Temperature and Carbon Assimilation Regulate the Chlorosome Biogenesis in Green Sulfur Bacteria

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ABSTRACT Green photosynthetic bacteria adjust the structure and functionality of the chlorosome—the light-absorbing antenna complex—in response to environmental stress factors. The chlorosome is a natural self-assembled aggregate of bacteriochlorophyll (BChl) molecules. In this study, we report the regulation of the biogenesis of the Chlorobaculum tepidum chlorosome by carbon assimilation in conjunction with temperature changes. Our studies indicate that the carbon source and thermal stress culture of C. tepidum grows slower and incorporates fewer BChl c in the chlorosome. Compared with the chlorosome from other cultural conditions we investigated, the chlorosome from the carbon source and thermal stress culture displays (a) smaller cross-sectional radius and overall size, (b) simplified BChl c homologs with smaller side chains, (c) blue-shifted Q y absorption maxima, and (d) a sigmoid-shaped circular dichroism spectra. Using a theoretical model, we analyze how the observed spectral modifications can be associated with structural changes of BChl aggregates inside the chlorosome. Our report suggests a mechanism of metabolic regulation for chlorosome biogenesis.

INTRODUCTION

Green photosynthetic bacteria, which include green sulfur bacteria and green nonsulfur bacteria, can live in low-light or extremely low-light environments (1–3). To efficiently harvest light in such conditions, green photosynthetic bacteria use the chlorosome, a natural optical antenna complex. The solar energy absorbed by the chlorosome is converted to molecular excitations and then transferred to the reaction center, where a transmembrane electron potential is generated. The chlorosome contains hundreds of thousands of bacteriochlorophylls (BChls) (4–6) that are self-assembled into supramolecular aggregates with few proteins associated. Several reports have implied the great potential of the chlorosome for its use in biohybrid and biomimetic devices (7–9).

Biogenesis of the chlorosome from green sulfur bacteria has been reported to be regulated by multiple environmental factors including light intensities (10), temperature (11,12), light intensities combined with temperatures (13), and sulfate concentrations (13,14). In this work, we study how the chlorosome in the green sulfur bacterium Chlorobaculum tepidum is adapted to the type of carbon source and thermal stress conditions. Carbon assimilation and electron transport are known to be a bottleneck of photosynthesis (15). To store solar energy, green sulfur bacteria use the reductive (reverse) tricarboxylic acid (TCA) cycle for carbon fixation (2,16,17). In contrast to the oxidative (forward) TCA cycle, which is energetically favorable, the reductive TCA cycle uses ATP and NAD(P)H produced via photosynthetic electron transport for assimilating inorganic carbon and synthesizing acetyl-CoA. Green sulfur bacteria are known to grow mixotrophically with acetate, pyruvate, and a few organic carbon sources (2). Studies indicate that mixotrophic-grown green sulfur bacteria operate mainly the reductive TCA cycle for assimilating pyruvate, because of low carbon flux from pyruvate to acetyl-CoA, and operate both oxidative and reductive TCA cycles for assimilating acetate (18) (Fig. S1 in Supporting Material). Thus, green sulfur bacteria exhibit better growth with acetate, whereas they require more energy to assimilate pyruvate.

In this study, we analyzed the structural and morphological modifications of the C. tepidum chlorosome upon changes in temperatures and nutrition; viz. organic carbon sources, using an array of techniques such as small-angle neutron scattering (SANS), ultraviolet (UV)-visible absorption, circular dichroism (CD), 77 K fluorescence emission spectra, and mass spectra. To obtain more structural insights on BChl packing in the chlorosome from the different cultures, we computed absorption and CD spectra of BChl aggregates composing the chlorosome.

MATERIALS AND METHODS

Materials

All solvents and reagents were obtained from standard commercial sources and used as received. The thermophilic green sulfur bacterium C. tepidum was cultured anaerobically and phototrophically using the medium as reported (2,18). The cultures were grown in low-intensity light (20 ± 0.2 μmol/m²/s) inoculated with 1% cell cultures grown at 50°C or 30°C in the late exponential growth phase. Cell cultures were monitored using optical density (OD) of a cell suspension at 625 nm, where the absorbance of photosynthetic pigments is minimal (18). The cell growth rate was

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The scattering data of the chlorosome were also analyzed with a biaxial ellipsoid of rotation fit (22). A scale factor, which is proportional to the square of (the particle volume) term and particle concentration (i.e., number of such particles), was included in the data fit.

