Dark Anaerobic Hydrogen Production in the Mutants of *Synechocystis* sp. Strain PCC6803-GT Defective in Lactate Dehydrogenase Activity and/or Alcohol Dehydrogenase Activity, Incubated in Buffer Solutions With or Without Glucose

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Received: September 5, 2014   Accepted: September 22, 2014   Online Published: October 20, 2014
doi:10.5539/ijb.v7n1p33         URL: http://dx.doi.org/10.5539/ijb.v7n1p33

Abstract

To redirect NAD(P)H consumed by dehydrogenases to NiFe-hydrogenase, the genes encoding for lactate dehydrogenase (*ddh*) and alcohol dehydrogenase (*adh*) were disrupted in a glucose tolerant mutant of unicellular cyanobacterium *Synechocystis* sp. strain PCC6803 (GT strain). The cells of GT strain, *ddh* deficient (*Δddh*) mutant and both *adh* and *ddh* deficient (*ΔadhΔddh*) mutant were harvested at the late-logarithmic growth phase of photoautotrophic culture in antibiotic-free BG-11 medium. Dark anaerobic hydrogen production in GT strain, *Δddh* mutant and *ΔadhΔddh* mutant in nitrate-free HEPES buffer solution without or with glucose has been studied by following the time courses of the number of moles of hydrogen, endogenous and exogenous glucose, lactate, acetate and ethanol per culture volume and the dry cell weight concentration. Dark incubation of cells of *Δddh* mutant and *ΔadhΔddh* mutant in HEPES buffer solution without glucose resulted 1.16-fold and 1.15-fold increases in the initial hydrogen production rates over GT strain, respectively, while 1.1-fold and 0.85-fold increases in the number of moles of hydrogen per culture volume at 96 h over GT strain, respectively. When hydrogen production experiments were performed in the glucose-added HEPES buffer solution, dark incubation of cells of *Δddh* mutant and *ΔadhΔddh* mutant resulted 1.4-fold and 1.6-fold increases in the initial hydrogen production rates over GT strain without glucose run, respectively, while 1.5-fold and 1.3-fold increases in the number of moles of hydrogen per culture volume at 96 h over GT strain without glucose run, respectively.

Keywords: alcohol dehydrogenase, cyanobacteria, dark anaerobic, glucose, hydrogen production, lactate dehydrogenase, mutant, *Synechocystis*

1. Introduction

The glucose tolerant mutant of unicellular cyanobacterium *Synechocystis* sp. strain PCC6803 (GT strain) is transformable and represents both photoautotrophic growth on carbon dioxide and heterotrophic growth on glucose (Rippka et al., 1979; Kufryk et al., 2002). During phototrophic growth, carbon dioxide is incorporated into ribulose bisphosphate (RuBP) in the Calvin cycle to form 3-P-glycerate that is further converted to glucose in glycogen for storage utilizing the ATP and NADPH produced by the light-dependent reactions. During heterotrophic growth, exogenous glucose is transported into the cell through glucose-fructose transporter and phosphorylated to form glucose 6-phosphate that is also converted to glycogen. Generations of ATP and NAD(P)H by glycogen decomposition by either aerobic respiration or anaerobic glycolysis have been applied for molecular hydrogen production on bidirectional NiFe-hydrogenase (Antal and Lindblad, 2005; Cournac et al., 2004). If hydrogen production in the dark is associated with breakdown of glycogen to pyruvate via oxidative pentose phosphate (OPP) pathway, glucose 6-phosphate dehydrogenase, 6-phosphogluconate dehydrogenase and glyceraldehyde-3-phosphate dehydrogenase contribute a supply of NAD(P)H for hydrogenase. Different from those enzymes, under the fermentative condition, D-lactate dehydrogenase (*ddh*) assimilates NADH by converting pyruvate to D-lactate (hereafter lactate) and alcohol dehydrogenase (*adh*) assimilates NADPH by converting acetaldehyde to ethanol. The *ddh* for lactate production and the *adh* for ethanol production compete NAD(P)H with NiFe-hydrogenase for hydrogen production. If disruption of the genes coding for a NAD(P)H-dependent enzyme results a variation in redox balance of GT strain, such mutation appears to present a good outlook for increasing hydrogen production.
A previous work on the growth of mutant of *Thermoanaerobacterium* strain defective of L-lactate dehydrogenase genes (ldh) on 100 μmol mL⁻¹ xylose shows the 2- and 2.5-folds increases in hydrogen production yield and hydrogen production rate, respectively (Li et al., 2010). When grown on glucose, hydrogen fermentation of ldh-deleted *Escherichia coli* was superior to that of wild type strain (Maeda et al., 2007). Dark incubation of a cyanobacterium *Synechococcus* sp. strain PCC7002 and that lacking the enzyme for the NADH-dependent reduction of pyruvate to D-lactate (AdhDHA) results 11 and 55 mole hydrogen per 10¹⁷ cells (=0.0384 and 0.215 μmol mL⁻¹), respectively (McNeely et al., 2010). In our previous publication, we reported the successful variation in the hydrogen production of GT strain by disrupting the genes coding for *ddh* with aiming at elevating the level in the amount of intracellular NADH for bidirectional hydrogenase (Yamamoto et al., 2012). The keyword search on the genome database of *Synechocystis* sp. strain PCC6803 utilizing CyanoBase (http://genome.kazusa.or.jp/cyanobase/) hits 41 genes for NADH and 9 genes for NADPH. Among them this study concerns *adh* genes in slr0942 that is a manager of carboxyls received from glycolysis system (Shimakawa et al., 2013). *Synechocystis* sp. strain PCC6803 can naturally convert pyruvate to acetate that is further reduced via adh to a small amount of ethanol in dark anaerobic condition. There is a report that shows a construction of a mutant of *Synechocystis* sp. strain PCC6803 with genes coding for *adh* and pyruvate decarboxylase (*pdc*) of *Zymomonas mobilis* and a production of 120 μmol mL⁻¹ ethanol phototrophically from carbon dioxide (Gao et al., 2012; Dehring et al., 2012). Although aldehyde is toxic to *Synechocystis* sp. strain PCC6803, cells are protected by converting it to ethanol.

