Modeling the electron transport chain of purple non-sulfur bacteria

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Introduction

Our current understanding of some of the most fundamental processes of life, such as the photosynthetic generation of cellular energy or the metabolic fixation of carbon dioxide and nitrogen, has been driven largely by investigating a unique group of facultative photosynthetic bacterial organisms, the purple non-sulfur bacteria (Rhodospirillaceae). The first purple bacterial reaction center (RC) was crystallized in 1985 from *Rhodopseudomonas viridis* (Deisenhofer et al., 1985), and high-resolution crystal structures of the light-harvesting complexes (LHCs) LH2 from *Rhodopseudomonas acidophila* (McDermott et al., 1995) and LH1 from *Rhodospirillum rubrum* (Karrasch et al., 1995) followed, thus providing a structural basis for the understanding of the photochemical reactions taking place in the photosynthetic apparatus. Furthermore, purple non-sulfur bacteria exhibit a remarkable metabolic versatility (Tabita, 1995) that, in recent years, has put a focus on studying regulatory mechanisms that govern gene expression in response to varying redox conditions of the environment (Bauer et al., 2003). Purple non-sulfur bacteria constitute an excellent model system for this purpose due to the presence of intracytoplasmic membranes (ICMs), harboring the photosynthetic RC and LHCs, which are produced depending on the environmental factors such as oxygen, light and growth substrates. The formation of ICM can easily be measured in vivo and provides a phenotypic marker for the activation of photosynthetic genes. For several species, complete genome sequences are now available (http://genome.jgi-psf.org/mic_home.html) and molecular methodologies for constructing mutants have been developed. During the past years, some conserved regulatory systems involved in the control of ICM expression have been identified, including the global and highly conserved RegA/RegB (PrrA/PrrB) two-component signal transduction system (Elsen et al., 2004), the FpsR (CrtJ) repressor (Elsen et al., 2005; Kovacs et al., 2005) and the Fnr-like transcriptional regulator FnrL (Zeilstra-Ryalls et al., 1997; Zeilstra-Ryalls and Kaplan, 1998). However, particularly for light regulation of ICM synthesis, the molecular signals that are involved in the initiation of signal transduction

Purple non-sulfur bacteria (Rhodospirillaceae) have been extensively employed for studying principles of photosynthetic and respiratory electron transport phosphorylation and for investigating the regulation of gene expression in response to redox signals. Here, we use mathematical modeling to evaluate the steady-state behavior of the electron transport chain (ETC) in these bacteria under different environmental conditions. Elementary-modes analysis of a stoichiometric ETC model reveals nine operational modes. Most of them represent well-known functional states, however, two modes constitute reverse electron flow under respiratory conditions, which has been barely considered so far. We further present and analyze a kinetic model of the ETC in which rate laws of electron transfer steps are based on redox potential differences. Our model reproduces well-known phenomena of respiratory and photosynthetic operation of the ETC and also provides non-intuitive predictions. As one key result, model simulations demonstrate a stronger reduction of ubiquinone when switching from high-light to low-light conditions. This result is parameter insensitive and supports the hypothesis that the redox state of ubiquinone is a suitable signal for controlling photosynthetic gene expression.
pathways leading to photosynthetic gene expression are poorly characterized or even still unidentified. In a remarkable forward looking perspective, Cohen-Bazire et al. (1957), who systematically investigated the overall effect of O₂ and light on synthesis of bacteriochlorophyll (BChl) and carotenoids in *Rhodobacter sphaeroides* and *R. rubrum*, concluded that the oxidation–reduction (redox) state of one component or several components of the electron transfer systems of photosynthesis and respiration functions as a cellular signal for the activation of photosynthetic genes. Despite the fact that at that time the composition of the electron transfer chains was unknown and the assumptions about basic photochemical events were wrong, their concept of redox signaling constituted a major guideline for the following biochemical and molecular biological analysis of regulatory mechanisms that control ICM synthesis in Rhodospirillaceae. This endeavor has been challenging until today. Recently, it has been hypothesized that the redox state of the membranous ubiquinone pool could play a major role in the transcriptional control of gene expression (Oh and Kaplan, 2001; Swem et al., 2006). However, direct experimental evidence in the form of a signal–response curve is lacking up to now, and quantitative data obtained *in vivo* of the responses of cellular redox states are scarce. Thus, despite the details now available about the molecular structures and mechanisms involved in these processes, we are still far from having a comprehensive and quantitative insight into the overall process. The estimation of how the redox states of the different components of the photosynthetic and respiratory electron transport systems will respond to a changing redox environment, for example, the availability of electron acceptors or light, is very hard to achieve by pure intuition and also difficult to determine experimentally. Therefore, mathematical modeling is an important tool for obtaining a more holistic understanding of redox metabolism and redox-mediated control of gene expression in these organisms.

In this work, we present a model of the electron transport chain (ETC) in purple bacteria that describes the qualitative effects of different redox conditions on the redox states of electron transfer components. Even though most conclusions drawn from the model simulations should be relevant for many representatives of the Rhodospirillaceae, specific parts of the model are tailored to *R. rubrum* for the following reasons: (i) The individual components of the ETC are sufficiently characterized and a large amount of physiological data are available from earlier studies (including NADH/NAD and ATP/ADP ratios provided by the work of Schön, 1969, 1971). (ii) The ratio of RCs and LH1 expression is fixed due to activation of *puf* and *puh* operons coding for the RC and LH1 polypeptides. Since *R. rubrum* does not possess LH2, no additional interfering regulatory circuits have to be considered in the model. (iii) With the development of a special growth medium for *R. rubrum* (M2SF medium) containing the two carbon sources succinate and fructose, Ghosh et al. (1994) provided a valuable experimental tool for addressing aspects of redox signaling. Compared to a standard medium (with succinate only), growth of *R. rubrum* with the M2SF medium results in drastically increased levels of photosynthetic membranes (PM) under semi-aerobic conditions *in the dark*, which so far were only obtainable under anaerobic low-light conditions. Here, we verify our previous hypothesis, that the effect of M2SF medium could be equivalent to redox signaling under low-light conditions, particularly at the level of the ubiquinone redox state (Grammel et al., 2003).

As stated above, although the model considered here is tailored to *R. rubrum*, most of the results are valid also for other representatives such as *Rb. capsulatus* and *R. sphaeroides* and it should be easy to adapt the model to address specific questions. The composition of the ETC in these organisms is homologous, with only minor differences in the nature of the quinone and cytochrome species.

We start the model-building procedure by compiling a list of relevant components and processes yielding first a stoichiometric model of the ETC. As we will show, even such a relatively simple model enables to identify all potential operational modes of the ETC. We will then incorporate kinetic terms to describe driving forces and reaction rates as well as—in a more coarse-grained manner—regulatory events. The primary goal of this kinetic model is to deliver robust qualitative predictions on the steady-state operation of the ETC and on how the redox states of ETC components behave under different environmental conditions.

Kinetic ETC models have already been constructed and analyzed previously, for example, for the respiratory electron transport phosphorylation in mitochondria (Korzeniewski, 1996a, b; Korzeniewski and Mazat, 1996; Beard, 2005) and for photosynthesis in higher plants and algae (Berry and Rumberg, 2000). However, although respiration in mitochondria and photosynthesis in plants share some similarities with the ETC in purple bacteria, these models cannot directly be used for our purpose. More recently, Geyer and Helms (2006) and Geyer et al. (2007) have proposed kinetic models for photosynthesis in *Rb. sphaeroides*, which were helpful, for example, in deriving stoichiometric relationships between ETC components. However, respiration and the coupling of the ETC with the central metabolism were not considered; hence, these models cannot be employed for simulating redox states of ETC components under varying growth conditions—a major objective of this work.

**Results**

**Stoichiometric model of the ETC**

**Components and processes**

Anoxogenic photosynthesis (under anaerobic conditions in the light) and respiration (under aerobic conditions) are the two major strategies of the Rhodospirillaceae for generating energy in the form of ATP. For both the metabolic regimens, the key processes take place in the membranous ETC, which couples the transfer of electrons, driven by redox potential differences to the generation of proton-motive force (∆p), thus catalyzing the synthesis of ATP. Figure 1 shows a scheme of the processes and components involved neglecting, for the sake of simplicity, that the Rhodospirillaceae possess actually two structurally and functionally different types of membranes. The cytoplasmic membrane (CM), harboring the components of the respiratory chain, and the ICMs, which can be isolated as chromatophores (Drews and Golecki, 1995) with the photosynthetic RCs and the LHCs. While some electron transfer
components such as NADH dehydrogenase, succinate dehydrogenase, cytochrome bc₁, ATP synthase and ubiquinone are constituents of both types of membranes, others such as RC, LHC and cytochrome oxidases appear to be exclusively located in either the photosynthetic ICM or respiratory CM, respectively. Thus, CM catalyzes respiratory phosphorylation whereas ICM exclusively catalyzes photophosphorylation.

