Therefore, the amino acids for new protein synthesis must originate from protein turnover, most prominently from phycobiliprotein degradation. Of the about 70 starvation-induced spots that could be visualized by two-dimensional gel electrophoresis of pulse-labelled cells, some are specific for nitrogen depletion but others also appear under conditions of sulfur deprivation (J. Sauer & K. Forchhammer, unpublished).

In addition to the reduction in the synthesis of the phycobiliproteins by nitrogen starvation, an effect previously demonstrated by Lau et al. (1977), the synthesis of a variety of other proteins was also strongly inhibited. Factors involved in sensing nutrient starvation in cyanobacteria have not yet been identified at the molecular level. The signal-transduction protein P$_{\text{II}}$ might be required in responses specific for nitrogen deprivation, since its phosphorylation state is maximal under conditions of nitrogen deprivation (Forchhammer & Tandeau de Marsac, 1995). Whether P$_{\text{II}}$ is indeed involved in the nitrogen deprivation response is currently under investigation. In phase 1, cells can quickly regenerate after the addition of a combined nitrogen source (Allen, 1984). The resynthesis of phycobiliproteins has been
suggested to be controlled at the level of transcription by the chromophore supply (Gilbert et al., 1996).

Prolonged nitrogen depletion causes *Synechococcus* PCC 7942 cells to differentiate further into a non-photosynthetic state. At the beginning of phase 2, PS II is already inactive but the cells can still generate ATP by PS I activity. The fact that most cyanobacterial chl a is associated with PS I (Manodori et al., 1984; Bryant 1986) implies that chl a degradation reflects the loss of PS I reaction centres. Concomitantly, a breakdown of the photosynthetic membranes, the thylakoids, can be visualized in these cells by electron microscopy (Wanner et al., 1986; G. Wanner, A. Bock & K. Forchhammer, unpublished). At the same time, the protein content of the cells decreases greatly, most likely by proteolytic processes. New protein synthesis still takes place, although at a reduced level, as deduced from the long
labelling period required to achieve sufficient incorporation of $[^{35}]$methylene. The synthesis pattern indicates an extensive alteration of gene expression. In a variety of bacteria, gene expression in stationary phase is altered due to the activity of specific sigma factors (Hengge-Aronis, 1993; Antelmann et al., 1997). In the genomic sequence of the unicellular cyanobacterium Synechocystis PCC 6803, at least eight genes encoding putative sigma factors have been found (Kaneko et al., 1996) and in the marine cyanobacterium Synechococcus PCC 7002, the presence of at least five sigma factors has been demonstrated, two of which are expressed under conditions of carbon and nitrogen starvation (Caslake et al., 1997). Whether a special sigma factor operates in the nitrogen-deprivation response of Synechococcus PCC 7942 remains to be demonstrated.

After completion of the degradative processes, Synechococcus cells enter phase 3 of chlorosis, which exhibits features of a dormant state. This contrasts with the stationary-phase physiology in various heterotrophic bacteria, where under prolonged starvation the number of surviving cells initially decreases exponentially and finally stabilizes at a low level. Such cultures are in a dynamic equilibrium between growth and death (Kolter, 1992). A dormant-like state in Synechococcus PCC 7942 is deduced from the following observations. (i) No photosynthetic activities could be measured in long-term starved cells. Although it is tempting to speculate that these chlorotic cells maintain a very low PS I activity that could energize the cells by cyclic electron transport, the method employed in our investigation was not sensitive enough to detect very low PS I activities after long-term-starvation periods. (ii) Respiration was abolished and the glycogen reserves were not utilized, indicating that they did not contribute to the energy supply of the cells. (iii) Only traces of protein synthesis could be detected, which was limited to a minor subset of proteins. If a small number of survivors were to live on lysis products, one would expect a protein synthesis pattern resembling that of growing cells. (iv) Analysis of single-cell viability suggested that the majority of cells maintained membrane potential and retained the capacity to regenerate in liquid medium. This resembles the 'viable but non-culturable' state, a term that is used when starved but viable cells fail to yield colonies on solid medium regeneration (discussed by Bloomfield et al., 1998). The molecular basis of the dormant state and of the regeneration processes merits further investigations. The capacity of Synechococcus PCC 7942 cells to sustain prolonged starvation periods could represent an important mechanism for cyanobacterial populations to survive in natural environments.

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