Molecular biology and genetics / Biologie et génétique moléculaires

**Why is it so difficult to construct Q\textsubscript{i} site mutants in *Chlamydomonas reinhardtii*?**

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Received 13 December 2007; accepted after revision 3 April 2008

Available online 9 May 2008

Presented by Philippe Morat

**Abstract**

The cytochrome b\textsubscript{6}f complex catalyses electron transfer from plastoquinol to a hydrosoluble acceptor (plastocyanin), while building up an electrochemical proton gradient. Oxidation and reduction of plastoquinol occur respectively at the Q\textsubscript{o} site (exposed on the luminal side of the thylakoid membrane) and at the Q\textsubscript{i} site (facing the stroma). The discovery of an additional c\textsuperscript{'}-type heme in the Q\textsubscript{i} site has cast a new light on the difficulties previously encountered to obtain mutants at this site. In this work, we critically examine our unsuccessful attempts to obtain Q\textsubscript{i} site mutants based on sequence and structure homology between cytochrome b\textsubscript{6}f and bc\textsubscript{1} complexes.

To cite this article: A. de Lacroix de Lavalette et al., C. R. Biologies 331 (2008).

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**Résumé**

Pourquoi est-il si difficile d’obtenir des mutants du site Q\textsubscript{i} chez *Chlamydomonas reinhardtii* ? Le complexe cytochrome b\textsubscript{6}f assure le transfert d’électrons depuis un transporteur liposoluble, la plastoquinone, vers un transporteur hydrosoluble, la plastocyanine ; ce processus est couplé à une translocation de protons de part et d’autre de la membrane du thylacoïde. L’oxydation et la réduction du plastoquinol (PQH\textsubscript{2}) s’opèrent respectivement au niveau des sites Q\textsubscript{o} (lumen) et site Q\textsubscript{i} (stroma). La découverte d’un hème de type c\textsuperscript{'} dans le site Q\textsubscript{i} jette une lumière nouvelle sur nos difficultés antérieures à obtenir des mutants de ce site. Dans cet article, nous réexaminons de façon critique notre travail de mutagenèse dirigée basée sur l’homologie de séquence et de structure entre les complexes cytochrome b\textsubscript{6}f et bc\textsubscript{1}. Pour citer cet article: A. de Lacroix de Lavalette et al., *C. R. Biologies* 331 (2008).

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**Keywords:** Cytochrome b\textsubscript{6}f complex; Q\textsubscript{i} site; Site-directed mutagenesis; Heme; Photosynthesis

**Mots-clés:** Complexe cytochrome b\textsubscript{6}f ; Site Q\textsubscript{i} ; Mutagenèse dirigée ; Hème ; Photosynthèse

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1631-0691/S – see front matter © 2008 Académie des sciences. Published by Elsevier Masson SAS. All rights reserved. doi:10.1016/j.crvi.2008.04.003
1. Introduction

Quinol-oxidizing complexes – the cytochrome $bc_1$ complex of mitochondria and bacteria or its chloroplast counterpart, the cytochrome $b_6f$ complex – are the most widespread protein complexes among energy-transducing membranes. They oxidize quinol while reducing a soluble electron carrier to generate an electrochemical gradient of protons across the membrane. According to the Q-cycle proposed by Mitchell [1], the oxidation of a plastoquinol ($PQH_2$) at the $Q_o$ site (lumen) releases two protons and two electrons, one of which reduces a high-potential chain formed by the Rieske protein and a $c$-type cytochrome – cytochrome $f$ in the case of cytochrome $b_6f$ complexes – both of which are membrane-bound. The second electron is transferred to cytochrome $b_L$ then to cytochrome $b_H$, which together constitute the low-potential chain. Re-oxidation of hemes $b_H$ and $b_L$ occurs at the $Q_i$ site (facing the stroma) through the reduction of a plastoquinone (PQ) molecule, generating a cyclic electron transfer around the $b$ heme chain.