Since the precise topological nature and stoichiometric composition of the two membrane compartments and their functional interactions are not yet elucidated in sufficient quantitative detail, we will not distinguish between the two types of membranes in our model, and consider instead one (combined) membrane pool. Accordingly, we also model the periplasmic space as one homogenous compartment. These assumptions simplify considerably the complicated real situation, however, as we will focus in this study on purely respiratory or purely photosynthetic growth where the main activity of the ETC is located either exclusively in the CM (respiration) or exclusively in the ICM (photosynthesis) we argue that a compartmentalized model would not change the main results of this study.

In the model, the membrane will function either as a respiratory ETC or as a photosynthetic ETC and functional interactions between respiration and photosynthesis cannot take place. The different stoichiometry of CM and ICM (in particular regarding the photosynthetic apparatus) is considered in the model by incorporating regulatory pathways governing the expression of membrane components.

In the following, we compile a list of redox carriers and enzyme complexes of the ETC considered in our model (Figure 1).

Diffusible redox carriers in the membrane are ubiquinone (Q) and cytochrome c₂ (c₂). While Q is localized in the membrane, c₂ is diffusible at the P side of the membrane close to the periplasmic space. Alternative quinones (rhodoquinone in R. rubrum; Imhoff and Bias-Imhoff, 1995) with a more negative standard redox potential than ubiquinone are known. However, since the role of these players is not completely understood, we will not consider them herein. Also, cytochromes other than c₂ are known (Vermeglio et al, 1995; Zannoni, 1995), which seem to be involved mainly in respiration. Here, we will consider only one (combined) cytochrome c pool whose representative will be c₂.

NADH is a universal cytoplasmic carrier of reducing equivalents. In our model, NADH will represent a ‘boundary’ metabolite: it couples the membranous ETC with the ‘rest’ of the central (redox) metabolism, either by feeding electrons into the ETC or by taking up electrons from the ETC.

The NADH-Dehydrogenase (NADH-DH) couples the transfer of two electrons from NADH to Q (yielding ubiquinol, QH₂) to the translocation of cytoplasmic protons $H_i^+$ into the periplasm ($H_p^+$; here, we assume a stoichiometry of four pumped protons). The net stoichiometry of the NADH-DH thus reads:

$$\text{NADH} + \text{Q} + 5H_i^+ \leftrightarrow \text{QH}_2 + 4H_p^+ + \text{NAD}^+$$  (1)

Under certain circumstances, the NADH-DH can also run in the reverse direction. Purple bacteria make use of this reverse electron flow particularly under phototrophic conditions.

The succinate dehydrogenase (Succ-DH) oxidizes succinate to fumarate and passes the two electrons to ubiquinone. Protons are not pumped in this process.

$$\text{Succinate} + H_2O \leftrightarrow \text{Fumarate} + \text{QH}_2$$  (2)

Succ-DH is a key enzyme of the citric acid cycle, and couples the central carbon metabolism to the ETC at the level of Q.

Reaction (2) may also run in the reverse direction, depending...
on the concentrations of the compounds involved. Succ-DH functions then as a fumarate reductase. Alternative fumarate reductases, which are differentially expressed during anaerobic respiration with fumarate as electron acceptor, will not be considered here as additional components. Other enzyme complexes that are able to oxidize metabolites and to feed electrons into the ETC (such as glycerol dehydrogenase) are also not taken into account as they can be assumed to be of low relevance for the environmental scenarios considered herein.

The cytochrome bc₁ complex (bc₁) transfers electrons from the ubiquinol pool to cytochrome c₂ and pumps protons via a mechanism known as the Q cycle (Trumpower, 1990). The net balance of this complicated cycle reads:

$$QH_2 + 2H_2^+ + 2c_2^{3+} \rightarrow 2c_2^{2+} + Q + 4H_2^+$$

(3)

Whereas NADH-DH, Succ-DH and bc₁ are operative under photosynthetic as well as respiratory growth conditions (and are contained in both types of membranes), the following two oxidases are only active under anaerobiosis since they mediate the irreversible transfer of electrons to the terminal acceptor oxygen (other acceptors such as DMSO can be utilized but are not relevant herein).

The cytochrome c oxidase oxidizes two c₂ and passes the electrons to oxygen, thereby pumping protons into the periplasm:

$$0.5O_2 + 2c_2^{3+} + 4H_2^+ \rightarrow 2c_2^{2+} + H_2O + 2H_2^+$$

(4)

Two major cytochrome c oxidases are known in purple bacteria. The cbb₃ oxidase having a high affinity to oxygen, and an aa₃ oxidase with a lower affinity (Zannoni, 1995). Cytochrome oxidases of the cbb₃ type are found, for example, in *Rb. sphaeroides, Rb. capsulatus* and *R. rubrum* (the latter based on gene sequence data, not shown), whereas from these three representatives only *Rb. sphaeroides* has been shown to possess the aa₃-type enzyme. Therefore, we will consider only the cbb₃ oxidase.

The ubiquinol oxidase (containing a b-type heme; Varela and Ramirez, 1990) transfers electrons directly from QH₂ to oxygen:

$$0.5O_2 + QH_2 + 2H_2^+ \rightarrow Q + H_2O + 2H_2^+$$

(5)

The ubiquinol oxidase functions also as a proton pump but the overall H₂O / e⁻ stoichiometry is lower than using bc₁ and cytochrome oxidase. Presumably for this reason, ubiquinol oxidase is mainly expressed under semi-aerobic conditions (and above and Materials and methods section). Photons (υ) are harvested in the LHC and then funneled as excitations to the RC where charge separations take place leading to the reduction of Q and oxidation of c₂:

$$2\nu + Q + 2c_2^{3+} + 2H^+ \rightarrow QH_2 + 2c_2^{2+}$$

(6)

Protons are not pumped during this process. The merit of the photosynthetic apparatus is the light-induced uphill (from high to low redox potential) transfer of electrons from c₂ to QH₂.

Finally, the ATP synthase complex converts generated Δp to chemical energy in the form of ATP. Assuming a H₂O / P ratio of 3, the stoichiometry reads:

$$ADP^3^- + P_i^+ + 3H_2^+ + 2H_2O \rightarrow ATP^4^- + H_2O + 3H_2^+$$

(7)

**Operational modes of the ETC**

So far we have collected the main components and the net stoichiometries of the electron transfer processes in the ETC of purple non-sulfur bacteria. This does not yet say much about how the different processes have to be combined in a way that the ETC can bring about a coherent function. The seven stoichiometric reactions (1–7) give rise to a stoichiometric reaction network that can be described formally by a stoichiometric matrix *N* (rows: species; columns: reactions; Nᵢᵢ := stoichiometric coefficient of species *i* in reaction *j*). We can then apply well-established tools from metabolic network analysis to understand how the ETC can operate in the steady state (Price *et al.*, 2004; Klamt and Stelling, 2006). A suitable approach from this methodology is the formalism of *elementary modes* (EMs) that enables to identify operational modes (Schuster *et al.*, 2000; Klamt and Stelling, 2006). EMs are defined as minimal subnetworks that can perform a function in the steady state. They correspond typically to pathways or cycles in the reaction network. Each EM is characterized by the participating reactions (and the relative fluxes they carry) and no subset of the reactions involved of an EM can sustain a steady-state flux distribution. Importantly, any potential steady-state behavior of a metabolic network can be described as a non-negative superposition of the EMs.

Since the ETC is a subsystem of the whole system ‘cell’, applying the approach of EMs requires the definition of model boundaries. In our case, we have to specify which compounds are coupled to other processes taking place outside the ETC. This concerns (from here, we omit molecule charges) NAD, NADH, ADP, ATP, succinate, fumarate, photons, Pᵢ, H₂O and O₂ which are considered to be *external* metabolites (Klamt and Stelling, 2006). For these metabolites, we do not demand a steady-state (mass balance) condition since they can be produced, consumed, exported or imported elsewhere in the metabolism. In contrast, reduced and oxidized forms of ubiquinone and cytochrome c₂ as well as the protons in the periplasm (H₂O) are considered as *internal* metabolites, that is they have to be kept in a balanced state. Accordingly, the stoichiometric matrix *N* for the internal species of the ETC reads (the seven columns (reactions) correspond to the equations (1–7)):

$$N = \begin{pmatrix}
1 & 1 & -1 & 0 & -1 & 1 & 0 \\
-1 & -1 & 1 & 0 & 1 & -1 & 0 \\
0 & 0 & 2 & -2 & 0 & -2 & 0 \\
4 & 0 & 4 & 2 & 0 & -3 & 0
\end{pmatrix} \rightarrow QH_2$$

(8)