In order to gain more insight into the effect of elimination of single or multiple NAD(P)H consumption reactions that compete NAD(P)H with NiFe-hydrogenase in dark anaerobic nitrate-free condition, in the present work, we disrupt not only genes coding for D-lactate dehydrogenase (*ddh*) but also genes coding for alcohol dehydrogenase (*adh*) in a glucose tolerant mutant of *Synechocystis* sp. strain PCC6803 and compare the hydrogen production of GT strain, *ddh* deficient (*Δddh*) mutant with both *adh* and *ddh* deficient (*ΔadhΔddh*) mutant in the HEPES buffer solution without and with glucose. The *adh* gene and *ddh* gene are selected and targeted to be deleted in order to modify the size of protein to be expressed and to alter the homeostasis network to generate NAD(P)H for NiFe-hydrogenase pathway.

### 2. Material and Method

#### 2.1 Mutant Constructions

The competent cells of *Escherichia coli* X1-Blue were used for transformation purposes. *E.coli* cells were cultured at 37 °C in 2 mL LB medium in 24.5 mL clear Pyrex glass test tube (TE-32, AGC TECHNO GLASS Co. Ltd.) which were shaken at 220 rpm in a shaking incubator. The wild-type host cyanobacterium used in this study was a GT strain, ddh deficient (*Δddh*; Prentki and Krisch, 1984), hence the disruption of *ddh* gene resulted in the 0.56 kb increase in the size of DNA. The *Δddh* mutant lacking both *adh* and *ddh* (slr1556::Km) was constructed by ligating PCR-amplified *ddh* fragment into a plasmid pUC19 at *Smal*. The product was then digested with *BglI* before the ligation of kanamycin resistance gene cassette from pUC4K. Sizes of *ldh* gene and *Km* gene were 1.002 kb and 0.543 kb, respectively, hence the disruption of *ldh* gene results in the 0.459 kb decrease in the size of DNA. The *ΔadhΔddh* mutant lacking both *adh* and *ddh* (slr1556::Km, slr0942::Sm/Sp) was constructed by ligating PCR-amplified *adh* fragment into a plasmid pMD19 at *EcoRV* site. The streptomycin/spectinomycin resistance gene cassette (*Sm*/*Sp*) from plasmid pHP45-Ω was inserted into the *EcoRV* restriction site on *adh* gene fragment. Sizes of *adh* gene and *Sm*/*Sp* gene were 0.981 kb and 2.0 kb (Prentki and Krisch, 1984), hence the disruption of *ldh* gene and *adh* gene resulted in the 0.56 kb increase in the size of DNA. Experimental technique to decrease or increase DNA size results in decrease or increase in the amount of NAD(P)H that is required for protein synthesis.

Transformant colonies of *Δddh* mutant and *ΔadhΔddh* mutant were selected from the plates containing kanamycin and both kanamycin and streptomycin, respectively, and continuously re-streaked to new plates. Antibiotic resistant cells were grown on BG-11 medium plates at increasing antibiotic concentrations and finally transferred into liquid medium (BG-11). The cells of *Δddh* mutant and *ΔadhΔddh* mutant were grown on BG-11 medium supplemented with 60 μmol mL⁻¹ kanamycin, and that with both 60 μmol mL⁻¹ kanamycin and 2.5 μmol mL⁻¹ streptomycin, respectively. Mutant cells were harvested and washed to remove antibiotics before the inoculation into to the photoautotrophic main cultures in the light.
2.2 Photoautotrophic Growth for Inoculum Preparation

Cells of GT strain, ∆ddh mutant and ∆adhΔddh mutant of *Synechocystis* sp. strain PCC6803 were grown photoautotrophically in 60 mL antibiotic-free BG-11 medium (initial pH 7.8) that was aerated by 6% carbon dioxide in the air in clear Pyrex glass test tube (TE-32, AGC TECHNO GLASS Co. Ltd.) at 34 ºC in water bath, surface of which was exposed to fluorescent lamps with a photosynthetic photon flux density (PPFD) of 100 µmol-phots m⁻² s⁻¹. Growth was monitored by measuring OD₇₃₀ of cell suspension. One unit of OD₇₃₀ represents the dry cell weight concentration (X) of 0.369 mg mL⁻¹. In preliminary experiments it was observed that cells were in late-logarithmic growth phase when OD 730 of cell suspensions was 7. Cultures of GT strain, ∆ddh mutant and ∆adhΔddh reached these OD730 cultivations, respectively. Cultures of OD730 at 7 were centrifuged at 3000 rpm for 10 min at 25 ºC.

2.3 Dark Anaerobic Incubation for Hydrogen Production

The cell pellets were washed, re-suspended and incubated in 10 mL of either 50 mmol L⁻¹ HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid)) buffer (pH 7.8) or 50 mmol L⁻¹ HEPES buffer (pH 7.8) with 5 mmol L⁻¹ glucose in 32 mL glass test tube with butyl rubber cap. HEPES was selected for pH control preventing a drastic drop in pH during fermentation. The initial dry cell mass weight concentration was set at 2 mg mL⁻¹ and cell suspension was purged with nitrogen gas for a few minutes to remove oxygen gas. Incubation of cell suspension in test tube was carried out under dark anaerobic conditions, shaking at 145 rpm in a reciprocating shaker. The reciprocating distance was 40 mm and the horizontal angle was about 30 ºC. All experiments were carried out in duplicate.

The number of moles of hydrogen per culture volume (yH₂) was determined by analyzing the composition of gas phase in 32 mL test tube utilizing gas chromatograph and thermal conductivity detector (GC-320, GL science Inc.; column, Molecular sieve 13X; carrier gas, nitrogen gas; column temperature, 37 ºC; injector temperature, 45 ºC; detector temperature 80 ºC).

The amount of cellular glycogen was estimated in terms of the number of moles of glucose equivalent. The number of moles of endogenous glucose per dry cell weight (mG) was determined as follows: the amount of dry cell weight per culture volume was determined according to previous report (Yamamoto et al., 2012); cell pellets were collected by centrifugation at 25 ºC, 3000 rpm for 10 min then washed 3 times by deionized water to eliminate extracellular carbon sources; intracellular glycogen was extracted and decomposed to glucose in 50 µL of 6 N HCl at 80 ºC for 30 min; glucose amount was determined by using Glucose CII Test Wako (Wako Pure Chemical Ind., Ltd.). The number of moles of endogenous glucose per culture volume (mG,X) was calculated from mG and X. For experiments with exogenous glucose, the number of moles of glucose per culture volume (cS) was analyzed by HPLC. Exogenous glucose dissolved in supernatant was analyzed by HPLC (Shimadzu LC-10AD and RID-6A) with a 6.0 × 150 mm column SZ5532 (Showa Denko Co.). Mobile phase was 70% acetonitrile with water. Mobile phase flow rate was 0.7 mL/min. Column temperature was set at 50°C.