While numerous data and mutation studies have provided a fairly good description of the $Q_o$ site from both cytochrome $b_6f$ [2,3] and $bc_1$ (for a review see [4]) complexes, much less is known about structure/function relationships within the $Q_i$ site. Some information on the localisation of the $Q_i$ site in $bc_1$ complexes were drawn from mutagenesis approaches and studies of inhibitor resistance in bacteria and yeast [4–6]. By contrast, information pertaining to the residues that form the $Q_i$ site of cytochrome $b_6f$ complex and to its structure has long remained scarce. Before the cytochrome $b_6f$ structure was solved at atomic resolution in both *Chlamydomonas reinhardtii* and *Mastigocladus laminosus* [7,8], our structural knowledge on this protein heavily depended on predictions based on the structure of the homologous $bc_1$ complex [9–11]. Indeed, the major strategy to design site-directed mutations of the cytochrome $b_6f$ complex was based on sequence homology between cytochromes $b_6f$ and $bc_1$. As will be described here, this strategy turned out to be poorly successful for the study of the $Q_i$ site.

Despite their similarities, $bc_1$ and $b_6f$ complexes feature fundamental functional and structural differences. In oxygenic photosynthesis, the electron pathway can switch from a linear mode, producing both ATP and reducing equivalents, to a cyclic one, where electrons transferred from cytochrome $b_6f$ to PSI on the luminal face of the membrane return to $b_6f$ complex on the stromal face, with production of ATP, but not of reducing equivalents. In addition, the $b_6f$ complex, by activating a protein kinase, plays a key role in state transitions, a major regulatory process of the photosynthetic apparatus that has no counterpart in the respiratory chains [3,12,13]. Finally, cytochrome $b_6f$, as compared to cytochrome $bc_1$, accommodates three additional cofactors, whose function still remains poorly understood: a chlorophyll $a$, a $\beta$-carotene and a heme named $c_1$, with $E_m = -100$ mV [14,15]. The discovery in the X-ray structure of this additional heme, which contributes to the formation of the $Q_i$ site, was a major surprise and gave an important boost to the study of structure/function relationships of cytochrome $b_6f$’s $Q_i$ site. Since this site has to accommodate both a substrate and heme $c_1$, its structure indeed is quite different from that of cytochrome $bc_1$, even if the polypeptide chain around $Q_i$ has the same topology in the $b_6f$ and $bc_1$ complexes. This could account for the lack of inhibitory effect of antimycin on cytochrome $b_6f$ and, in fine, for strong discrepancies between the two complexes that have long remained unexplained.

The scarcity of mutagenesis data also originates from the fact that cyanobacteria do not tolerate severe alterations in the function of the $b_6f$ complexes, while site-directed mutagenesis of cytochrome $b_6f$ subunits remains technically difficult in land plants. Since the green alga *C. reinhardtii* shows dispensable photosynthesis and is amenable to chloroplast gene transformation, it is best suited for structure/function relationship studies of the $Q_i$ site [16,17].

In *C. reinhardtii* the cytochrome $b_6f$ complex comprises four major subunits [15]: the Rieske protein, binding a Fe$_2$S$_2$ cluster, is encoded by the nuclear gene *petC* while the chloroplast genes *petA*, *petB* and *petD* respectively code for cytochrome $f$ – a $c$-type cytochrome with $E_m$ of +330 mV [14] –, cytochrome $b_6$, which binds the $b_H$, $b_L$ and $c_1$ hemes, with $E_m$s of −84 mV, −158 mV and −100 mV [14,15] – and subunit IV. Sequence comparisons have shown that cytochrome $b_6$ and subunit IV from cytochrome $b_6f$ are respectively homologous to the N- and C-terminal parts of mitochondrial and bacterial cytochrome $b$ [18]. The complex also comprises four additional small subunits, the products of the chloroplast genes *petG* and *petL* and of the nuclear genes *PETM* and *PETN* [15,19–22], which have no homologues in the cytochrome $bc_1$ complex.