For this stoichiometric configuration of the ETC, in total, nine EMs can be computed (e.g. with *CellNetAnalyzer*; Klamt *et al.*, 2007). The stoichiometry of these modes is summarized in Table I, a schematic representation is given in Figure 2. The EMs have been ordered with respect to the Δp-generating process, namely photosynthesis (EM1 and EM2), oxidative respiration (EM3–EM8) and fumarate reduction (EM9). Most EMs correspond to well-known functions of the ETC in purple...
bacteria. The photosynthetic lifestyle can be described by just two EMs: EM1 is the cyclic photophosphorylation that produces ATP at the expense of absorbed photons by cycling electrons from ubiquinol to cytochrome c2 (via bc1) and back to ubiquinol (via the RC). EM1 reproduces correctly the fact that the cyclic photophosphorylation does not consume electrons in the overall stoichiometry. EM2 reflects another important function of the ETC under photosynthetic conditions, namely reverse electron flow: Δp is generated via photosynthesis and then utilized for an active transfer of electrons from succinate via ubiquinol to cytochrome c2 and back to ubiquinol (via the RC). EM1 reproduces correctly the fact that the cyclic photophosphorylation does not consume electrons in the overall stoichiometry. 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Figure 2  The fundamental operational modes (elementary modes) of the electron transport chain in purple non-sulfur bacteria. Stoichiometries (i.e. relative reaction rates) have been omitted and can be seen in Table I. In the upper left corner, the full network is depicted (compressed version of Figure 1).
Kinetic model of the ETC

The kinetic model—described in detail in the Materials and methods section and in Tables III and IV—comprises the electron transfer processes of the ETC as depicted in Figure 1. The key question in the development of the ETC model was how to describe kinetically the complicated electron transfer processes. Here, we followed the approach of Korzeniewski (1996a, b) and Korzeniewski and Mazat (1996) by modeling the entirety of elementary steps taking place at an enzyme complex as one (overall) redox reaction. The overall stoichiometries of these redox reactions correspond to the equations (1–6) and their driving forces are redox potential differences between the redox couples involved (see Materials and methods). Korzeniewski and co-workers modeled the respiratory chain of mitochondria that exhibits some homologies to the ETC of Rhodospirillaceae, though the latter possesses additionally the photosynthetic phosphorylation cycle.

For quantifying redox potentials (and redox potential differences), one requires the concentration ratios of the reduced/oxidized forms of the relevant redox couples, namely [QH2]/[Q], [c2+]/[c2−], [NADH]/[NAD] and [succinate]/[fumarate] (see Materials and methods). Whereas [Q], [QH2], [c2+], and [c2−] are state variables in the model, we cannot proceed in the same way with [NADH], [NAD], [succinate] and [fumarate] as these compounds are involved in many other metabolic reactions taking place outside the ETC. Here, we face a general problem in modeling cellular processes: how to delimit the modeled processes from the rest of the cell? We employ the following strategy: for each of the growth regimens considered—for example, aerobic growth in the dark or anaerobic growth in light—we clamp the concentration ratios of external metabolites such as [NADH]/[NAD] to a specific value (obtained from measurement data). The advantage is that we do not need to model the whole cell to get a complete balance for these compounds. However, in a simulation, the assumed (steady state) ratio is immediately fixed after changing the environmental conditions. For this reason, we will focus herein on the steady-state behavior of the kinetic model (which is nevertheless fundamentally different to the approach of EMs since steady-state concentrations of metabolites are explicit state variables.). There are also other reasons why our model allows only for steady-state analysis. As we consider only overall reactions inside the protein complexes, any reaction is immediate and does not reflect the time course of events inside the protein that may lead to wrong simulation results for the very early response of the ETC. Moreover, the buffer capacity (captured by the parameter piV; see Materials and methods section), which determines how fast the ETC can adapt after changing metabolite concentrations, is insensitive against the steady-state data used to fit the experiments (see below) and thus badly determined.

Though the main application of our kinetic model is to simulate the steady-state operation of the ETC under different environmental conditions, we need to include basic regulatory events such that the photosynthetic apparatus consisting of LHC and RC is synthesized under anaerobic conditions. As already mentioned above, the composition of the ETC and the formation of photosynthetic membranes in purple bacteria are mainly governed by the external signals oxygen and, to a lower extent, light intensity and the supply of reducing equivalents by the carbon substrate. Three major regulators (RegB/RegA (PrbB/PrrA), ForL and PsrR (CrlU)) have been identified that control the expression of genes involved in the synthesis of the photosynthetic gene cluster (PGC; Alberti et al., 1995) and other ETC components such as ubiquinone, c2 as well as of the ubiquinol and the cytochrome (cbb3) oxidase (see Oh and Kaplan, 2001; Bauer, 2004 and Materials and methods). In a coarse-grained manner, we included kinetic terms in the differential equations for LHC, RC and other ETC components that account for the combined effects of the three regulators (see Materials and methods).

Interestingly, none of the above-mentioned regulators is able to sense light directly. Instead it has been suggested that different light intensities are indirectly measured via the redox state of the ubiquinone pool (Swem et al., 2006). In fact, we want to examine whether our model supports this hypothesis.

In total, the model comprises 33 time-dependent variables of which 10 are states (expressed by ordinary differential equations); the rest depends algebraically upon the state variables (Table III). The model has 55 parameters (Table IV); 17 of them could be directly taken from literature values. The other parameters were estimated by fitting the measurement data—taken from different growth scenarios—as given in Table II. We implemented and simulated the ETC model with self-written MATLAB scripts and implemented it additionally with the Systems Biology Toolbox (Schmidt and Jirstrand, 2006; http://www.sbttoolbox.org/) for MATLAB. The model file for the Systems Biology Toolbox can be downloaded from http://www.mpi-magdeburg.mpg.de/projects/cna/etcmodel.html.

Model analysis and simulations

In the simulations, we focused on four different environmental regimens (Table II): aerobic (R1) and semi-aerobic (R2) in dark conditions; anaerobic under high-light (R3) and low-light (R4) conditions. The four growth regimens are thus characterized by the imposed oxygen tension and mean irradiance, for which we have chosen typical values reported in the literature. As explained above, for each simulation, the three concentration ratios [NADH]/[NAD], [succinate]/[fumarate] and [ATP]/[ADP] have to be clamped (but can be varied in different simulations). We abbreviate these ratios in the following with R_{NADH}, R_{succ} and R_{ATP}, respectively. The respective ratios chosen for the four different growth regimens are also shown in Table II. Regarding R_{succ}, we assumed unity since no appropriate measurements could be found in the literature. For R_{ATP}, some values are available in Schöhn (1969) showing comparable values for aerobic and photosynthetic growth of R. rubrum on succinate. We therefore opted to use an average value of 2.7 for all conditions. Regarding the ratio of R_{NADH}, Schöhn (1971) presented data for three of the four different growth regimens; an estimate was made for low-light growth (R4). In addition to R_{NADH}, R_{succ} and R_{ATP}, we found a number of measurements of ETC components in the literature, which have been used for parameter fitting (see Table II and Figure 3, where the circles indicate a sufficiently good fitting result).
Although the chosen values for $R_{\text{NADH}}$, $R_{\text{Succ}}$ and $R_{\text{ATP}}$ should reflect the situations for the different growth regimens, their precise value will not only depend on the imposed light intensity and oxygen concentration (as assumed in Table II) but also on other external factors such as the reduction degree and energy content of the substrate. Therefore, it makes sense to vary these ratios and to study how this affects the steady-state behavior of the ETC in the different growth regimens.

**Model simulations: varying the cellular [NADH]/[NAD] ratio**

$R_{\text{NADH}}$ is an important measure of the overall redox state of the cell and is strongly coupled with the ETC and therefore of paramount importance for our investigations. For studying the response of the ETC to different levels of $R_{\text{NADH}}$, we varied this value (in all growth regimens) within a physiologically meaningful range between 0.01 and 2.01 and computed the resulting steady states in the system. In this way, we obtained response curves of ETC components with respect to the cytoplasmic redox state (Figure 3), which will be discussed below.

Note that $R_{\text{NADH}}$ will not only influence the redox state of ETC but also depends on the redox state of the ETC. However, here we consider $R_{\text{NADH}}$ as an independent variable so that a certain value of $R_{\text{NADH}}$—for example, measured in a particular experiment—can be directly related to the corresponding electron flows and (redox) states of components in the ETC.

**Aerobic growth (regimen R1)**

As expected, in the fully aerobic case we have the lowest redox charge of the ubiquinone and $C_2$ pool of all growth regimens (see Figure 3: $Q_{\text{charge}}$ ($QH_2/Q_{\text{total}}$) and $c_2_{\text{charge}}$ ($c_2^+/c_2_{\text{total}}$)). We can conclude that a low cytoplasmic redox potential, for example, generated by a strongly reduced substrate, will not limit the operation of the ETC; the opposite is the case as a higher value for $R_{\text{NADH}}$ increases the electron flow. The reason is that oxygen withdraws the electrons—fed in by the NADH-DH and Succ-DH—quickly from the ETC. We also notice that the transport of electrons to oxygen is mainly catalyzed via the $cbb_3$ oxidase as the ubiquinol oxidase is not expressed under strong aerobic conditions (Swem and Bauer, 2002) (see Figure 3).