RESULTS
Grow C. tepidum at various temperatures and with different carbon sources

We first investigated C. tepidum grown in 12 different cultural conditions with varied temperatures (constant temperature, temperature up-shifted, and temperature down-shifted) and carbon sources (HCO$_3^-$, HCO$_3^-$/acetate, and HCO$_3^-$/pyruvate) (Table S1). The culture growth rates, Q$_g$ absorption maxima, room temperature fluorescence emission maxima, and hydrodynamic diameters of the chlorosome are summarized in Table S2. The results indicate that the growth rate for cultures supplied with various carbon sources follows the trend HCO$_3^-$/acetate cultures > HCO$_3^-$ cultures > HCO$_3^-$/pyruvate cultures. Moreover, the overall size of the chlorosome from HCO$_3^-$/pyruvate cultures is smaller than the chlorosome from HCO$_3^-$/acetate cultures. Culture 6 (the temperature down-shifted culture grown on pyruvate/HCO$_3^-$), which in our understanding experiences maximal environmental stress, in addition to slowest growth rate and smallest overall size, exhibited a substantial blue-shift of the Q$_g$ absorption maximum and the room temperature fluorescence peak. In the rest of the article, we compare the structural, optical, and functional properties of the chlorosome from Culture 6 with chlorosomes from other cultures and also with chlorosomes from Chloroflexus aurantiacus.

Structural information

To acquire more structural insights on the chlorosome from Culture 6 and other cultures grown on various carbon sources and temperatures, we employed LC/MS to analyze the BChl c homologs in the chlorosome and SANS to investigate the morphology of the chlorosome.

**LC/MS measurements.** We analyzed the ratio of BChl c homologs in the chlorosome via LC/MS. Fig. 1 shows the data from 6 of 12 cultures. Fig. 1 A shows the high-performance liquid chromatography chromatogram for BChl c homologs in the chlorosome with detection wavelength at 640 nm, and Fig. 1 B reports the percentage of BChl c homologs in the chlorosome analyzed by mass peak area of LC/MS. As reported in the literature (12,23,24), major BChl c homologs in the C. tepidum chlorosome are [8-ethyl, 12-methyl]-farnesyl-BChl c (m/z 793.5), [8-ethyl, 12-ethyl]-farnesyl-BChl c (m/z 807.5), [8-propyl, 12-ethyl]-farnesyl-BChl c (m/z 823.5), [8-isobutyl, 12-ethyl]-farnesyl-BChl c (m/z 839.5), and [8-neopentyl, 12-ethyl]-farnesyl-BChl c (m/z 853.5). Fig. 1 A shows the coelution of [8-ethyl, 12-ethyl]-farnesyl-BChl c/[8-propyl, 12-ethyl]-farnesyl-BChl c (Rt 5.35 min), [8-propyl, 12-ethyl]-farnesyl-BChl
c/[8-isobutyl, 12-ethyl]-farnesyl-BChl c (Rt 5.77 min), and [8-isobutyl, 12-ethyl]-farnesyl-BChl c/[8-neopentyl, 12-ethyl]-farnesyl-BChl c (Rt 6.30 min). Also, Fig. 1A indicates that Culture 6 exhibits simpler BChl c homolog distributions with very small peak in the retention time at 5.8 min (corresponding to BChl c homologs with larger substituents at C-8) (12), and Fig. 1B shows that more than 70% of BChl c homologs in the chlorosome from Culture 6 are [8-ethyl, 12-ethyl]-farnesyl-BChl c (m/z 807.5), in contrast to more heterogeneous distribution of BChl c homologs in other cultures. Culture 6 contains simpler BChl c homologs, which is akin to the distribution of BChl c homologs in C. aurantiacus, which only has [8-ethyl, 12-methyl]-BChl c because of the absence of genes encoding C-8², C-12¹ methyltransferases (25).