Here, we critically re-examine our previous attempts to obtain mutants of the $Q_i$ site, impossible to interpret in the absence of the 3-D structure. Mutant construction was based on sequence and structure homology between cytochromes $b_6f$ and $bc_1$, which display, as described previously, fundamental functional and structural differences. The recent X-ray structure of $b_6f$ complex [8]...
has allowed a successful strategy of mutagenesis, but even then we faced some paradoxes, which we will try to discuss.

2. Results

2.1. Before the structure

In cytochrome $bc_1$ the $Q_i$ site is formed by cytochrome $b$ and most mutations altering the functionality of the $Q_i$ site target the amino-terminal part of cytochrome $b$ or the $de$ loop connecting helices D and E [4]. Cytochrome $b_6$ and subunit IV from cytochrome $b_6f$ are homologous to the N- and C-terminal parts, respectively, of mitochondrial and bacterial cytochrome $b$ [18]. It was therefore expected that both subunits would contribute to the formation of the $Q_i$ site.

Two residues in helix A from cytochrome $b_6$ of $C. reinhardtii$, bG37 and bF41, were targeted since substitutions of the corresponding residues in Rhodobacter’s cytochrome $b$ ($G48$ and $A52$ respectively) had provided interesting results [23]: in Rhodobacter, substitution of $G48$ by an Ala residue preserved a wild-type phenotype in the $G48A$ mutant, but replacements with Val or Asp residues led to impaired re-oxidation of $b_H$ causing the loss of photosynthetic growth capability or to failure to assemble the $bc_1$ complex respectively; when Rhodobacter residue $A52$, equivalent to residue bF41 in $C. reinhardtii$, was changed into a Val, the resulting mutant strain displayed a 5-fold slowing down of $Q_i$ site activity and became resistant to antimycin, a specific inhibitor of the $bc_1$ $Q_i$ site that has no effect on the $b_6f$ complex [23].

By site-directed mutagenesis of plasmid pWB encompassing the $petB$ coding sequence [24], we substituted the conserved Gly37 of cytochrome $b_6$ by Val or Asp residues in order to change the local bulkiness and polarity respectively, while residue Phe41 was mutated to Gly to reduce the local bulkiness (Fig. 1). In addition, we also substituted for Ala the residue Phe44, located one helix turn further from the conserved Gly37.

These mutated $petB$ genes were introduced to the chloroplast genome of $C. reinhardtii$ by biolistic transformation, according to two strategies used in parallel (see Section 4). In the first one, deletion strains – here $\Delta petB$ [24] – were used as recipient and transformed clones were selected on minimum medium for the recovery of photosynthetic capability, when the mutations preserve the function of the cytochrome $b_6f$ complex. In the second one, the mutations were first associated with the $aadA$ [25] cassette, conferring resistance to the antibiotic spectinomycin, inserted into a neutral site [26]. Transformed clones, derived from a wild-type recipient strain, could be recovered on spectinomycin-supplemented medium, even when defective for cytochrome $b_6f$ complex activity.

Two mutants (bF41G and bF44A) were recovered after transformation of strain $\Delta petB$, following the capability of the deleted strain to recover photosynthesis. Characterisation of these mutant strains did not reveal...
any drastic changes, since they accumulated the $b_\text{h}_6f$ complex to a wild-type level, were still sensitive to the NQNO inhibitor of the $Q_\text{i}$ site, and showed normal turnover of the complex [27].

By contrast, no phototrophic clones could be recovered upon transformation of the $\Delta petB$ strain with $petB$ genes carrying substitutions of the Gly37 residue, suggesting that these mutations are detrimental to photosynthesis. Consistently, spectinomycin-resistant transformants obtained from the wild-type recipient strain displayed fluorescence induction kinetics typical of mutants with impaired cytochrome $b_6f$ activity [26, 28], while Western blot analysis revealed that cytochrome $b_6f$ subunits fail to accumulate. Since similar substitutions in the $bc_1$ complex have widely different effects, our results pointed to some structural differences between the $Q_\text{i}$ sites of the two complexes, as was already suggested by their different sensitivity to inhibitors [29].