To relate the results of the kinetic model to the EM analysis (Table I; Figure 2), we also computed for each scenario the relative contribution of the nine EMs (Figure 4; see Materials and methods section). Regarding the aerobic regimen, we can summarize that the ETC is mainly driven by the two elementary modes EM6 and EM8 and the relative proportion of EM8 (which uses the NADH-DH) increases with $R_{\text{NADH}}$ (Figure 4). Neither furmate reduction (EM9) nor reverse electron transport (EM3 and EM4) takes place; these modes become only active if $R_{\text{NADH}}$ is extremely high (and/or $R_{\text{Succ}}$ extremely low) or vice versa, respectively.

If we consider identical $R_{\text{NADH}}$ values for all four growth regimens, aerobic growth results in the highest $\Delta p$. However, this condition is not necessarily valid for the real physiological situation. Since experimental data show that $R_{\text{NADH}}$ is usually low under aerobic and higher under anaerobic growth (Table II), we might nevertheless observe a higher $\Delta p$ for photosynthetic conditions. Importantly, and this applies for all growth regimens, the $\Delta p$ values that were computed in the

| Table II | Four growth regimens simulated with the ETC model together with published measurements of model variables used for parameter fitting |
|----------|------------------------------------------------------------------------------------------|
|          | R1: aerobic growth in the dark | R2: semi-aerobic growth in the dark | R3: anaerobic growth in high light | R4: anaerobic growth in low light |
| Oxygen tension (%) | 9.9 | 0.1 | 440 (95.6) (Holt and Marr, 1965) | 7.3 (1.59) (Holt and Marr, 1965) |
| Mean irradiance, μE/s m² (W/m²) | 0 | 0 | 1.5 | 1.5 |
| $R_{\text{Succ}}$: [Succinate]/[Fumarate] | 1a | 1a | 1a | 1a |
| $R_{\text{ATP}}$: [ATP]/[ADP] | 2.7a (Schön, 1969) | 2.7a (Schön, 1969) | 2.7a (Schön, 1969) | 2.7a (Schön, 1969) |
| $R_{\text{NADH}}$: [NADH]/[NAD] (Schön, 1971) | 0.05a (Schön, 1971) | 0.71 (Schön, 1971) | 0.21 (Schön, 1971) | 0.71a (Schön, 1971) |
| Measurements (in parentheses: name of the corresponding model variable (see Table III)) | | | Q_{\text{total}} (Qt) | 1.31 (Carr and Exell, 1965) | 2.52 (Carr and Exell, 1965) |
| | | | $QH_2/Q_{\text{total}}$ ($Q_{\text{charge}}$) | 0.17b (Zannoni, 1995) | 0.53 (Redfarn, 1967) |
| | | | Bacteriochlorophyll (Bchl) | 0.05 (Grammel et al, 2003) | 1.22 (Grammel et al, 2003) |
| | | | | 2.17 (Holt and Marr, 1965) | 8.20 (Holt and Marr, 1965) |
| | | | $cbb_3$-Oxidase ($Oxi_t$) | 1.5c (Swem and Bauer, 2002) | 4.5c (Swem and Bauer, 2002) |
| | | | Ubiquinol oxidase ($UbiquinOxi_t$) | 0.5c (Swem and Bauer, 2002) | 1.5c (Swem and Bauer, 2002) |
| | | | μ (muc) | 0.154 (Grammel et al, 2003) | 0.10 (Holt and Marr, 1965) |
| | | | | 0.14 (Holt and Marr, 1965) | 0.10 (Holt and Marr, 1965) |

For oxygen tension and mean irradiance (sun light), typical values have been chosen (unit ‘E’ (Einstein) denotes 1 mol photons and 1 μE/s m² = 0.217 W/m²). References for measurements are given. Measurement units are μmol/gDW (for BChl and $Q_{\text{total}}$), h⁻¹ (for μ) and dimensionless for the others (see also Table III).

a Estimated values (see explanations in main text).

b In reference Zannoni (1995), the reduction degree of ubiquinone for aerobic conditions was estimated between 4 and 30%; here we chose the average.

The values for the relative concentrations of $cbb_3$ and ubiquinol oxidases have been estimated from expression ratios given in Swem and Bauer (2002).
A model of a bacterial electron transport chain
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Table III  State (S) and algebraic (A) variables of the electron transport chain model

| Name (unit)               | Type | Differential/algebraic equation                                                                 | Remarks                                                                 |
|---------------------------|------|--------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------|
| Qt (μmol/gDW)             | S    | Qt' = Ubiostich · c2_synth – mue · Qt                                                         | Total concentration of ubiquinone                                       |
| QH2 (μmol/gDW)            | S    | QH2 = 1000 · (NADH_DH_flux – bc1_flux + RC_flux + Succ_DH_flux) – mue · QH2                   | Concentration of reduced ubiquinone (ubiquinol)                         |
| Q (μmol/gDW)              | A    | Q = Qt – QH2                                                                                   | Free oxidized ubiquinone                                                |
| Q_charge (−)              | A    | Q_charge = QH2/Qt                                                                              | Reduction degree of ubiquinone                                          |
| c2t (μmol/gDW)            | S    | c2t = c2_synth – mue · c2t                                                                   | dpsi = psi_p                                                             |
| c2red (μmol/gDW)          | S    | c2red = 1000 · (2 · bc1_flux – Oxi_flux – 2 · RC_flux) – mue · c2red                           | Total concentration of cytochrome c2                                   |
| c2ox (μmol/gDW)           | A    | c2ox = c2t – c2red                                                                              | Concentration of oxidized c2                                          |
| c2_charge (−)             | A    | c2_charge = c2red                                                                              | Reduction degree of c2                                                  |
| c2_synth (μmol/gDW h)     | A    | c2_synth = K1 · in(h(pO2,Km,E2),ln(h(pO2,Km,E3)) + K1c · mue)                                 | Synthesis rate of c2 (Q and bc1) are synthesized in a fixed ratio       |
| Oxi_t (−)                 | S    | Oxi_t = K3 · hill(pO2,Km,E3) · in(h(pO2,Km,E3))                                                | (Relative) concentration of cbb3 oxidase reduced in cytochrome b3      |
| Oxi_red (−)               | A    | Oxi_red = Oxi_t/(Q/QH2 – exp((pmf + dpsi – Em_b3 + Em_c2)/ZZ) + 1)                            | (Relative) concentration of ubiquinone oxidase reduced in b3-type cytochrome; analogous assumptions as for Oxi_red |
| Oxi_flux (mmol/ gDW h)    | A    | Oxi_flux = K · Oxi_red · c2red · hill(pO2,Km,Oxi,Exp_Oxi)                                    | Flux through cbb oxidase                                               |
| UbOxi_t (−)               | S    | UbOxi_t = K4c · mue + K4 · hill(pO2,Km4,E3) · in(h(pO2,Km4,E3) – mue · UbOxi_t)               | Relative concentration of ubiquinone oxidase                            |
| UbOxi_red (−)             | A    | UbOxi_red = UbOxi_t/(Q/QH2 – exp((pmf – Em_b_UbiOxi + Em_Q)/ZZ) + 1)                          | (Relative) concentration of ubiquinone oxidase reduced in b3-type cytochrome; analogous assumptions as for Oxi_red |
| UbOxi_flux (mmol/ gDW h)  | A    | UbOxi_flux = K · UbOxi_red · QH2 · hill(pO2,Km_UbiOxi,Exp_UbiOxi)                             | Flux through ubiquinone oxidase                                         |
| bc1 (μmol/gDW)            | S    | bc1 = bc1_synch · c2_synth – mue · bc1                                                       | Concentration of bc1 complexes (monomers, not dimers)                  |
| bc1_flux (mmol/ gDW h)    | A    | bc1_flux = K · bc1 · (Em_c2 + ZZ · ln(c2ox/c2red) – (Em_Q + ZZ/2 · ln(Q/QH2))/2 · pmf + dpsi) | Flux through bc1 complex (see Materials and methods section)            |
| LHC (μmol/gDW)            | S    | LHC = PSC_synth − mue − LHC                                                                     | Light-harvesting complex (here only LH1 considered; see text)           |
| RC (μmol/gDW)             | S    | RC = PSC_synth – mue · RC                                                                       | Concentration of reaction centers                                      |
| Bchl (μmol/gDW)           | A    | Bchl = 4 · RC + 32 · LHC                                                                       | Bchl concentration (R. rubrum)                                          |
| PSC_synth (μmol/ gDW h)   | A    | PSC_synth = K2 · in(h(pO2,Km2,E2) · hill(Q_charge,Km2,E2) · in(h(Rc,Km2,E2))                 | Synthesis rate of photosynthetic complexes; with saturation term        |
| light_harvesting (me/gDW h) | A    | light_harvesting = K · LHC · hill(mean_irradiance,LH Lightint, Exp_lightint)                  | Photon absorption rate in LHC                                           |
| RC_flux (mmol/gDW h)      | A    | RC_flux = K · RC · hill(light_harvesting(Rc,Km,Lh,Exp,Lh))                                   |                                                                         |
| NADH_DH_flux (mmol/gDW h) | A    | NADH_DH_flux = K · NADHDH · (Em_Q + ZZ/2 · ln(Q/QH2) – (320 · ZZ/2 · ln(Q/QH2))/2 · pmf)      | Flux through NADH-DH                                                   |
| Succ_DH_flux (mmol/gDW h) | A    | Succ_DH_flux = K · SuccDH · (Em_Q + ZZ/2 · ln(Q/QH2))/2 – (31 · ZZ/2 · ln(1/Rsucci))        | Flux through Succ-DH                                                   |
| ATPSynthase_flux (mmol/gDW h) | A    | ATPSynthase_flux = K · ATPS · (exp(kk/(RR – TT)) – 1)) / (exp(kk/(RR – TT)) + 1) with kk = (3 · pmf – FF – 30.5 · RR – TT · ln(Ratp/Phos)) | Flux through ATP synthase                                             |
| mue (h−1)                 | S    | mue = mue_max · hill(AtPSynthase_flux,Km_flux,Exp_mue)                                        |                                                                         |
| Hp (mM)                   | S    | Hp = pV · (4 · NADH_DH_flux + 4 · bc1_flux + 2 · Oxi_flux + 2 · UbOxi_flux – 3 · ATPSynthase_flux) – proton_leak – mue · Hp | Growth rate; here: ATP-limited Proton concentration in the periplasm    |
| pHp (−)                   | A    | pHp = log10(Hp/1000)                                                                          | pH in periplasm                                                        |
| DeltaPH (−)               | A    | DeltaPH = pHp − pHp                                                                           | Δp (proton-motive potential)                                            |
| pmf (mV)                  | A    | pmf = 1/(1 – psi_p · 2.303 · ZZ · DeltaPH)                                                    | ΔΨ (membrane potential)                                                |
| dpsi (mV)                 | A    | dpsi = psi_p · pmf                                                                            | Expression taken from Korzeniewski (1996a)                             |
| proton_leak (mM/h)        | A    | proton_leak = K · leak · (pmf – 160)                                                          |                                                                         |