Extracellular concentrations of lactate (cL) and acetate (cA) in a 500 µL portion of supernatants were analyzed by HPLC (JASCO PU-2080 Plus and UV-2075 Plus). Mobile phase was 18 mmol L⁻¹ KH₂PO₄, pH of which was adjusted at 2.3 with H₃PO₄. Flow rate of mobile phase was 0.7 mL min⁻¹. Column temperature was 30 ºC. A 10 µL of supernatant sample was injected by using 10 µL syringes. The wave length of 210 nm was used for detection of metabolites. Ethanol (cE) in supernatant was analyzed by HPLC (Shimadzu LC-10AD and RID-6A) with a 6.0 × 150 mm column SZ5532 (Showa Denko Co.). Column temperature was set at 50°C. Mobile phase was 5 mmol L⁻¹ sulfuric acid.

3. Results

3.1 Construction of ∆ddh Mutant and ∆adhΔddh Mutant

Figure 1 represents the partial restriction maps of (a) the *ddh* in slr1556 cloned into the *SmaI* site of pUC19, the slr1556::Kmᵣ containing the kanamycin resistance gene inserted into the *BglII* site in the *ddh* in slr1556 and (b) the *adh* in slr0942 cloned into the *EcoRV* site of pMD19 and the slr0942::Smᵣ/Spᵣ containing the streptomycin/spectinomycin resistance gene inserted into the *EcoRV* restriction site on *adh* gene. Wild-type *Synechocystis* sp. strain-GT was transformed with slr1556::Kmᵣ and with both slr1556::Kmᵣ and slr0942::Smᵣ/Spᵣ.
Photographs in Figure 2 show results of agarose-gel electrophoretic analyses of PCR products for evaluating the achievement of deletion of (a) ddh genes and (b) both adh and ddh genes. Photograph (a) shows Lane 1: size marker DNA, Lane 2: PCR product for ddh primers and pUC19-slr1556:Km', Lane 3: PCR product for ddh primers and genomic DNA from GT strain, and Lane 4: PCR product with ddh primers and genomic DNA from Δddh mutant. Photograph (b) shows Lane 1: size marker DNA, Lane 2: PCR product for ddh primers and pUC19-slr1556:Km', Lane 3: PCR product for adh primers and pMD19-slr0942::Sp'Smr', Lane 4: PCR product for ddh primers and genomic DNA from GT strain, Lane 5: PCR product for adh primers and genomic DNA from GT strain, Lane 6: PCR product for ddh primers and DNA from ΔadhΔddh mutant and Lane 7: PCR product for adh primers and genomic DNA from ΔadhΔddh mutant. The successful segregation of Δddh is confirmed, ((a) Lane 4 or (b) Lane 6) is the same size as that for primers and DNA fragment from pUC19-slr1556:Km' ((a) or (b) Lane 2). Double mutation is also confirmed by the result showing that the size of PCR product for adh primers and genomic DNA from ΔadhΔddh mutant ((b) Lane 7) is the same size as that for primers and DNA fragment from pMD19-slr0942::Sp'Smr' (b) Lane 3).

Figure 2 Agarose-gel electrophoretic analyses of PCR products for evaluating the achievement of (a) Δddh mutation and (b) ΔadhΔddh mutation

3.2 Characteristics of Photoautotrophic Growth

Precultures for photoautotrophic growth of GT strain, Δddh mutant and ΔadhΔddh mutant were performed on BG-11 medium, BG-11 medium containing 60 μmol mL⁻¹ kanamycin and BG-11 medium containing 60 μmol mL⁻¹ kanamycin and 2.5 μmol mL⁻¹ streptomycin, respectively. At 96 h of precultures, cells were harvested, washed and then inoculated into main cultures in which cells were grown photoautotrophically in antibiotic-free BG-11 medium. Growth of cells on BG-11 was monitored by optical density at 730 nm wavelength. Growth curves for GT strain, Δddh mutant and ΔadhΔddh mutant show that cells of GT strain and Δddh mutant were in logarithmic growth phase from 0 h to 24 h and from 0 h to 40 h, respectively, while ΔadhΔddh mutant was in lag phase from 0 h to 24 h and in logarithmic growth phase from 24 h to 48 h (Figure 3).
Figure 3 Photoautotrophic growth curves of GT strain (○), Δddh mutant (■) and ΔadhΔddh mutant (▲) on antibiotic-free BG-11 medium. Pre-culture cells of GT strain, Δddh mutant and ΔadhΔddh mutant were grown on BG-11 medium, BG-11 medium with 60 mg mL⁻¹ kanamycin and BG-11 medium with both 60 mg mL⁻¹ kanamycin and 2.5 mg mL⁻¹ streptomycin, respectively. Cells were washed by sterilized distilled-water to remove antibiotics prior to main culture. The horizontal line pointing the level of $X$ at 2.58 mg mL⁻¹ represents the culture-harvest ion time when OD₇₃₀ of cell suspensions is 7

The lag time for the growth of ΔadhΔddh mutant appears to represent a period for adaptation of the mutant from antibiotic containing preculture medium to antibiotic-free main culture medium. After one week cultivation, dry cell weight concentrations of Δddh mutant and Δadhddh mutant reached the same level as GT strain.

Carbon dioxide is a single carbon source for cell growth, hence the slope of growth curve represents the activity of carbon dioxide fixation. Total amount of carbon dioxide incorporated into dry cell weight during 168 h cultivation is found not to be affected by deletion of adh or ddh genes. The inoculum cells of GT strain, Δddh mutant and ΔadhΔddh mutant for dark anaerobic hydrogen production were harvested at 36, 45, and 60 h, respectively.

The OD₇₃₀ at the cell-harvest ion time was fixed, hence inoculum cells for hydrogen production experiment were confirmed to be exposed to same PPFD from incident light to transmitted light. The slope of growth curves (dlnX/dt) of GT strain, Δddh mutant and ΔadhΔddh mutant of the $X$ at 2.58 mg mL⁻¹ are 0.0187, 0.0318 and 0.0238 h⁻¹ respectively. The cell formula of Synechocystis sp. strain PCC6803 shown in a reference is CH₁₆₂N₀.₂₂O₀.₄₀P₀.₁₁ (Yu et al., 2013), hence the rates of carbon dioxide assimilation by inocula of GT strain, Δddh mutant and ΔadhΔddh mutant are calculated as 2.09, 3.55 and 2.66 µmol mL⁻¹ h⁻¹, respectively and the rates of NADPH consumption at the Calvin cycle are calculated as 4.18, 7.10 and 5.32 µmol mL⁻¹ h⁻¹, respectively.