SANS measurements. We employed SANS for probing the morphology (i.e., size and shape) of the chlorosome. Our previous studies highlighted the power of SANS in elucidating structural features critical to the function of chlorosomes (7,20,21,26). Fig. 2A shows scattering data of the chlorosome from Cultures 1, 3, 4, 6, 7, and 9, with the data merged from three experimental configurations covering the q-range 0.001–0.70 Å⁻¹. All chlorosome samples for SANS measurements were prepared in 100% D₂O buffer (20 mM Tris-HCl at pH 7.8). Fig. 2B shows that chlorosomes from all cultures exhibit a rod-like shape, indicating that the morphology of the chlorosome from Culture 6 is similar to the chlorosome from other cultures. Table 1 lists the values of the parameters of modified Guinier analysis for a rod-like particle (Eq. 1) (7,20) performed on the

![FIGURE 1](A) High-performance liquid chromatography chromatogram of BChl c homologs in the chlorosome from Cultures 1, 3, 4, 6, 7, and 9 with detection wavelength at 640 nm and the elution pattern of each BChl c homolog. (B) Percentage of BChl c homologs in the chlorosome determined by mass peak area.

![FIGURE 2](A) SANS profiles of the chlorosome isolated from constant temperature cultures at 50°C (Culture 1 and Culture 3), temperature down-shifted to 30°C cultures (Culture 4 and Culture 6), and constant temperature cultures at 30°C (Culture 7 and Culture 9). SANS data fitted with modified (B) Guinier’s model and (C) biaxial ellipsoid of rotation.

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Biophysical Journal 105(6) 1346–1356

Kuo-Hsiang Tang et al.
chlorosome from all cultures. The cross-sectional radius of gyration of the chlorosome from Culture 6 is significantly smaller compared with other cultures. The results follow the trend observed in the dynamic light-scattering measurements (Table S2).

Fig. 2 C shows the scattering data analyzed with a biaxial ellipsoid of rotation fit (22). The cross-sectional radius of the chlorosome analyzed with modified Guinier model (R-value in Table 1) agrees with the biaxial ellipsoid of rotation fit (perpendicular semiaxis value in Table 2), further indicating that the chlorosome from Culture 6 appears to have smaller cross-sectional radius (i.e., more flat) compared with the chlorosome from other cultures.

Taken together, the LC/MS and SANS data show that the chlorosome from Culture 6 exhibits simpler BChl c homologs and smaller cross-sectional radius compared with the chlorosome from other cultures.

**Optical properties**

**Absorbance and CD measurements.** The structural studies described above indicate a different morphology of the chlorosome from Culture 6 compared with the chlorosome from other cultures. To understand further the changes in the internal structure of the chlorosome from the stressed culture, we compared the chlorosome from two temperature down-shifted cultures: Culture 4 (grown on HCO$_3^-$/acetate) and Culture 6 (grown on HCO$_3^-$/pyruvate). Fig. 3 A shows the UV-visible spectra of Culture 4 and Culture 6, in which Culture 6 displays a blue-shifted $Q_y$ maximum and smaller cross-sectional radius (i.e., more flat) indicating that the chlorosome from Culture 6 appears to have smaller cross-sectional radius compared with the chlorosome from other cultures.

**Fluorescence emission spectra.** To characterize the stress-associated changes in the excitation energy transfer, we compared low-temperature (T = 77 K) fluorescence emission spectra of the whole cell and the chlorosome from Culture 4 and Culture 6. For the chlorosome spectra, the rate of excitation transfer from BChl c in the chlorosome (780–784 nm) to BChl a in the baseplate complex can be estimated with the ratio of the peak area (the baseplate complex/chlorosome). In the whole-cell spectra, this ratio (the baseplate complex or the Fenna–Matthews–Olson protein/chlorosome) can also be affected by energy transfer from the baseplate to the Fenna–Matthews–Olson protein and reaction center.

Green sulfur bacteria are known to exhibit redox-dependent energy transfer (1,2). Thus, cultures and chlorosomes were incubated in 100-mM freshly prepared dithiothreitol in the anaerobic chamber overnight as described previously (12). Fig. 5 A shows a stronger relative emission from the baseplate energy domain of the stressed chlorosome, with peak ratio 25% more compared with the chlorosome from Culture 4 (Fig. S2). This can be an indication of an enhanced chlorosome peak intensity. The chlorosome from Culture 6 displays smaller intensity of the $Q_y$ absorption band (Fig. 3 B), implying that fewer BChls are contained in the chlorosome. In addition, we observed that the $Q_y$ maxima of the Culture 6 chlorosome and the *C. aurantiacus* chlorosome are almost identical (Fig. 3 C), with the Culture 6 chlorosome exhibiting a broader peak. The latter fact is likely due to various BChl c homologs present in the Culture 6 chlorosome (27).