Parameters are written in italic. hill expresses standard Hill kinetics: hill(S,K,c) = S/S′ + (K′ + K). Furthermore, exp(s) = e^s, ln is the natural and log10 the decimal logarithm. gDW stands for ‘gram dry weight.’ Several equations are derived or explained in the Materials and methods section.
### Table IV Parameters of the electron transport chain model

| Symbol             | Value                      | Remarks                                                                 |
|--------------------|----------------------------|-------------------------------------------------------------------------|
| **mean_irradiance**| see Table II               | In \( \mu \text{E}/\text{s m}^2 \); value depends on chosen growth regimen (Table II) |
| \( pO_2 \)         | see Table II               | Oxygen tension (% saturation); value depends on chosen growth regimen (Table II) |
| \( \text{Rnad} \)  | see Table II               | \( \text{[NADH]/[NAD]} \); value depends on chosen growth regimen (Table II) |
| \( \text{Ratp} \)  | see Table II               | \( \text{[ATP]/[ADP]} \); value depends on chosen growth regimen (Table II) |
| \( \text{Rsucc} \) | see Table II               | \( \text{[Succinate]/[Fumarate]} \); value depends on chosen growth regimen (Table II) |
| \( FF \)           | 0.0965 KJ/mol mV           | Faraday constant                                                        |
| \( RR \)           | 0.0083 KJ/mol K            | Gas constant                                                            |
| \( TT \)           | 298 K                      | Temperature                                                             |
| \( ZZ \)           | 25.631 mV                  | \( ZZ = RR \cdot TT/FF \)                                             |
| \( E_{mQ} \)       | 70 mV                      | Midpoint redox potential of ubiquinone                                  |
| \( E_{mC_2} \)     | 340 mV                     | Midpoint redox potential of cytochrome \( c_2 \) (Garcia et al., 1987)   |
| \( E_{mB,UbiOxi} \)| 250 mV                     | Midpoint redox potential of cytochrome \( b \) in ubiquinol oxidase (Varela and Ramirez, 1990) |
| \( E_{mB_3} \)     | 540 mV                     | Midpoint redox potential of cytochrome \( b_3 \) in \( cbb_3 \) oxidase; analogous to Korzeniewski (1996b) |
| \( pHi \)          | 7                          | Cytosolic pH                                                            |
| \( \text{Phos} \)  | 0.001 M                    | Cytosolic phosphate concentration                                      |
| \( \text{psi}_p \) | 0.7                        | Relative contribution of \( \Delta \Psi \) (membrane potential) to \( \Delta \phi \) (estimated from Wraight et al. (1978); \( \Delta \phi = 1 \rightarrow \Delta \Psi/\Delta \phi = 0.7 \)) |
| \( pIV \)          | 10 mM/mmol/gDW             | Expresses the change of periplasmic proton concentration (mM) if 1 mmol cytosolic protons per gDW is pumped over the membrane; this value depends on buffer capacities and volume ratios (estimated; but steady-state results are insensitive against this parameter) |
| \( \text{mue}_\text{max} \) | 0.17 h\(^{-1}\)        | Maximal growth rate \( \text{(for } R. \text{rubrum); from own experiments; see also Grammel et al. (2003) and Holt and Marr (1965)} \) |
| \( E_{\text{pH}} \) | 7                          | Half the rate of ATP synthase if the cell grows with maximal growth rate \( (0.17 \text{ h}^{-1}) \); estimated from a stoichiometric model (Klamt et al., 2002) for photosynthetic growth on succinate (a similar value follows for respiratory growth) |
| \( K_m_{\text{mue}} \) | 8.5 mmol/gDW h            | Approximated concentration ratio \( Qt/c_2 \) (Garcia et al., 1987) |
| \( K_{\text{Ubistoich}} \) | 25                      | Approximated concentration ratio \( bc1/c_2 \) (Garcia et al., 1987) |
| \( K_{\text{bc1stoich}} \) | 0.6                      | Approximated concentration ratio \( bc1/c_2 \) (Garcia et al., 1987) |
| \( K_{\text{ATPS}} \) | 20.0 mmol/gDW h           | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{NADHDH}} \) | 3.35 mmol/gDW h mV      | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{SucchDHDH}} \) | 0.02 mmol/gDW h mV      | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{leak}} \) | 0.08 mmol/h mV            | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |
| \( K_{\text{Oxi}} \) | 13000 mmol/h \( \mu \text{mol} \) | Rate constant for photon absorption by LHCs \( K_m \) value for light absorption; we assume light saturation at 50 W/m\(^2\) (Geyer and Helms, 2006), i.e. \( K_m = 25 \text{ W/m}^2 = 115 \mu\text{E}/\text{s m}^2 \) |

Parameter values without references or explanations have been estimated by fitting the measurements given in Table II.
model simulations are in a physiologically meaningful range (180–200 mV), even though this has not been taken as a constraint for parameter fitting. This underlines the validity of our model. The model also reproduces correctly that the synthesis of the photosynthetic apparatus and pigments is strongly repressed because neither the oxygen concentration is low nor the ETC is in a reduced state.

Semi-aerobic growth (regimen R2)
The semi-aerobic state is comparably only loosely defined, due to experimental difficulties as well as species-dependent differences. For *R. rubrum*, Grammel et al (2003) demonstrated that at oxygen tensions below 0.5%, ICM expression is induced and is accompanied with a specific enzymatic profile.

The simulation of this growth regimen shows that when oxygen becomes limiting, the electron flow to oxygen via the cytochrome oxidase is decelerated. As a consequence, ubiquinone and $c_2$ are reduced stronger compared to the aerobic case (Figure 3). This effect certainly contributes to higher $R_{NADH}$ values seen under semi-aerobic conditions (see Table II). The regulatory machinery increases the synthesis of the $cbb_3$ cytochrome oxidase and of the ubiquinol oxidase. Since the

![Figure 3](image-url)
electron flow to the final acceptor oxygen now occurs via both oxidases, the elementary modes EM5 and EM7 are now acting concurrently with EM6 and EM8 (Figure 4) thus avoiding an over-reduction of the ETC, though at the expense of lowering the P/O ratio and therefore ∆p. With ascending cytosolic redox charge $R_{\text{NADH}}$, the ubiquinol oxidase carries an increasing number of electrons to oxygen because QH$_2$ becomes available in high amounts. As for regimen R1, reverse electron transport can only be observed when choosing extreme values for $R_{\text{NADH}}$ and/or $R_{\text{Succ}}$. Furthermore, since $R_{\text{NADH}}$ is higher under semi-aerobic conditions, the input of electrons into the ETC occurs now with a higher proportion via the NADH-DH (EM7 and EM8). In contrast, the flux through Succ-DH (EM6) is decreased being in agreement with the fact that the TCA flux in *R. rubrum* is reduced under semi-aerobic and anaerobic conditions (Grammel et al., 2003).

In addition, as shown in Figure 3, the induction of ICM synthesis under oxygen-limiting conditions in the dark is correctly reproduced by the model: a considerable amount of BChl and photosynthetic complexes is synthesized, matching the published data.