### 3.3 Characteristics of Dark Anaerobic Nitrate-Free Hydrogen Production in the Absence of Glucose

In the first series, photoautotrophic cells of GT strain, Δddh mutant or ΔadhΔddh mutant in late-logarithmic growth phase was incubated at 2.58 mg mL⁻¹ dry cell weight concentration in dark anaerobic nitrate-free HEPES buffer solution without glucose. Similar to photoautotrophic culture, the solution for hydrogen production was free from antibiotics. Figure 4 shows the time courses of the number of moles per culture volume of (a) hydrogen ($y_{H₂}$) and (b) endogenous glucose in glycogen ($mGX$), and the concentrations of (c) dry cell weight ($X$), (d) lactate ($c_l$) and (e) acetate ($c_A$). Ethanol concentration of run with GT strain, Δddh mutant or ΔadhΔddh mutant in the absence of glucose was not in detectable level, hence time course data for it were not plotted in Figure 4.

Hydrogen production characteristics in the absence of glucose: If subscript 1 shows the state at 96 h, the $y_{H₂,1}$ values of runs with GT strain, Δddh mutant and ΔadhΔddh mutant are 1.8, 2.0 and 1.5 µmol mL⁻¹, respectively (Figure4(a)). The $y_{H₂,1}$ of Δddh mutant is 1.11 times that of GT strain. The $y_{H₂,1}$ of Δadh mutant shown in our previous report (Yamamoto et al., 2012) was 1.5 times that of GT strain. Previous work and present work utilized the inocula from logarithmic growth phase and late-logarithmic growth phase, respectively. Growth phase is suggested to be of utmost important factor for hydrogen production. The elimination of adh activity resulted to decrease $y_{H₂,1}$ that was 0.83 times that of GT strain, The highest $y_{H₂,1}$ was observed in run with Δddh mutant, second highest is in run with GT strain and the lowest is in run with ΔadhΔddh mutant. This descending order is same as the ascending order of the modified DNA size shown in Table 1. The DNA size of ΔadhΔddh mutant is 0.56 kb larger than that of GT strain, hence expression of increased DNA size appears to increase the NAD(P)H consumption for protein and amino acid synthesis and reduce the hydrogen production. On the other hand, the DNA size of Δddh mutant is 0.459 kb smaller than that of GT strain, hence expression of decreased DNA size
appears to decrease the NAD(P)H consumption for protein and amino acid synthesis and increase the hydrogen production.

The time courses of $y_{H2}$ show that the production of hydrogen by those three strains was highest right after inoculation and deactivated with time. The $y_{H2}$ versus time data fit the equation:

$$y_{H2} = \frac{y_{H2,f}}{k} \left[ 1 - \exp(-kt) \right]$$

in which $y_{H2,f}$ is the attainable level of $y_{H2}$, $k$ is the deactivation constant for hydrogen production and $r_{H2,0}$ is the initial hydrogen production rate. The curves in Figure 4(a) for GT strain, Δddh mutant and ΔadhΔddh mutant are calculated with the values of $y_{H2,f}$ and $k$ in Table 1. The calculated $y_{H2}$ agrees well with the observed $y_{H2}$. The descending order of $y_{H2,f}$ in Table 1 is same as the descending order of the $y_{H2,1}$.

The $r_{H2,0}$ of Table 1 is highest for run with Δddh mutant, comparable second highest for run with ΔadhΔddh mutant and lowest for run with GT strain. The descending order of $r_{H2,0}$ is similar to that of the consumption rate of NAD(P)H in the Calvin cycle just before inoculation (Table 1). Shift of cellular environment from illuminated aerobic nitrate-containing BG-11 medium to dark anaerobic nitrate-free HEPES buffer solution appears to redirect the accumulated NAD(P)H molecules in the Calvin cycle to NiFe-hydrogenase.

Table 1. Kinetic parameters of dark anaerobic hydrogen production in GT strain, Δddh mutant and ΔadhΔddh mutant in HEPES buffer solution without or with glucose

<table>
<thead>
<tr>
<th>Strain</th>
<th>Increase in DNA size [kb]</th>
<th>NAD(P)H production rate in the Calvin cycle before inoculation [μmol mL⁻¹ h⁻¹]</th>
<th>HEPES buffer additives</th>
<th>$r_{H2,0}$ [μmol mL⁻¹ h⁻¹]</th>
<th>$k$ [h⁻¹]</th>
<th>$y_{H2,f}$ [μmol mL⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>GT strain</td>
<td>0</td>
<td>4.18</td>
<td>w/o glucose</td>
<td>0.0309</td>
<td>0.0167</td>
<td>2.28</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>with glucose</td>
<td>0.0901</td>
<td>0.0184</td>
<td>5.61</td>
</tr>
<tr>
<td>Δddh mutant</td>
<td>-0.495</td>
<td>7.10</td>
<td>w/o glucose</td>
<td>0.0359</td>
<td>0.0213</td>
<td>2.31</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>with glucose</td>
<td>0.0430</td>
<td>0.0145</td>
<td>3.48</td>
</tr>
<tr>
<td>ΔadhΔddh mutant</td>
<td>0.56</td>
<td>5.32</td>
<td>w/o glucose</td>
<td>0.0355</td>
<td>0.0365</td>
<td>1.48</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>with glucose</td>
<td>0.0502</td>
<td>0.0299</td>
<td>2.45</td>
</tr>
</tbody>
</table>

The elimination of ddh activity or both adh and ddh activities results in 1.16-fold increase or 1.15-fold increase in the $r_{H2,0}$. Considering that the NiFe-hydrogenase converts 1 μmol NAD(P)H to form 1 μmol hydrogen, this result shows that the amount of NADH consumed by lactate dehydrogenase of GT strain is about 16% of the total amount of NADH and NADPH consumed by NiFe-hydrogenase of GT strain. The initial lactate production rate was 0.000833 μmol mL⁻¹ h⁻¹ in GT strain and almost 0 μmol mL⁻¹ h⁻¹ in Δddh and ΔadhΔddh mutants (Figure 4d). Stoichiometry of D-lactate dehydrogenase and NiFe-hydrogenase suggests that the elimination of above lactate production rate potentially increases hydrogen production rate from 0.0309 μmol mL⁻¹ h⁻¹ to 0.0359 μmol mL⁻¹ h⁻¹. The observed increase is 6 times more than the estimated increase by ddh elimination. Redirection of carbon metabolism at pyruvate from lactate production to acetyl-CoA production is found to amplify NAD(P)H production.