The structure of the cytochrome $bc_1$ complex that became available in the late 1990s [9, 11, 30] (PDB accession 1bcc and 3bcc) allowed us to select other residues putatively contributing to the formation of the $Q_\text{i}$ site. We chose the residues Arg207 at the C-terminus of cytochrome $b_6$, which we substituted for Asp, carrying an opposite charge, as well as residues Asp35 and Phe120, at the N-terminus and in the $fg$ loop of subunit IV, which we modified to Asn and Asp respectively, in order to change both bulkiness and polarity (Fig. 1). The wild-type $\Delta petB$ or $\Delta petD$ strain – this latter carrying a deletion of the $petD$ gene [24] – was transformed with site-directed mutagenised plasmid pWB or pWQ [24], with and without the $aadA$ selectable marker [25]. Again, we could not recover phototrophic clones upon transformation of the $\Delta petB$ or $\Delta petD$ deletion strains and, indeed, the mutants selected for resistance to spectinomycin lacked any accumulation of the $b_6f$ complex (Fig. 2).

We also substituted a Val for Cys35, a residue specifically conserved in all cytochrome $b_6$, but only present in those very few cytochromes $b$ that retain a heme upon denaturation by SDS (see Ref. [31]). Meanwhile, Cys35 was identified by X-ray diffraction as the residue to which heme $c_i$ is covalently attached [8]. The C35V mutant was unable to grow phototrophically because this substitution prevented the accumulation of cytochrome $b_6f$ subunits [32].

2.2. After the structure

We have already pointed out how the unexpected discovery of the $c_i$ heme in the $Q_\text{i}$ site of the cytochrome $b_6f$ complex [7, 8] has given a new impetus to the study of structure/function relationships in cytochrome $b_6f$. The crystallographic structure provided a sound structural background to the construction of new mutants altering the $Q_\text{i}$ site. We chose three residues in the proximity of the $c_i$ heme: Leu36, Phe40 and Pro41 of subunit IV (Figs. 1 and 3). Leu36, standing at the entrance to the $Q_\text{i}$ site, was substituted for Ala to enlarge the entrance of this site, while Tyr/His or Leu residues were substituted for Phe40 that may shield the iron of heme $c_i$ from the external medium so as to provide a possible axial ligand to iron of heme $c_i$ or to test its protective role. We also tested whether residue Pro41 imposed
Fig. 3. View of Q\textsubscript{i} site (1Q9 pdb data). Mutated residues are located in the structure; in orange: mutations leading to non-assembled complexes and non-photosynthetic strains (on cytochrome b\textscript{6}: bC35V, bG37V, bR207D and on subunit IV: dD35N, dF120D), in green: mutations leading to photosynthetic strains (on cytochrome b\textscript{6}, bF41G, bF44G), and in yellow: mutations leading to photosynthetic strains that could only be selected on spectinomycin (on subunit IV: dL36A, dF40L/H/Y and dP41A).

Structural constraints to the environment of heme c\textsubscript{i} by mutating it for an Ala residue.

Our attempts to recover phototrophic clones upon transformation of the \(\Delta\)petD strain were unsuccessful, leading us to suppose that also these mutations were detrimental for photosynthesis. We then used the wild-type strain as a recipient for transformation experiments with plasmids carrying the specific mutations associated with the aadA cassette. Unexpectedly, the transformed clones recovered on antibiotic-supplemented medium retained wild-type fluorescence induction kinetics, even after they reached homoplasmy for the mutation (when all copies of the polyploid chloroplast genome carry the mutation while all wild-type versions of the petD gene have been eliminated). Paradoxically, the mutations therefore were not harmful to photosynthesis, even though the mutated genes were not able to rescue deletion mutants upon transformation.

3. Discussion

For more than ten years, we have tackled the ‘mystery’ of the Q\textsubscript{i} site. The resolution of the 3-D structure of the cytochrome \(b_6f\) complex with the discovery of an additional heme in the Q\textsubscript{i} site has partially solved this puzzle and shed a new light on the molecular organisation of this site. Heme c\textsubscript{i} has quite disconcerting features, such as a single thioether linkage to the protein, no amino acid side chain serving as an axial ligand, and, possibly, a molecule of water or an hydroxide ion as a fifth putative ligand. Since we are not able yet to draw a precise model for the role of the c\textsubscript{i} heme, the critical analysis of previous alterations to the Q\textsubscript{i} site may help us to understand its peculiar structure and organisation.