Anaerobic growth under high light (regimen R3)

When oxygen supply is switched off and light is provided (here with high intensity), a complete reorganization of the composition of the ETC and of the electron flows take place. The repressing signal of oxygen has completely disappeared and photosynthetic membranes and complexes are now synthesized with a higher rate compared to the semi-aerobic case, whereas the expression of oxidases is repressed to a constitutive level. Cyclic photosynthesis (EM1) is now the dominating operational mode in the ETC generating ATP constantly without consumption of electrons. Therefore, electrons fed into the ETC via Succ-DH must leave the ETC again for avoiding over-reduction. In fact, the reaction rate of NADH-DH is now negative and the overall stoichiometry together with the rate of Succ-DH corresponds to the ∆p-driven reverse electron transport expressed by EM2 (Figure 4). One might argue that an input of electrons into the ETC and a subsequent energetically expensive reverse flow should be avoided by the cell; however, there are situations where the Succ-DH is necessarily in use, for example, if succinate is the substrate. Note also that a higher reduction level of NADH increases the electron pressure toward the ETC, repressing the reverse flow of electrons from Succ-DH to NADH-DH. For very large $R_{\text{NADH}}$ values (beyond the range considered here), reverse electron flow (EM2) would be replaced by fumarate reduction (EM9).

The reduction degree of ubiquinone and $c_1$ is significantly larger than in the aerobic regimen R1, even if we would consider the same level for $R_{\text{NADH}}$ (Table II shows that $R_{\text{NADH}}$ tends to be higher under photosynthetic conditions). This is not surprising since oxygen is not available as an electron sink. However, the ETC is less reduced compared to the semi-aerobic regimen (although the validity of this relation strongly depends on which oxygen tension is considered to be ‘semi-aerobic’). Growing $R_{\text{NADH}}$ values increase the pressure of electrons into the ETC and the redox state of the ubiquinone (and accordingly the expression of photosynthetic genes) is therefore monotonically dependent on $R_{\text{NADH}}$.

In contrast, another result of our simulations shows that there is a maximum in ∆p for photosynthetic conditions: when low $R_{\text{NADH}}$ values are increased, ∆p goes up as well because the ETC gets filled with electrons required for driving the photosynthetic cycle. Then, $R_{\text{NADH}}$ reaches an optimum value above which ∆p decreases as now the ETC becomes saturated.

![Figure 4](image-url) Contribution of the nine elementary modes (Figure 2 and Table I) in the steady-state operation of the kinetic model. As in Figure 3, a response curve for a range of NADH/NAD ratios ($R_{\text{NADH}}$) for the four different growth regimens (Table II) were considered. (For details regarding the computation of these contributions, see Materials and methods.)
increasingly over-reduced, impeding the recruitment of Q by the RC. Furthermore, as already mentioned above, $\Delta p$ under photosynthetic conditions does not reach the high level of the aerobic regimen R1, when identical values for $R_{\text{NADH}}$ are considered in both cases. However, one might choose different $R_{\text{NADH}}$ values (small for R1 and larger for R3, e.g. 0.03 and 0.9, respectively) and a higher $\Delta p$ would result for photosynthetic growth. Hence, no general statement can be made whether photosynthetic or respiratory growth generates a larger $\Delta p$.

**Anaerobic growth under low light (regimen R4)**

Under low-light conditions, photons become the limiting resource and the overall rate of photosynthesis (see variable $\text{RC_flux}$ in Figure 3) decreases. The bacteria react upon low light with an increased synthesis of photosynthetic membranes and complexes that, however, can only partially compensate for the reduced photosynthetic flux. The driving force for the amplified expression of the photosynthetic gene cluster can be attributed to the significantly increased reduction degree of ubiquinone. So far, no direct experimental evidence has been given that $\text{QH}_2/Q_{\text{total}}$ increases when diminishing the light intensity. It is even not clear by intuition: a lowered influx of photons implies that the light-induced transfer of electrons from $c_2$ to Q is decelerated that could lead to an accumulation of reduced $c_2$ (which is indeed the case) but not necessarily of $\text{QH}_2$. Our model gives not only a prediction it also gives an explanation for this behavior: due to the lowered photosynthesis rate, the $\Delta p$ value drops down to about 181 mV. As an implication, the resistance of the ETC against the influx of electrons from NADH via NADH-DH decreases as well (see equation (12) in Materials and methods) leading eventually to a higher reduction degree of ubiquinone. The effect is amplified by the fact that the cytosolic redox state seems to be reduced stronger under low-light conditions.

Importantly, the qualitative behavior of the ubiquinone pool displayed is independent of any model assumption made on the regulation, that is it is a direct effect of the reduced light intensity (and not an indirect effect induced by the activity of regulatory pathways). One might also argue that our result is a matter of the chosen parameter set. Using an evolutionary optimization algorithm (Rechenberg, 1984; Rettinger, 1995), we therefore tried to find a parameter set that brings about the opposite result, that is: that the ubiquinone pool is more oxidized under low-light conditions. Even when running the algorithm over days, we could not identify such a parameterization indicating that our findings are indeed a qualitative and structural property of the system.

The fact that the ubiquinone pool gets more reduced under low-light conditions has strong implications: it provides the mechanistic basis for employing the redox state of ubiquinone as an integrated signal for sensing oxygen availability and light intensity that can be used to trigger regulatory events. Our model thus supports the hypothesis that the ubiquinone redox state could be involved in controlling the expression of photosynthetic genes (Oh and Kaplan, 2001; Swem et al., 2006). However, one should also be aware of another important aspect when considering the specific amount of BChl and photosynthetic complexes. The growth rate (in our model coupled with the rate of ATP synthesis) usually decreases under limiting low-light conditions implying also a lowered growth-related dilution of photosynthetic components. This may have a profound effect on specific concentrations of photosynthetic components (the latter are not actively degraded): having a cell culture A whose specific content of BChl is twice that of a cell culture B but the growth rate of A is half that of B, then the specific pigment synthesis rate of both cultures is the same. Hence, although the expression rate of photosynthetic genes is certainly increased, a higher specific concentration is partially achieved by the reduced growth rate.

Similar as for high-light conditions, there exists a maximum $\Delta p$ when increasing $R_{\text{NADH}}$ values, though the optimum occurs at a lower $R_{\text{NADH}}$ level compared to regimen R3 (Figure 3).

A further observation we can make is that for larger $R_{\text{NADH}}$ values (see Figure 4; $R_{\text{NADH}}>1.95$), the reverse electron transport (EM2) stops and is then replaced by fumarate reduction (EM9), which can be seen for high-light conditions only for even larger $R_{\text{NADH}}$ values.

**Model simulations: varying the cellular $[\text{ATP}]_{\text{p}}/[\text{ADP}]$ ratio**

As we did in the last section for $R_{\text{NADH}}$, we also varied $R_{\text{ATP}}$ in a physiologically meaningful range from 1.5 to 3.5 (and fixed $R_{\text{NADH}}$ and $R_{\text{Succ}}$ to the values given in Table II). As the main result, increasing $R_{\text{ATP}}$ and keeping the other two ratios constant implies a higher $\Delta p$ for the growth regimens R1–R3: the rate of ATP synthesis is decreased because of the high concentration of ATP. As a consequence, the ubiquinone and $c_2$ pools get more oxidized because the (forward) flows through NADH-DH, $bc_1$ and the oxidases are reduced as these complexes pump protons against $\Delta p$. These effects establish a negative feedback (also known from respiratory control): a high ATP concentration reduces the proton pumping and thus ATP-producing activities in the ETC. Unexpectedly, this behavior appears to be different for photosynthetic growth under low-light conditions (R4): as for the other regimens, $\Delta p$ increases with $R_{\text{ATP}}$ and ubiquinone and $c_2$ are less reduced; however, the rate of the cyclic photosynthesis increases as well further contributing to a higher $\Delta p$, leading eventually to a higher rate of ATP synthesis. The reason is that larger $R_{\text{ATP}}$ values shift the ETC to a more oxidized state (as reverse electron flow is facilitated by a higher $\Delta p$) in which the photosynthetic cycle can be driven more efficiently (remember that the ETC is over-reduced under low-light conditions).

**Model simulations: varying the cellular $[\text{succinate}]/[\text{fumarate}]$ ratio**

Finally, we varied $R_{\text{Succ}}$ over two orders of magnitude from 0.1 to 10 and kept $R_{\text{NADH}}$ and $R_{\text{ATP}}$ constant. It turned out that, under all growth regimens, most steady-state values exhibit a remarkable insensitivity against $R_{\text{Succ}}$. This concerns in particular the reduction degrees of ubiquinone and $c_2$, which remain almost constant over the full range of simulated $R_{\text{Succ}}$. Clearly, a strong change occurs in electron fluxes carried by NADH-DH and Succ-DH, respectively: under aerobic conditions, increasing $R_{\text{Succ}}$ enhances the electron influx into the ETC via Succ-DH (EM5 and EM6 contribute stronger), whereas
the influx of electrons via NADH-DH is thereby lowered (reduced activity for EM7 and EM8). Analogously, under photosynthetic conditions, reverse electron flow takes place with a higher rate due to a higher influx of electrons from succinate.