A 1.28-fold increase in Δddh mutant and a 2.19-fold increase in ΔadhΔddh mutant over GT strain in the deactivation constant $k$ are seen in the results (Table 1). The presence of dehydrogenases in GT strain appears to be important to stabilize the dynamical redox adaptation of cells against continuous consumption of NAD(P)H on NiFe-hydrogenase.

The attainable level of $y_{H2}$ by Δddh mutant is comparable with that of GT strain, while that by ΔadhΔddh mutant is 0.65 times that of GT strain. For hydrogen production in the absence of exogenous glucose, stabilization is of first importance for high production of hydrogen by ΔadhΔddh mutant. Our result of Δddh mutant in Synechocystis sp. strain PCC6803 differs from that of ΔddhA mutant in Synechococcus sp. strain PCC7002 (McNeely et al., 2010) that shows 5-fold increase in hydrogen production from wild type strain, however the attainable level of hydrogen ($y_{H2,f}$) in this experiment is 10 times that shown by their experiment.

Endogenous glucose consumption characteristics in the absence of glucose: The $m_{c}X$ of runs with three strains decreased or kept constant with time (Figure 4b)). Although hydrogen productions in three strains were highest at
initial, extremely low decrease in the \( m_c X \) at initial was seen in run with \( \Delta ddh \) mutant. The initial rates of endogenous glucose consumption of GT strain, \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) mutant were 0.00208, 0.000417 and 0.00125 \( \mu \text{mol mL}^{-1} \text{ h}^{-1} \), respectively. The initial hydrogen production of mutants is unrelated to endogenous glucose consumption. The decrease in the \( m_c X \) with time is highest from 24 to 48 h in runs with GT strain and \( \Delta ddh \) mutant, and from 0 to 24 h in run with \( \Delta adh\Delta ddh \) mutant. The yields of hydrogen on endogenous glucose (\( Y_{H2G} \)) in GT strain and in \( \Delta ddh \) mutant evaluated in this period are 6.67 and 10.0, respectively. The \( Y_{H2G} \) in \( \Delta adh\Delta ddh \) mutant is constant at 26 from 0 h to 72 h.

No decrease in \( m_c X \) is seen from 48 to 96 h in run with GT strain, from 0 to 24 h and 48 to 72 h in run with \( \Delta ddh \) mutant and from 24 to 72 h in run with \( \Delta adh\Delta ddh \) mutant. The molecules of NAD(P)H for NiFe-hydrogenase are found to be mainly supplied by oxidative reactions other than glycolysis such as turnover of proteins, membrane lipids and PHB.

The decrease in \( m_c X \) with time is highest from 24 to 48 h in runs with GT strain and \( \Delta ddh \) mutant, and from 0 to 24 h in run with \( \Delta adh\Delta ddh \) mutant. The yields of hydrogen on endogenous glucose (\( Y_{H2G} \)) in GT strain and in \( \Delta ddh \) mutant evaluated in this period are 6.67 and 10.0, respectively. The \( Y_{H2G} \) in \( \Delta adh\Delta ddh \) mutant is constant at 26 from 0 h to 72 h.

The present results are obtained in runs with inocula from late-logarithmic growth phase. If logarithmically growing cells are inoculated into dark anaerobic nitrate-free HEPES buffer solution, the \( m_c X \) increases in the first 24 h and then decreases after 24 h (data not shown). The inoculum \( m_c X \) levels of GT strain, \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) mutant were 0.21, 0.17 and 0.14 \( \mu \text{mol mL}^{-1} \), respectively. Figure 4 (b) shows that the \( m_c X \) at 96 h of runs with GT strain, \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) mutant were 0.06, 0.07, and 0.07 \( \mu \text{mol mL}^{-1} \), respectively. The \( m_c X \) of \( \Delta adh\Delta ddh \) mutant is relatively stable with time.

Cell weight consumption characteristics in the absence of glucose: Concentration of dry cell weight of GT strain or \( \Delta adh\Delta ddh \) mutant slightly decreased with time, while that of \( \Delta ddh \) mutant was almost not varied with time (Figure 4(c)).

Lactate production characteristics in the absence of glucose: The production of lactate in GT strain started right after dark incubation Although the deletion of \( ddh \) genes is confirmed in Figure 2, the time courses of lactate concentration of runs with \( \Delta ddh \) mutant and with \( \Delta adh\Delta ddh \) mutant show the production of trace amount of lactate. Lactate concentrations of runs with \( \Delta ddh \) and \( \Delta adh\Delta ddh \) mutants at 96 h were extremely low as 0.03 and 0.01 \( \mu \text{mol mL}^{-1} \), respectively. These trace amount represent the generation of \( \Delta ddh \) phenotype deficient cells from \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) mutant during photosynthesis on antibiotic-free BG-11 medium and dark incubation on antibiotic-free HEPES buffer solution.

The specific production rate of lactate by GT strain in the first 24 h was 0.000397 \( \mu \text{mol} \mu \text{mol mg}^{-1} \text{ h}^{-1} \). The increases in lactate concentration in the incubation of \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) on glucose in the first 24 h were 0.01 \( \mu \text{mol mL}^{-1} \) and 0 \( \mu \text{mol mL}^{-1} \), respectively, hence the fraction of \( \Delta ddh \) phenotype deficient cells in \( \Delta ddh \) cells and \( \Delta adh\Delta ddh \) cells are analyzed to be 0.5 and 0, respectively. Lactate concentrations at 96 h of runs with \( \Delta ddh \) and \( \Delta adh\Delta ddh \) mutants were extremely low as 0.03 and 0.01 \( \mu \text{mol mL}^{-1} \), respectively. The initial yield of lactate on endogenous glucose of GT strain was 0.4, which shows that the relatively large portion of degraded-glycogen is utilized not for the synthesis of lactate. Although hydrogen production rate and endogenous glucose consumption rate decreased with time, lactate production rate was accelerated after 48 h. The carbon for lactate production after 48 h appears to be supplied from not only endogenous glucose but also other carboenous compounds.