Table 2 Scattering parameters of the chlorosome determined with modified Guinier analysis.

<table>
<thead>
<tr>
<th>Chlorosome</th>
<th>$I(0)$</th>
<th>$R_c$ ($\AA$)*</th>
<th>$R$ ($\AA$)†</th>
<th>$R_c \times Q_{\text{max}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.22 ± 0.04</td>
<td>126 ± 4</td>
<td>178 ± 6</td>
<td>0.92 ± 0.05</td>
</tr>
<tr>
<td>3</td>
<td>0.38 ± 0.03</td>
<td>123 ± 3</td>
<td>174 ± 4</td>
<td>0.91 ± 0.04</td>
</tr>
<tr>
<td>4</td>
<td>0.78 ± 0.03</td>
<td>127 ± 3</td>
<td>180 ± 4</td>
<td>0.93 ± 0.04</td>
</tr>
<tr>
<td>6</td>
<td>0.14 ± 0.02</td>
<td>92 ± 4</td>
<td>130 ± 6</td>
<td>0.92 ± 0.03</td>
</tr>
<tr>
<td>7</td>
<td>0.15 ± 0.03</td>
<td>122 ± 3</td>
<td>173 ± 4</td>
<td>0.90 ± 0.03</td>
</tr>
<tr>
<td>9</td>
<td>0.51 ± 0.04</td>
<td>126 ± 4</td>
<td>178 ± 6</td>
<td>0.92 ± 0.04</td>
</tr>
</tbody>
</table>

* $R_c$ is the cross-sectional radius of gyration for a particle.
† $R$ is the radius of a rod-like particle of infinite length defined as $R = \sqrt{2} \times R_c$. 

**Table 2** Scattering parameters of the chlorosome determined with biaxial ellipsoid of rotation fit, estimated volume of the chlorosome, and estimated number of BChl c in the chlorosome.

<table>
<thead>
<tr>
<th>Chlorosome</th>
<th>Perpendicular semiaxis</th>
<th>Parallel semiaxis</th>
<th>Volume ($\times 10^3 \text{ nm}^3$)</th>
<th>Number of BChl c ($\times 10^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>19.2 ± 0.4</td>
<td>221.1 ± 41.2</td>
<td>344.2 ± 77.9</td>
<td>165 ± 40</td>
</tr>
<tr>
<td>3</td>
<td>18.8 ± 0.3</td>
<td>184.1 ± 32.4</td>
<td>274.2 ± 56.7</td>
<td>129 ± 27</td>
</tr>
<tr>
<td>4</td>
<td>18.9 ± 0.5</td>
<td>241.4 ± 24.5</td>
<td>363.4 ± 55.8</td>
<td>173 ± 28</td>
</tr>
<tr>
<td>6</td>
<td>15.4 ± 0.5</td>
<td>162.8 ± 30.0</td>
<td>163.8 ± 40.3</td>
<td>77 ± 21</td>
</tr>
<tr>
<td>7</td>
<td>18.9 ± 0.6</td>
<td>182.4 ± 25.0</td>
<td>275.6 ± 54.8</td>
<td>132 ± 27</td>
</tr>
<tr>
<td>9</td>
<td>18.7 ± 0.4</td>
<td>171.0 ± 22.0</td>
<td>252.0 ± 43.0</td>
<td>119 ± 20</td>
</tr>
</tbody>
</table>
chlorosome-to-baseplate energy transfer in Culture 6, despite the blue-shift of the chlorosome absorption peak. The emission spectra from the whole cells show that the exciton population of the baseplate in stressed culture is smaller (Fig. 5 B), with peak ratio 50% less compared with the chlorosome from Culture 4 (Fig. S3). We hypothesize that the temperature/nutrition stress induces structural changes in the light-harvesting complex beyond the chlorosome, which may enhance the energy funneling from the baseplate. However, the detailed analysis of this process goes beyond the scope of the current study.