We also observed that $\Delta p$ is slightly decreased with growing $R_{\text{Succ}}$. This can be explained by the fact, that the Succ-DH does not contribute to the generation of $\Delta p$ (in contrast to NADH-DH whose forward operation is reduced).

**Model simulations: aerobic growth in light**

We also tested how the model behaves if oxygen and light are available concurrently. The simulation results for this situation should be interpreted with caution because (i) phototoxic effects induced by oxygen radicals, which may occur under these conditions and inhibit growth, are not considered by our model and (ii) modeling the modulation of respiration by light and vice versa might require a compartmentalized model with cytoplasmic and intracytoplasmic membranes. However, we mention here that our model behaves qualitatively correct, that is the synthesis of the photosynthetic apparatus is repressed at high oxygen tension irrespective of the availability of light. Photosynthesis may interfere with respiration only when a photosynthetically grown culture is exposed to oxygen; the repression of the photosynthetic genes will abolish this effect when the photosynthetic units are diluted out. In the steady state, respiration is the dominating operational mode and the ETC behaves virtually as in the aerobic dark case.

**Parameter sensitivities and model robustness**

As an important step in quantitative modeling, we discuss the robustness of our in silico results with respect to parameter uncertainties. As mentioned, several parameters are well defined from literature values (e.g. midpoint redox potentials of ETC components) and the rest has been estimated by fitting the measurements given in Table II. We cannot exclude that other parameter sets fit the measurements also reasonably well. However, albeit we examined quantitative (steady state) results from model simulations, most conclusions drawn from the kinetic model are mainly qualitative (e.g. ‘ups and downs’) and are thus per se robust against parameter variations as long as none of the parameters is close to a bifurcation point.

For a rough assessment of the parameter sensitivities, we perturbed each estimated parameter $p_j$ (except exponents in kinetic equations) by a relative level of 10% and analyzed how the model variables $x_i$ (we considered state and algebraic variables) change their steady-state value $x_i^{ss}$. As a main result, only in about 6% of all cases the model variables changed more than 10% (i.e. only in those cases the approximated relative parameter sensitivities

$$\frac{\Delta x_i^{ss}}{x_i^{ss}}/\Delta p_j/p_j$$

are larger than unity) and thus the model shows a reasonable robustness. Parameters exhibiting larger sensitivities are those involved in the synthesis of ETC components, in particular parameters controlling the synthesis of the photosynthetic apparatus. In addition, several model variables were sensitive with respect to the parameter $K_{\text{ATP}}$, the rate constant of ATP synthase. This is not surprising since the rate of ATP synthase has a strong effect on $\Delta p$ which in turn influences the thermodynamic span and thus the rate of several other reactions in the ETC. Finally, as expected, for semi-aerobic growth several variables show a higher sensitivity for the $K_m$ values (oxygen saturation) of the two oxidases.

In contrast, very low sensitivities for the steady states can be seen for the two parameters describing properties of the membrane compartment, namely $K_{\text{leak}}$ (quantifying the strength of proton leak) and $pV$ (dependent on buffer capacity and periplasmic volume; see Materials and methods). Most likely, these parameters will have different values for the two membrane types CM and ICM. However, the low sensitivities justify that we have considered identical values for respiratory and photosynthetic growth.

Importantly, the two model variables $\text{Ubi}_{\text{charge}}$ and $c_2_{\text{charge}}$, which reflect the overall redox state of the ETC and which have been discussed intensively in the previous sections, exhibit generally a comparably low sensitivity against parameter variations.

**Discussion**

In this work, we have presented and analyzed a stoichiometric and kinetic model for the photosynthetic and respiratory operation of the ETC in purple non-sulfur bacteria. Although the stoichiometric model is relatively small (seven reactions) and does not consider any kinetics, it helps to identify the potential functional behaviors of the ETC under photosynthetic, respiratory and fermentative conditions. Apart from the well-known cycles and pathways, the model revealed two operational modes representing reverse electron flow under respiratory conditions. So far, the functional role of reverse electron flow has been mainly discussed for photosynthetic growth only. In fact, due to thermodynamic constraints, reverse electron flow under aerobic conditions plays only a minor role; however, it might become important for certain (possibly temporary) metabolic states, for example, if a great imbalance in the current redox potentials of NADH and succinate occurs. The stoichiometric ETC network model thus demonstrates how stoichiometric modeling helps in identifying—in an unbiased way—meaningful functions of a biochemical network.

Clearly, the stoichiometric model can only show the potential behavior of the ETC. To study the actual driving forces, electron flows and redox states occurring in the ETC under different environmental conditions, we constructed a kinetic model that also includes known regulatory pathways governing the expression of ETC components. To our knowledge, it is the most comprehensive model of the ETC in purple bacteria with respect to the components and processes considered. The reaction steps taking place inside the enzyme complexes have been described with relatively simple, yet thermodynamically justified kinetic laws. We do not claim that our model reproduces ETC states accurately when absolute quantitative units are considered, since available parameters and measurements do not permit this. Nevertheless, it
provides valuable semiquantitative and qualitative insights into the behavior of the ETC, in particular on the redox states of key components that are often difficult to measure. The most important prediction we could derive with our model is that the ubiquinone pool becomes reduced stronger during photosynthetic growth when diminishing the light intensity to a low (limiting) level. Only this behavior qualifies the ubiquinone redox state as a suitable signal for regulating the synthesis of photosynthetic genes as experimental studies have indicated (Oh and Kaplan, 2001; Swem et al., 2006).

We mention that light induction of photosynthetic gene expression has also been found to be dependent on photosynthetic electron flow in plant chloroplasts, although the present results about the regulatory role of the redox state of the plastoquinone pool in chloroplast gene expression are conflicting (see for example, Pfannschmidt et al., 1999; Matsuo and Obokata, 2002). We think that the modeling approach chosen here is, in principle, extendable by topological constraints for addressing also electron transport phenomena in plant cells, including transfer of reducing equivalents between different energy-converting organelles.

The redox state of ubiquinone is a highly integrative signal. According to our simulations, oxygen availability has the strongest influence, whereas the light intensity, the redox state of cytosolic metabolites exchanging electrons with the ubiquinone pool (such as NADH and succinate) and the ATP concentration do further modulate this signal. The advantage of using an integrated signal for regulation is that the cell requires only one ‘measurement device’ for sensing different environmental cues. However, combining different signals into one may also lead to some unexpected (for the cell possibly unfavorable) side effects. As we have shown previously for R. rubrum, the usage of a special C-substrate combination (fructose and succinate) instead of a standard (succinate) medium, significantly increases the expression of photosynthetic membranes and pigments under semi-aerobic conditions in the dark (Ghosh et al., 1994). We have suggested that growth with this substrate combination results in a stronger reduction of the ubiquinone pool, thus interfering with the signaling pathways controlling photosynthetic gene expression (Grammel et al., 2003). From our simulations, we can conclude that an increase of \( R_{\text{NADH}} \) and/or a decrease of \( R_{\text{ATP}} \) is most likely responsible for this behavior, whereas the reduction level of ubiquinone is relatively insensitive with respect to \( R_{\text{Succ}} \).

Another interesting result of our model is that for anaerobic photosynthetic growth, there is an optimal value of \( R_{\text{NADH}} \) resulting in the highest possible \( \Delta p \), and the optimal \( R_{\text{NADH}} \) value depends on the light intensity. Both a too low and a too high reduction degree of NADH limit cyclic photosynthesis because both oxidized and reduced forms of ubiquinone and \( c_2 \) are required for driving this cycle. This implies that there is an optimal electron load in the ETC. In contrast, such biphasic phenomena cannot be observed under aerobic conditions: there is a monotone dependency of \( \Delta p \) for arbitrarily high values of \( R_{\text{NADH}} \). Clearly, these considerations assume that the values for \( R_{\text{ATP}} \) and \( R_{\text{Succ}} \) are not directly coupled with \( R_{\text{NADH}} \).

Our kinetic ETC model in its present form provides a mathematical description of the membrane ETC as a first building block, which can be refined or extended in a modular way by additional blocks such as cytoplasmic redox events or a more detailed representation of the mechanisms of redox signaling and redox control of photosynthetic gene expression. The possible refinement of the model by introducing two membrane compartments has already been discussed above.

In the long run, mathematical models, such as the one described in this work, promise to enable a comprehensive theoretical investigation of the redox metabolism of purple bacteria at the systems level.

**Materials and methods**

**Kinetic modeling of electron flows in the ETC**

The kinetic ETC model comprises the processes depicted in Figure 1. For the kinetic description of the complicated electron transfer processes, we followed the approach of Korzeniewski (1996a, b) and Konrzeniewski and Mazat (1996) by modeling the entirety of elementary steps taking place at an enzyme complex as one (overall) redox reaction. The overall stoichiometries of these reactions correspond to the equations (1–6) and their driving forces are redox potential differences between the two redox couples involved.