Acetate production characteristics in the absence of glucose: Acetate concentration is about one-tenth of lactate concentration (Figure 4(e)). Acetate is not detected in the incubation of \( \Delta adh\Delta ddh \) mutant. The production of acetate delays for 24 h in the incubations of GT strain and \( \Delta ddh \) mutant. No difference is observed in the acetate production in GT strain and in \( \Delta ddh \) mutant from 24 h to 72 h. The acetate production rate in this period was 0.00417 \( \mu \text{mol mL}^{-1} \text{ h}^{-1} \). After 72 h, acetate production was activated in GT strain but turned to consumption in \( \Delta ddh \) mutant. The acetate consumption after 72 h in \( \Delta ddh \) mutant appears to show an oxidation of acetate to carbon dioxide in incomplete TCA cycle. In \( \Delta ddh \) mutant, acetate appears to be redirected to compensate the damaged cell constituting materials for survival. Acetate consumption in \( \Delta ddh \) mutant after 72 h is not seen in GT strain that represents high activity in endogenous glucose consumption. In GT strain, degradation of endogenous glucose likely contributes to be directed to biochemical synthesis for survival. Different from mutant, GT strain processed actively in glycolysis by converting glycogen lactate and acetate.

Ethanol production characteristics in the absence of glucose: Ethanol concentration at 96 h of runs with GT strain, \( \Delta ddh \) mutant and \( \Delta adh\Delta ddh \) mutant was extremely low as 0.7, 1.0 and 0.4 \( \mu \text{mol mL}^{-1} \), respectively. These results show that the specific rates of ethanol production by GT strain and \( \Delta ddh \) mutant were 0.00283 and 0.00404 \( \mu \text{mol} \mu \text{mol mg}^{-1} \text{ h}^{-1} \).
mg⁻¹ h⁻¹, respectively. The ethanol production of Δddh mutant was higher than that of GT strain. The excess NADH molecules that were generated by elimination of lactate hydrogenase activity appears to be redirected to alcohol dehydrogenase. Drop of ethanol concentration of run with ΔadhΔddh mutant from that with Δddh mutant partially verifies the successful deletion of the genes coding for adh. Incomplete elimination of ethanol production in run with ΔadhΔddh mutant shows the generation of Δadh phenotype deficient cells from ΔadhΔddh mutant.

3.4 Characteristics of Dark Anaerobic Nitrate-Free Hydrogen Production in the Presence of Glucose

In the second series, photoautotrophic cells of GT strain, Δddh mutant and ΔadhΔddh in late-logarithmic growth phase were harvested and incubated with X at 2.58 mg mL⁻¹ in dark anaerobic nitrate-free antibiotic-free HEPES buffer solution with 5 mmol mL⁻¹ exogenous glucose. Figure 5 shows the time courses of (a) the number of moles of hydrogen per culture volume (\(y_{H_2}\)), (b) exogenous glucose concentration (\(c_S\)), (c) the number of moles of endogenous glucose per culture volume (\(m_GX\)), (d) dry cell weight concentration (\(X\)), (e) lactate concentration (\(c_L\)), (f) acetate concentration (\(c_A\)) and (g) ethanol concentration (\(c_E\)). Different from curves in Figure 4, curves in Figure 5 represent the heterotrophic growth of GT strain, Δddh mutant and ΔadhΔddh mutant on exogenous glucose. Production of lactate, acetate and ethanol in GT strain were also become eminent.
Figure 5. Time courses of culture variables for dark anaerobic hydrogen production of cells of GT strain (○), Δddh mutant (■) and ΔadhΔddh mutant (▲) in HEPES buffer solutions supplemented with 5 μmol mL⁻¹ glucose. Quantities measured include (a) the moles of hydrogen per culture volume, (b) the concentration of exogenous glucose, (c) the moles of endogenous glucose in glycogen per culture volume, (d) the concentration of cell mass, (e) the concentration of lactate, (f) the concentration of acetate and (g) the concentration of ethanol. Experiments were carried out in duplicate.
Hydrogen production characteristics in the presence of glucose: Figure 5(a) shows that the $y_{H2}$ versus time data of runs with exogenous glucose are fitted by Eq.(1). The parameters $r_{H2,o}$, $k$ and $y_{H2,f}$ for curve-fittings are tabulated in Table1. The calculated $y_{H2}$ agrees relatively well with the observed $y_{H2}$. The $y_{H2}$ of glucose-supplemented runs with GT strain, Δddh mutant and ΔadhΔddh mutant were 5.61, 3.48 and 2.45 μmol mL$^{-1}$, respectively, which were 2.46, 1.50 and 1.65-fold, respectively, over runs without glucose. It becomes eminent that exogenous glucose elevates dark anaerobic hydrogen production in GT strain, Δdh mutant and ΔadhΔddh mutant. The $r_{H2,o}$ of Table 1 is highest for run with GT strain, lowest for run with Δdh mutant and second lowest for run with ΔadhΔddh mutant. This order is quite different from that of runs without glucose. Intracellular electrons increased at the dark anaerobic inoculation is likely directed to NiFe-hydrogenase in the absence of glucose, however, they are directed not only NiFe-hydrogenase but also many metabolic pathways including cell growth, acid fermentation that are triggered in the presence of glucose. Hence $r_{H2,o}$ of Δdh mutant under exogenous glucose solution couldn’t show highest activity. Highest $y_{H2,f}$ is found to be achieved utilizing GT strain in the presence of exogenous glucose. The attainable level of $y_{H2}$ by Δdh mutant and ΔadhΔddh is less than that of GT strain. When glucose was present, the highest $y_{H2,f}$ was GT strain, second highest is Δdh strain and the lowest is ΔadhΔddh mutant. The elimination of adh activity resulted to decrease $y_{H2,f}$ that was 0.49 times that of GT strain in runs with exogenous glucose.

Exogenous glucose consumption characteristics in the presence of glucose: Curves in Figure 5(b) show that consumption activity of exogenous glucose by GT strain is higher than that by Δdh mutant and ΔadhΔddh mutant. Glucose consumption of GT strain was active in the first 48 h, while that of Δdh mutant and that of ΔadhΔddh mutant were inactive in the first 72 h. Conversions of glucose at 96 h by GT strain, Δdh mutant and ΔadhΔddh mutant were 0.76, 0.19 and 0.23, respectively. The observed block of exogenous glucose assimilation by the disruption of ddh genes was not anticipated.