Model

BChl aggregates in the chlorosome are organized as a mixture of tubular and lamellar shapes (36). Recent cryoelectron microscopy studies suggest that inside the chlorosome tubular aggregates (rolls) are composed in larger concentric assemblies (35,37). Fig. 6 illustrates our schematic understanding of aggregate packing in an idealized chlorosome, where $L_{1-3}$ are three semiaxes of the ellipsoidal body and $R$ is a radius of a tubular aggregate. Therefore, the chlorosome length $L_1$ determines the maximal length of rolls in the structure, $L_3$ limits the maximal radius of a roll, and the ratio $L_2/L_3$ controls the contribution of lamella versus roll shapes.

Among several existing models for BChl aggregation in chlorosomes (35,38–41), we used the one introduced by Ganapathy and coworkers (35,41). In addition to conventional NMR and cryoelectron microscopy studies, this structure has also been supported using two-dimensional polarization fluorescence microscopy (42). We have shown recently that the timescales characterizing exciton dynamics in this structures agree with the ones obtained in time-resolved measurements of the chlorosome (43,44). In this model, the BChl stacks form a two-dimensional lattice shown in Fig. 7 A. Subsequently, the lattice is folded into a roll. We considered two types of folding: Structure I, BChl stacks form concentric rings (Fig. 7 B); and Structure II, BChl stacks are parallel to the symmetry axis of the roll (Fig. 7 C). The former structure was predicted for a mutant chlorosome that consists of BChl $d$ pigments (35), and the same group of researchers suggested that the latter structure describes a wild-type chlorosome (41). The latter structure also qualitatively agrees with the model proposed previously (38) and used for the description of CD spectra of chlorosomes (45). For the purpose of simplicity in the following discussion, we neglect the distance variation
between syn-anti dimers, which can introduce a shift of the absorption band but cannot result in the change of the type of the CD spectra.

From the biaxial ellipsoid of rotation fit for SANS measurements and the chlorosome model shown in Figs. 6 and 7, we estimate the volume of the chlorosome and the amount of BChl c in different cultures and report the results in Table 2. Herein, we assume that the spacing between BChl aggregates is equal to 2.1 nm (35) and that approximately a half of the chlorosome volume is occupied by BChls (1). The number of BChl c homologs in the chlorosome has also been estimated by thin section and freeze-fracture electron microscopy (2,400–12,000 BChls in the C. aurantiacus chlorosome) (4), fluorescence correlation spectroscopy (215,000 ± 80,000 BChls in the C. tepidum chlorosome) (5), confocal fluorescence microscopy (∼140,000 BChls in the C. tepidum chlorosome and ∼96,000 BChls in the C. aurantiacus chlorosome) (6), and atomic force microscopy (34,000–240,000 BChls in the chlorosome of various green photosynthetic bacteria) (46). Although our estimations are qualitative, the values we obtain are comparable to other reported results. The number of BChls in the stressed Culture 6 is approximately a half of the number in other cultures (Table 2).

The shift of the Qy absorption maximum of the chlorosome can originate from the difference in electronic excitation spectra of single BChl c homologs and also modifications of BChl assemblies on different length scales. In contrast, single-molecule contribution to CD spectra of the chlorosome is very small and most of the signal is due to the supramolecular assembly.

Absorption and CD spectra of molecular aggregates

We used single BChl rolls as a representative model for the absorption and CD spectra of the chlorosome, assuming that the absorption signal is dominated by the rolls with the largest number of molecules. The latter assumption is relaxed for CD spectra because the signal is stronger for rolls with a larger curvature. Thus, we limited the maximal size of representative rolls to be approximately a half of the size of a chlorosome obtained from the SANS measurements (Table 1).

To predict possible changes in the BChl aggregation in the chlorosome from the stressed culture, we varied the

FIGURE 5 77-K fluorescence spectra of (A) the chlorosome from Culture 4 and Culture 6 (A) and (B) the whole cells from the same cultures. Spectra were normalized at the maxima of the chlorosome.

FIGURE 6 Schematic illustration of BChl packing in the chlorosome. The chlorosome is considered as an ellipsoidal body with semi-axes $L_1$, $L_2$, $L_3$. Two-dimensional aggregates of BChls are packed in a form of concentric rolls and curved lamellae.