The mutated amino acids can now be located on the 3-D structure (Fig. 3) \[8\] in order to ascertain the structural effects of the mutations on the Q\textsubscript{i} site environment and to correlate it with the corresponding phenotype.

Gly37 of cytochrome b\textsubscript{6} (Fig. 3) residue coloured in red on the cytochrome b\textsubscript{6} helix A in cyan), is at 3.7 Å from the heme \(b_H\) porphyrin cycle, with which it may be in Van der Waals contact. Accommodating a Val residue in mutant bG37V may require a displacement of helix A, which is possible in cytochrome \(b_{6f}\), since the corresponding mutant does not display any effect on the assembly. By contrast, the thioether linkage between c\textsubscript{i} heme and Cys35 at the stromal end of helix A requires a precise protein environment and this displacement probably prevents cytochrome \(b_6f\) assembly, as suggested by the phenotype of the mutant. Similarly, substitution of Cys35 (Fig. 3, residue coloured in orange on the cytochrome b\textsubscript{6}, helix A in cyan) led to a phenotype much very similar to that of \(b_H\) mutants in \(C.\text{reinhardtii}\) \[32,33\]. This Cys residue is conserved in all species carrying a heme covalently bound to the protein moiety, either on chloroplast cytochrome \(b_6\) or on cytochrome \(b\) from thermophilic bacilli \[34\].
The detrimental effect of substitution for an Asp of residue Arg207 (Fig. 3, residue coloured in orange on the cytochrome \( b_6 \), helix D in cyan), which interacts with the propionate group of heme \( c_1 \), may be related to the radical change in the \( Q_i \) local charge. Arg207 participating to a network that involves propionate, water molecules and plastoquinone, the substitution probably modifies the interaction between plastoquinone and heme \( c_1 \).

Substitutions of the two Phe residues of cytochrome \( b_6 \) in positions 41 and 44 (Fig. 3, residues coloured in green on the cytochrome \( b_6 \), helix A in cyan) led to a wild-type phenotype. These residues are far from the \( Q_i \) site and point rather towards the \( b_1 \) heme. A reduction in their bulkiness did alter neither the folding nor the accumulation of the complex [27].

Two other mutations were carried out on subunit IV (Figs. 1 and 3), both of which yielded non-phototrophic mutants. Residue Asp35 (residue coloured in orange on the subunit IV, helix E in yellow) is in strong electrostatic interaction with residue Arg95 from the PetN subunit. Its substitution with Asn by removing the negative charge changes the local polarity, which seems important for the conformation of the site and the stability/assembly of the complex, as shown by the phenotype of the mutant. It is more difficult to explain the detrimental effect of the Phe120 \( \rightarrow \) D mutation, since residue dPhe120 (coloured in orange on the subunit IV, \( f_g \) loop in yellow) is located quite far away from the \( Q_i \) site and does not seem to interact with other cofactors. However, Phe120 is close to a positively charged patch (Arg112 on cytochrome \( b_6 \), Tyr119 and Arg125 on subunit IV); its replacement by a negatively charged residue would alter the charge network. Hydrogen bridges may rigidify the structure, preventing efficient assembly of the complex.