Endogenous glucose production/consumption characteristics in the presence of glucose: The time courses of the $m_cX$ show that, during 96 h incubation, the $m_cX$ of mutant increased during 24 h and become constant after 48 h whereas the $m_cX$ of GT show accumulation of endogenous glucose from the beginning to 72 h (Figure 5(b)). Although hydrogen production in GT strain was highest at initial, an increase in the $m_cX$ at initial was seen in run with GT strain. The initial rate of endogenous glucose accumulation of GT, Δdh mutant and ΔadhΔddh mutant were 0.00384, 0.00250 and 0.00032 μmol mL$^{-1}$ h$^{-1}$, respectively. The decrease in $m_cX$ with time is not observed from 48 to 96 h in runs with Δdh mutant and ΔadhΔddh mutant, they were constant. The endogenous glucose of GT strain was increasing during fermentation with a sharp drop after 72 h. The present results are obtained in runs with inocula from late-logarithmic growth phase.

Cell weight production/degradation characteristics in the presence of glucose: Dry cell weight concentration of GT strain increased in the first 24 h, kept constant between 24 and 48 h, and decreased quite slightly after 48 h. The initial increase in dry cell weight concentration is associated with the reduction in exogenous glucose concentration. This association confirms the heterotrophic growth of GT strain on glucose. The rate of decrease in dry cell weight concentration after 48 h is comparable with the initial decrease in dry cell weight concentration of run without glucose. The time course of dry cell weight concentration of ΔadhΔddh mutant is comparable with that of Δdh mutant. Dry cell weight concentrations of Δdh mutant and ΔadhΔddh mutant increased in the first 24 h, and decreased after 24 h.

Interestingly, when cells of Δdh mutant and ΔadhΔddh mutant grew in the first 24 h, concentrations of exogenous glucose were not varied. No decrease in the number of moles of endogenous glucose per culture volume in the first 24 h of runs with Δdh mutant and ΔadhΔddh mutant is seen in Figure 5(c). Growth of Δdh mutant and ΔadhΔddh mutant appears to be triggered by exogenous glucose. Combination of the time courses of exogenous glucose, endogenous glucose and dry cell weight suggests that carbon source for cell growth of Δdh mutant is not exogenous glucose or endogenous glucose but unidentified intracellular organic compounds.

Lactate production characteristics in the presence of glucose: Lactate concentration at 96 h of run with GT strain in the presence of glucose was 4.74 μmol mL$^{-1}$ which is 18 times more than that in the absence of glucose. This result shows that exogenous glucose is converted to lactate by GT strain. The yield of lactate on dry cell weight of GT strain was 5.61 μmol mg$^{-1}$ which is 18 times more than that of GT strain. The yield of lactate on dry cell weight of Δdh mutant and ΔadhΔddh mutant was 3.48 and 2.45 μmol mL$^{-1}$, respectively, hence the fraction of Δdh phenotype deficient cells in Δdh mutant and ΔadhΔddh mutant are analyzed to be 0.00362 and 0.00844, respectively. Lactate concentrations at 96 h of runs with with Δdh and ΔadhΔddh mutants were extremely low as 0.05 and 0.06 μmol mL$^{-1}$, respectively. The deletion of ddh genes in this work possibly altered the intracellular balance between NAD(P)H and NAD(P)$^+$ towards shifting-up NAD(P)H level. This NAD(P)H shifting-up is important because NADH and NADPH are essential reductive compounds for hydrogen production on NiFe-hydrogenase. The increase in the level of NAD(P)H results in the increase in the level of ATP and
and subsequently inactivates phosphofructokinase. Assimilation of exogenous glucose by Δddh mutant appears to be blocked by the inactivation of phosphofructokinase, brought by the elimination of the genes coding for ddh.

Acetate production characteristics in the presence of glucose: Acetate production is highly activated by exogenous glucose in GT strain. Exogenous glucose stimulated acetate production also in Δddh mutant and ΔadhdΔddh mutant. After 96 h, acetate concentrations of runs with GT strain, Δddh mutant and ΔadhdΔddh mutant in the presence of exogenous glucose were 17.19, 8.66 and 4.83 μmol mL⁻¹, respectively, that were about more than 100 times those in the absence of exogenous glucose. Acetate is one of the terminal carbon release compound from glucose consumption. Glucose consumption and acetate production of GT strain were higher than those of Δddh and ΔadhdΔddh mutants.

Ethanol production characteristics in the presence of glucose: Ethanol concentration at 96 h of runs with GT strain, Δddh mutant and ΔadhdΔddh mutant was extremely low as 0.70, 1.01 and 0.47 μmol mL⁻¹, respectively. Extremely low production of ethanol by the incubation of Δddh mutant from antibiotic-free photautotrophic culture verifies the successful deletion of the genes coding for adh. The yield of ethanol on dry cell weight of GT strain in the first 24 h is 0.45 μmol mg⁻¹. The increase in ethanol concentration in the incubation of ΔadhdΔddh mutant on glucose in the first 24 h was 0.17 μmol mL⁻¹, hence the fraction of Δadh phenotype deficient cells in ΔadhdΔddh cells are analyzed to be 0.180.

If stability of hydrogen production is evaluated in low value of the parameter k shown in Table 1, incubation of Δddh mutant in the presence of exogenous glucose represents most stable hydrogen production condition. This stabilization is not harnessed in the attainable level of hydrogen (γH₂,0), which is highest in run with GT strain in the presence of exogenous glucose. When viewed over the parameters in this study, the increase in the initial hydrogen production rate (γH₂,0) is found to be crucial to elevate the level of γH₂,f. Elimination of ddh and adh genes is effective to increase the level of γH₂,0 in the absence of exogenous glucose, however, it becomes ineffective in the presence of exogenous glucose. Redox balance in the presence of exogenous glucose is found to be different from that in the absence of exogenous glucose.

4. Discussion

Our results show that exogenous glucose shifts up initial rate of hydrogen production in GT strain, Δddh mutant and ΔadhdΔddh mutant, and that, in the absence of exogenous glucose, initial hydrogen production rates of Δddh mutant and ΔadhdΔddh mutant are higher than that of GT strain, but in the presence of glucose, that of Δddh mutant and ΔadhdΔddh mutant are lower than that of GT strain. The highest attainable level of the number of moles of hydrogen per culture volume of 5.61 μmol mL⁻¹ is achieved with GT strain in the presence of glucose due to heterotrophic growth of GT strain by assimilating exogenous glucose and producing endogenous glucose, lactate and acetate. The lowest attainable level of hydrogen at 2.28-2.45 μmol mL⁻¹ is achieved with GT strain or Δddh mutant in the absence of glucose and with ΔadhdΔddh mutant in the presence of glucose.