FIGURE 7 Model of BChl aggregation. (A) BChl pigments form two-dimensional lattice (35) with the lattice parameters $a = 6.25$ Å, $b = 9.79$ Å, $\theta = 122^\circ$, and $\alpha = 35^\circ$. Binding of BChls is schematically illustrated. Arrows show the orientation of molecular Qy transition dipoles. (B) and (C) represent two suggested ways for aggregate folding in rolls. (B) Structure I suggested for a mutant chlorosome with BChl stacks forming concentric rings (41). (C) Structure II proposed for a wild-type of bacteria (41), in which BChl stacks are parallel to the roll symmetry axis of the roll.
structural parameters of the aggregates: the lattice constants \(a\) and \(b\), the angle \(\theta\) between the lattice vectors, and the orientation of the transition dipole. We also modified the shape of the aggregates by changing the length and the radius of the rolls, and also by transforming rolls to lamellae.

The absorption and CD spectra of the aggregates were computed using conventional relations (47), in which the aggregate Hamiltonian was constructed within the point dipole approximation. For the value of the monomer transition dipole, we used \(|\mu|^2 = 30\) Debye (45). The calculated stick spectra were broadened using a Lorentz line with a half width of 100 cm\(^{-1}\).

We find that for both Structures I and II, variations of the lattice constants within 3–5% or reorientations of the monomer transition dipoles within 3–5 degrees of the original values can result in a shift of the absorption spectra of the order of 100 cm\(^{-1}\), which is comparable to the measured value. However, to change the type of the CD spectra, the structural modifications should be more drastic. This result remains valid for all rolls we computed with the radius in the range \(R = 3–11\) nm and the length in the range \(L = 10–130\) nm.

The CD spectra of rolls with a large aspect ratio \(L/R\) are of the mixed type (–/+/–) for both Structures I and II. Reducing the aspect ratio for the Structure II, we observe a transition between mixed-type and Type II (–/+ ) CD spectra, which agrees with the previous study (45). Fig. 8 shows the scaling of the transition length with the radius of a roll. Two types of the spectra for rolls of the same radius and different lengths are shown in Fig. 8 B. For Structure I, we obtain a transition to Type I (+/–) for the roll length \(-10\) nm independent of its radius (Fig. 9). The latter type of CD spectra was reported for single mutant chlorosomes (48). It should be noticed that for some aspect ratios, the rolls with the same values of \(L\) and \(R\) but different structures (I or II) give different type of CD spectra. For example, a roll with \(R = 9\) nm, \(L = 80\) nm, and Structure II has a Type II CD spectrum, whereas the roll of the same shape but Structure I has a mixed type of the spectrum. Thus, different types of CD spectra can also be explained by different folding of BChl aggregates.

Transitions between different types of CD spectra are also observed if we allow for the formation of curved lamella structures. In Fig. 10, we compare the CD spectrum of a roll and a curved lamella. The radius of the roll is \(R = 7\) nm, and the length is \(L = 100\) nm. The lamella is represented by a half of the roll with the same parameters cut along the symmetry axis. The spectra of both structures are calculated using the same formula (Eq. 13 in the report by Somsen et al. (47)) and normalized to the number of molecules. For both types of BChl aggregation, we obtained a transition between the mixed-type and Type II spectra. Within our model (Fig. 6), the contribution of lamellae and roll structures in CD spectra of the chlorosome should be controlled by the \(L_2/L_1\) ratio. Although our SANS measurements do not provide information on how this ratio varies in the chlorosome from different cultures, changes in the cross-sectional radius of the chlorosome support that possibility.

To summarize the simulation results, we conclude that the shift of the \(Q_y\) absorption maximum can be assigned to small variations in BChl aggregation. The changes in CD spectra should be associated with the modifications of the shape of the aggregate. In addition to a variation of the roll length proposed previously (45) as a possible cause for the change of CD type, we suggest that the transition can be due to formation of lamellae or different folding of aggregates (Structure I vs. Structure II). We notice that the structural modifications resulting in the change of the CD type cannot describe the blue-shift of the absorption peak.