Our successful attempts to obtain \( Q_i \) site mutants were based on the available 3-D structure. However, we could not rescue the \( \Delta \)petD deletion strain for photosynthetic growth upon transformation with the mutated plasmids. Paradoxically, after selection for resistance to antibiotics, the dL36A, dF40H/Y/L and dP41A mutants nevertheless turned out to be capable of phototrophic growth. The molecular basis of this paradoxical observation is not fully understood yet and was first hypothesised to reflect the complex organization of the \( Q_i \) site. As evoked above, the full structure of this site is particularly complex, since it requires interactions between two subunits, cyt. \( b_6 \) and subunit IV, two cofactors, hemes \( b_H \) and \( c_1 \) and a liposoluble electron carrier, plastoquinone. It may thus be very sensitive to changes in the proteic environment. However, those mutations altering the conformation of the \( Q_i \) site were not detrimental for the assembly of cytochrome \( b_6 f \), since wild-type levels of the complex are found in the mutants recovered on spectinomycin-supplemented medium (data not shown). Mutants defective for cytochrome \( b_6 f \) activity are light-sensitive, probably because the fruitless charge separation occurring in photosystems I and II leads to the formation of deleterious radical species. Radicals should be produced at high levels during the initial selection step of phototrophic transformants derived from deletion strains under high light. By contrast, their concentration should remain low when transformants are selected on spectinomycin-supplemented medium under low light from a wild-type recipient strain, which is capable to transfer electrons from PSII to PSI. This suggests an increased sensitivity to radical species (like the reactive oxygen species, ROS) of the mutants with altered \( Q_i \) site, which would be specific of this site, since mutants with altered conformation of the \( Q_o \) site do not present the same phenotype. This hypothesis may indicate a possible protective function for the \( c_1 \) heme, which is currently under investigation.

4. Materials and methods

4.1. Strains, media, and growth conditions

A wild-type strain derived from strain 137C [35] and the deletion strains \( \Delta \)petD and \( \Delta \)petB [24] were used as recipient strains in chloroplast transformation experiments. WT and mutant strains were grown on Tris-acetate-phosphate (TAP) medium (pH 7.2) at 25°C, under dim light (5–6 \( \mu \)E m\(^{-2} \) s\(^{-1} \)), unless otherwise indicated.

4.2. Site directed-mutagenesis

Plasmids pDWQ and pDWB, encompassing the regions coding for subunit IV and cytochrome \( b_6 \) respectively, have been described previously [24]. Mutations on cytochrome \( b_6 \), Cys35Val, Gly37Val, Phe41Gly, Phe44Gly, Arg207Asp and on subunit IV Asp35Asn and Phe120Asp were introduced in plasmids pWB or pWQ, respectively, by site directed mutagenesis according to Kunkel [35].

Mutations Leu36Ala, Phe40Leu/His/Tyr and Pro41Ala were introduced in plasmid pWQ by PCR-mediated site-directed mutagenesis [36]. The \( aadA \) cassette conferring resistance to spectinomycin [25] was then inserted into the pWB or pWQ plasmids carrying the various mutations, respectively at the NsiI or EcoRV neutral restriction sites [2,26].
4.3. Chloroplast transformation in C. reinhardtii

Wild-type as well as ΔpetB and ΔpetD strains, respectively deleted for petB or petD genes, were transformed by tungsten particle bombardment according to Boynton et al. [37]. Transformants were selected either on minimum medium under dim light (60 µE m⁻² s⁻¹) or on TAP medium supplemented with spectinomycin (100 µg ml⁻¹) under low light (10 µE m⁻² s⁻¹). They were submitted to several rounds of subcloning on selective medium until they reached homoplasmia, assessed by RFLP analysis of specific PCR amplification fragments.

4.4. Preparative and analytical techniques

For SDS-PAGE, membrane (or whole cells) proteins were resuspended in 100 mM dithiothreitol and 100 mM 4.4. Preparative and analytical techniques fragments. Preparative and analytical techniques fragments. fragments.

4.5. Fluorescence measurements

Fluorescence measurements were performed at room temperature on a home-built fluorimeter, using a light source at 590 nm. The fluorescence response was detected in the far-red region in the near IR region [26].

Acknowledgements

The authors are grateful to Yves Choquet and Jean-Luc Popot for their critical reading of the manuscript. Thanks are due to Daniel Picot for the fruitful discussion and the help in drawing Fig. 3. F.Z. whishes to thank Francis-André Wollman for his fundamental input in this study. A.L.L. was the recipient of a fellowship from the ‘Ministère de l’Enseignement supérieur et de la Recherche’, France.

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