In cyanobacterial research, conversion of glucose 6-P to pyruvate is assumed to occur through OPP pathway. Complete decomposition of 1 μmol endogenous glucose of reserved glycogen to 5/3 μmol pyruvate and 1 μmol carbon dioxide through OPP pathway produces 11/3 μmol NAD(P)H. The stoichiometry of ddh and adh represent that conversion of 1 μmol pyruvate to 1 μmol lactate consumes 1 μmol NADH and that conversion of 1 μmol pyruvate to 1 μmol ethanol via acetaldehyde also consumes 1 μmol NADPH.

Conversion of 1 μmol pyruvate to 1 μmol acetyl-CoA and 1 μmol carbon dioxide on pyruvate oxidoreductase (nif J) generates 1 μmol reduced ferredoxin from oxidized ferredoxin. Conversion of acetyl-CoA to acetate via acetyl-P on phosphate acetyltransferase (pta) and acetate kinase (ack A) (Juntarajumnong et al., 2007) in the acetate-forming direction requires 1 μmol ADP and 1 μmol Pi for oxidation. Complete oxidation of 1 μmol pyruvate to 3 μmol carbon dioxide through incomplete TCA cycle via acetyl-CoA produces 36/5 μmol NADPH.

Deletion of NAD(P)H-consuming dehydrogenase pathways redirects more NAD(P)H to NiFe-hydrogenase and affects to increase hydrogen production rate. Successful mutant constructions are confirmed by a drastical drop in lactate production and ethanol production by corresponding mutants. Although extremely low amounts of lactate and ethanol were detected in run with Δddh mutant and ΔadhdΔddh mutant, these are regarded as the appearance of wild type strain since antibiotics were not utilized for main culture and dark anaerobic incubation of those mutants. Production rates of lactate and ethanol by mutants in Figures 4 and 5 are 0 μmol mL⁻¹ h⁻¹, even though trace amounts of lactate and ethanol were detected for antibiotic-free conditions. During the first 24 h, acetate productions in GT strain, Δddh mutant and ΔadhdΔddh mutant are negligible small. Global metabolism of this strain accumulates NAD(P)H that is generated by glycolysis through OPP pathway, oxidation of pyruvate through incomplete TCA cycle and turnover of proteins, membrane lipids and PHB, and consumes it on NiFe-hydrogenase,
ddh, adh and other oxidative reactions to keep redox homeostasis. At the beginning of dark incubation, the inoculum cells of Δddh mutant and ΔadhΔddh mutant from photosynthesis contain high amount of NAD(P)H. The incubation of cells in nitrate-free solution in the dark triggers turnover of glycogen, proteins, membrane lipids and PHB. Turnover reactions are accompanied by accumulation of NAD(P)H. The excess NAD(P)H is utilized for hydrogen production, lactate production and ethanol production via acetate production. In GT strain, the summation of initial consumption rates of NAD(P)H by NiFe-hydrogenase and lactate dehydrogenase is 0.0317 μmol mL⁻¹ h⁻¹ that is higher than the theoretical initial production rate of NAD(P)H at 0.0250 μmol mL⁻¹ h⁻¹ from OPP pathway and incomplete TCA cycle, hence assumption is raised that, NAD(P)H for NiFe-hydrogenase is found to be supplied not only by glycolysis but also by turnover of proteins, membrane lipids and PHB.

In Δddh mutant under exogenous glucose-free condition, even though the activity of ddh is absent, the theoretical initial production rate of NAD(P)H from OPP pathway and incomplete TCA cycle at 0.0050 μmol mL⁻¹ h⁻¹ is only 14% of NAD(P)H consumed by NiFe-hydrogenase. At the beginning of dark incubation, the major NAD(P)H generating reaction for hydrogen production in Δddh mutant is found to be turnover reactions other than glycolysis. Under exogenous glucose condition, in the first 24 h, the carbon flux at glucose 6-P is directed to glycogen synthesis, cell growth, lactate production, acetate production and ethanol.

In ΔadhΔddh mutant under exogenous glucose-free condition, summation of theoretical production rates of NAD(P)H by OPP pathway and incomplete TCA cycle is 0.0150 μmol mL⁻¹ h⁻¹ that is 42% of NAD(P)H consumed by NiFe-hydrogenase. About a half of NAD(P)H is found to be supplied by turnover of proteins, membrane lipids and PHB.

Although hydrogen production rate is improved, other competitive pathway such as nitrate reductase may play a major role in electron sink. A previous work reported a significant improvement in hydrogen production up to more than 20-folds on nitrate-deprived BG-11 medium (Baebprasert et al. 2011).

Our experimental results present a knowledge that the deletion of ddh genes and/or adh genes is useful to increase reductive potential of Synechocystis sp. strain PCC6803-GT against cellular property of redox homeostasis. Exposure of cells to dark anaerobic nitrate-free environment triggers the adaptation of cells to new redox homeostasis state, hence above reductive potential is unstable with time.

The ΔadhΔddh mutant constructed in this study is characterized by a 37% decrease in the initial amount of endogenous glucose per culture volume, a 15% increase in the initial hydrogen production rate, 45% decrease in the attainable level of hydrogen, stable level in dry cell weight concentration, stable level in endogenous glucose, almost no production of lactate and no production of acetate. An addition of reducing sugar to HEPES buffer solution containing ΔadhΔddh mutant appears to elevate the amount of endogenous glucose per culture volume, which is promising to elevate hydrogen production.

5. Conclusions

This study provides additional information as to the elevation of dark anaerobic hydrogen production in a glucose tolerant mutant of Synechocystis sp. strain PCC6803 by deleting the genes coding for lactate dehydrogenase and alcohol dehydrogenase that compete NAD(P)H with hydrogen production of NiFe-hydrogenase. Although mutation cannot affect the attainable level in number of moles of hydrogen per culture volume, it results a 15% increase in the initial hydrogen production rate. This finding have led to a mechanism that the initial turnover of proteins, membrane lipids and PHB is related to the mutation. Glycogen turnover was reduced in mutants. Glucose is a good substrate for hydrogen production improvement. However, lower affinity of glucose consumption by mutant cells inhibits mutants to turn glucose for glycogen accumulation. Although the homeostasis of Synechocystis sp. strain PCC6803 in terms of alternations of its metabolic functions limit metabolic engineering approaches to increase hydrogen production, mutants certainly have a great potential for responding to the shift in environments. More data will be needed before the combined effects of mutation and some environmental supplement become apparent.

References


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