**Electronic excitation spectra of single BChl homologs**

In addition to the structural analysis for the spectra described previously, we verified computationally that the changes in the frequency of the \(Q_y\) electronic excitation of single BChls associated with the methylation at C-8 and C-12 positions cannot account for the blue-shift of the absorption peak observed in the experiments. The calculations were done using time-dependent density functional theory as implemented in the Turbomole quantum chemistry package (49). We used the PBE0 (50) hybrid functional with the triple-\(\zeta\) basis sets def2-TZVP (51). To simplify the

**FIGURE 8** Structure II: Length/radius dependence of CD spectra. (A) The characteristic length of a roll that corresponds to the transition between the mixed (–/+–) and Type II (–/+ ) CD spectra is shown as a function of a roll radius. (B) Two types of CD spectra are shown for rolls of the radius \(R = 7\) nm and the lengths \(L = 100\) nm and \(L = 60\) nm marked by black and grey stars in (A). The relative intensities are normalized to the number of molecules. The wavenumber scale shows the frequency red shift from the monomer transition due to BChl aggregation.
calculations, we substituted the farnesyl chain with a methyl group in all homologs.

The computed frequencies of BChl c Qy transitions, shown in Fig. 11, are blue shifted by ~2500 cm⁻¹ (~15% of the excitation frequency) compared with the spectra of BChl c measured in methanol. This difference can be associated with the solvent effect and also to the systematic error of density functional theory calculations (52). The computed electron excitation frequency varies by ~100 cm⁻¹ for different homologs. Weighted by the homolog composition obtained from mass spectral analyses (Fig. 1B), these can account only for ~10% (10 cm⁻¹) of the blue-shift of the Culture 6 absorption spectrum as compared with that of Culture 4.

**DISCUSSION**

Photosynthetic organisms use the energy and reducing equivalents generated via photosynthetic electron transport to produce building blocks of cells (i.e., via carbon metabolism) and undergo other cellular metabolic processes (e.g., nitrogen metabolism). It is known that carbon assimilation is a bottleneck of photosynthesis, and that enhancement of carbon assimilation is essential to the improvement of photosynthesis (15,53,54). The presented work shows that changing temperatures and carbon sources together can regulate the biosynthesis of BChl c homologs and the self-assembly of the chlorosome.

In this report, we studied *C. tepidum* grown with various carbon sources at different temperatures and noted that Culture 6, which is a temperature down-shifted culture grown on pyruvate/HCO₃⁻, has some unique properties compared with the other cultures. These are as follows:

(a) Combination of carbon assimilation and temperature shift exhibits more effects on chlorosome biogenesis (i.e., Culture 6) compared with changing only one environmental factor.

(b) Slower growth of Culture 6 may be due to more energy consumption in metabolic reactions.

(c) As Culture 6 requires more energy and reducing equivalents produced from light-induced electron transport to produce acetyl-CoA and biomass via the reductive TCA cycle, less resource is available to synthesize BChl c homologs, which require significant energy and carbon sources (55,56). Thus, fewer BChl c homologs are incorporated in the chlorosome from Culture 6.

(d) Culture 6 incorporates most BChl c homologs with a smaller substituent (i.e., an ethyl group) at C-8 attached to the chlorin ring in the chlorosome and produces a chlorosome with blue-shifted Qy absorption maxima. The composition of C-8 substituent has been suggested to contribute to the Qy transition band (34). Furthermore, *bchQ* and *bchR* encode C-8 and C-12 methyltransferase, respectively, and *bchQ(−), bchR(−), and bchQ(−)bchR(−) C. tepidum* mutants were reported to exhibit a blue-shifted Qy maxima (57). We suggest that the blue-shift can be mainly assigned to small variations in the structure of BChl aggregates composing the chlorosome.

(e) Culture 6 producing most of BChl c homologs with the ethyl group at C-8 and C-12 in the chlorosome suggests that C-12 methyltransferase is expressed, as C-12 can be either a methyl or ethyl group, and that C-8 methyltransferase is down-regulated as the C-8 substituent, which can be an ethyl, propyl, isobutyl, and neopentyl group, is largely an ethyl group.

**FIGURE 9** Structure I: Length dependence of CD spectra. Two types of CD spectra are shown for rolls of the radius *R* = 7 nm and the lengths *L* = 100 nm and *L* = 20 nm. The transition between the mixed-type and Type I spectra occurs independently of the roll radius at about *L* = 20 nm.

**FIGURE 10** CD spectra: Roll to lamella transition. CD spectra of single rolls (radius *R* = 7 nm and length is *L* = 100 nm) as compared with the spectra of curved lamellae of the same curvature radius and length. The curved lamellae are represented by halves of the rolls cut along the symmetry axis. The relative intensities are normalized to the number of molecules. (A) Structure I shows transition between the mixed-type and Type II spectra. (B) Structure II shows transition between the mixed-type and Type II spectra.
In addition to affecting methylation at C-8 and C-12 substituents, Culture 6 incorporates almost exclusively farnesyl alcohol (C15:3) plus very small amount of geranylgeranyl alcohol (C20:4) as the esterifying alcohol, whereas Culture 4 and other cultures have slightly more geranylgeranyl alcohol (data not shown). It can be also explained as Culture 6 has fewer energy and reducing equivalents available to synthesize a longer esterifying alcohol (i.e., a geranylgeranyl alcohol group). However, all cultures investigated in this study synthesize farnesyl alcohol as the major esterifying alcohol of BChl c, as reported in literature (10,12,13,23), so the change of the esterifying alcohol of BChl c in Culture 6 is not as significant as the changes of C-8 and C-12 substituents.

The chlorosome from Culture 6 exhibits a Type II (−/+−) CD spectrum, whereas the chlorosome from other cultures we investigated in this report displays a mixed-type (−/+−−) CD spectrum. Previous theoretical studies suggest that the variation of CD spectra may arise from different lengths of cylindrical BChl aggregates (45,58). As alternative mechanisms, we propose that the change in CD type can occur because of different folding of aggregates or formation of curved lamellae.

Culture 6 synthesizes a chlorosome with a smaller cross-sectional radius, which may partially arise from fewer BChl c homologs being synthesized and incorporated in the chlorosome. The smaller cross-sectional radius combined with the difference in the CD spectra may further indicate that the roll versus curved lamellae composition of the chlorosome from Culture 6 is different from that of other cultures.

The chlorosome from Culture 6 exhibits some similar properties to the chlorosome from C. aurantiacus, including a Type II (−/+−) CD spectra (7,21,30), Qy absorption maxima (~740 nm), and simpler BChl c homologs with smaller C-8 and C-12 substituents. The functional advantage of Culture 6, which grows under stressed conditions, producing a chlorosome with a blue-shifted Qy absorption maxima still remains to be understood.

Finally, we would like to emphasize that for ambient light conditions, carbon assimilation could be more sensitive to the amount of reaction centers and BChl a biosynthesis than the light-harvesting antenna complexes. Although this question goes beyond the scope of this article, we may provide a qualitative analysis based on our data reported herein, whereas details will need to be further investigated. Previous studies by Blankenship (59), Bryant (60,61), and their coworkers estimate the presence of 150–200 Fenna–Matthews–Olson trimers and 25–40 reaction centers per chlorosome molecule. Assuming that compared with other cultures the surface concentration of reaction centers in Culture 6 (the stressed culture) remains the same and the area of the baseplate scales is proportional to the surface area of the chlorosome, we estimate that Culture 6 contains ~40% fewer reaction centers compared with Culture 4, which should be in the range of 15–25 reaction centers per chlorosome.

CONCLUSION

The biogenesis of the chlorosome has been previously reported to be regulated by light intensity, temperature, and electron sources, but no attention has been given to the issue of carbon assimilation, which is the bottleneck of photosynthesis. The present work suggests that carbon metabolism together with temperature changes regulate the biogenesis of the C. tepidum chlorosome, resulting in substantial structural modification of the chlorosome at different length scales. These structural variations may arise from the amount and types of BChl c homologs being synthesized and incorporated in the chlorosome. Moreover, our molecular modeling studies provide more detailed structural insight into the spatial arrangement of BChls in the chlorosome.

SUPPORTING MATERIAL

Two tables and three figures are available at http://www.biophysj.org/biophys/supplemental/S0006-3495(13)00845-X.

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FIGURE 11 Computed Qy transition frequencies of BChl homologs. The transition frequencies are computed using time-dependent density functional theory. The following homologs are shown: m/z = 793.5 (8-ethyl, 12-methyl), m/z = 807.5 (8-ethyl, 12-ethyl), m/z = 823.5 (8-propyl, 12-ethyl), m/z = 839.5 (8-isobutyl, 12-ethyl), and m/z = 853.5 (8-neopenyl, 12-ethyl).
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