The full model with all its (ordinary) differential equations, algebraic relationships and parameters is documented in Table III (model variables) and Table IV (parameters). In the following, we explain some selected equations.

A central variable of the ETC model is the proton-motive force (\( \Delta p \)). The well-known equation for \( \Delta p \) accounts for the electrical gradient (membrane potential \( \Delta W \)) and the concentration gradient (\( \Delta H^+ \)) between the protons in the periplasmic and the cytoplasmic space:

\[
\Delta p = \Delta W - 2.303 \cdot R \cdot T \cdot \text{pH} - P_{\text{H}} = \Delta W - 59mV \cdot \Delta p
\]

\( R \): gas constant; \( T=298K \); \( F=\text{Faraday constant}; \) \( \Delta W=\text{pH}_c-\text{pH}_p \), \( \Delta H^+ \) and \( \text{pH} \) are the pH values in the periplasmic and cytoplasmic space, respectively. Contrary to \( \Delta W \), the membrane potential \( \Delta W \) is not defined by the proton gradient alone, instead it is an integral value that reflects the distribution of all ions at both sides of the membranes. Though only approximately correct (Schuster et al., 1999), we follow Korzeniewski (1996a, b) in assuming that there is a linear relationship between \( \Delta p \) and \( \Delta W \), that is that we can approximate \( \Delta W \) from \( \Delta p \) via a constant factor, which could be derived from measurements of Wraight et al. (1978) and indicates that 30% of the \( \Delta p \) is contributed by \( \Delta W \) and 70% by \( \Delta H^+ \). We assume additionally that the cytosolic \( \text{pH} \) is constant (\( \text{pH}=7 \)). For bookkeeping the concentration of protons in the periplasmic space, we use a parameter \( \text{pH}^+ \) expressing which amount of \( H_+^+ \) in [mmol/gDW] must be pumped over the membrane to increase the concentration of \( H_+^+ \) by 1 mM. Clearly, the value of this parameter will strongly depend on many factors such as volume ratios and the buffer capacity of the periplasm, and can only be estimated. However, buffer capacities do not change the steady-state behavior studied herein (but may change the transient behavior). In fact, sensitivity analysis shows that the model results are virtually invariant against varying this parameter over orders of magnitudes. The model considers proton leak also accounting for the escape of protons from the periplasmic space.

As mentioned above, electron transfer processes in complexes such as NADH-DOH or \( \text{bc}_{1} \) are modeled as overall redox reactions. The driving force of redox reactions is the redox potential difference \( \Delta E \) between the two participating redox couples (electron donor (D) and acceptor (A)) which, according to the Nernst equation, can be quantified as

\[
\Delta E = \Delta E_m + \frac{RT}{2F} \ln \left( \frac{[D \text{red}]}{[D \text{ox}]} \right)
\]

\( \Delta E_m \) (or \( \Delta E^0 \)) is the difference in the standard redox (midpoint) potentials of the two redox couples A and D: \( \Delta E_m = E^0_{\text{red}} - E^0_{\text{ox}} \) and \( z \) is the number of transferred electrons. The redox potential difference is directly coupled with the Gibbs energy change of the redox reaction

\[
\Delta G = -zF \Delta E
\]
In accordance with Korzeniewski (1996a,b) and Korzeniewski and Mazat (1996), we assume that the reaction rate of the redox reaction is proportional to the thermodynamic span (hence proportional to $-\Delta G$) and the amount of available enzyme. $\Delta p$ and/or $\Delta \Psi$ must be additionally taken into account if protons are pumped and/or electrons are translocated vertically in the membrane. We exemplify the rate equations for the case of the NADH-DH and the bc$_1$ complex.

NADH-DH transfers two electrons from NADH to Q. For the redox potential difference, we obtain

$$\Delta_{\text{NADH-DH}} = \Delta_{\text{NADH/QH}_2} - \frac{RT}{2F} \ln \left( \frac{[Q]}{[\text{NADH}]} \right)$$

(11)

We know that $\Delta_{\text{NADH/QH}_2} = 390$ mV since $E_{\text{QH}_2}^0 = 70$ mV and $E_{\text{NADH}}^0 = -320$ mV. The concentrations of Q and QH$_2$ are state variables in the model. The ratio [NADH]/[NAD] is fixed prior to simulation (see below). With equation (11) we can quantify the thermodynamic span of the electron transfer from NADH to QH$_2$; however, we have to consider that the NADH-DH couples the electron transfer by pumping four protons against $\Delta p$ that reduces the driving force. We thus obtain the following expression for the reaction rate of NADH-DH

$$r_{\text{NADH-DH}} = -k_{\text{NADH-DH}} \Delta_{\text{NADH-DH}} - 4\Delta p$$

(12)

which can conveniently be written as

$$r_{\text{NADH-DH}} = k'_{\text{NADH-DH}} (\Delta_{\text{NADH-DH}} - 2\Delta p)$$

(12')

Since $\Delta_{\text{NADH/QH}_2}$ is relatively high, the NADH-DH proceeds usually in the forward direction. However, under certain conditions (high levels of $\Delta p$ and QH$_2$ and a low level of NADH) the reaction rate may also become negative. This would correspond to the reverse electron flow as discussed in the main text (EM2-EM4 in Table 1; Figure 2) where electrons flow from QH$_2$ to NAD under consumption of $\Delta p$.

To evaluate equation (11), the concentration ratio [NADH]/[NAD] is required. Since NADH and NAD are involved in many other metabolic reactions in the cell, it is practically impossible to consider NADH and NAD as state variables in our kinetic model. Therefore, we employ the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the same strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy: for each of the considered growth regimens—for example, aerobic growth in the dark or anaerobic growth in light—we follow the following strategy:

The rate equation for the bc$_1$ complex is similar as for NADH-DH:

$$r_{\text{bc}_1} = -k_{\text{bc}_1} [\text{bc}_1] \Delta_{\text{bc}_1}$$

$$= k_{bc_1} [\text{bc}_1] F(2\Delta E_{bc_1} - 4(2.303RT\Delta pH) - 2\Delta \Psi)$$

(13)

or

$$r_{\text{bc}_1} = k_{bc_1} [\text{bc}_1] (\Delta E_{bc_1} - 2(2.303RT\Delta pH) - 2\Delta \Psi)$$

(13')

There are, however, two differences (i) in contrast to NADH-DH, the concentration of bc$_1$ is considered explicitly since it is a state variable in the model and subject to regulatory events (see below). (ii) During one Q cycle (eq. (3)), two electrons are transported against the membrane potential (from Q$_{p}$ (Qo) to Q$_{n}$ site of bc$_1$), whereas four protons are transported against the concentration gradient $\Delta p$. We note that equation (13) is equivalent to the term

$$r_{\text{bc}_1} = k'_{\text{bc}_1} [\text{bc}_1] (\Delta E_{bc_1} - 2\Delta p + 2\Delta \Psi)$$

used in the mitochondrial models of Korzeniewski (1996a,b) and Beard (2005) since $\Delta p = 2(2.303RT\Delta pH + 2\Delta \Psi)$.

The rate equation for the Succ-DH is analogous to NADH-DH (eq. (12)), except that the negative term disappears because protons are not pumped over the membrane. The rate equations for the ubiquinol oxidase and cytochrome (cbb$_3$) oxidase have been modeled analogously to Korzeniewski (1996b) by assuming that the redox state of the b-type cytochromes in the oxides is in near equilibrium with ubiquinone and $c_2$, respectively (see Table III). Then, electrons flow from these intermediate states to oxygen with Michaelis–Menten kinetics.

For the ATP synthase, we adopt the kinetic approach of Korzeniewski (1996b), which ensures that the rate of ATP synthesis becomes saturated for large $\Delta p$ values while thermodynamics still being consistent

$$r_{\text{ATPS}} = \frac{e^{\beta/RT} - 1}{e^{\beta/RT} + 1}$$

(14)

with

$$\gamma = -\Delta G_{\text{ATPS}} = \Delta G_{\text{bc}_1} - RT \ln \left( \frac{[\text{ATP}]}{[\text{ADP}][\text{Pi}]} \right) - 3 \cdot \Delta \Psi$$

The $\Delta G_{\text{bc}_1}$ is the Gibbs energy change of ATP hydrolysis under biological standard conditions ($-30.5$ kJ/mol) and $\Delta \Psi$ is the proton electrochemical gradient ($\Delta \Psi = -F\Delta p$). The phosphate concentration $[\text{Pi}]$ is considered to be constant and the ratio [ATP]/[ADP] is clamped as described for [NADH]/[NAD].

Finally, we need to describe the key reaction steps taking place in the photosynthetic apparatus. The latter consists of LHC and RC (for a detailed description of these complexes, see Nicholls and Ferguson, 2002). There are two types of LHC known as LH1 and LH2. LH1 consists of $32$ BChl and $16$ carotenoid molecules and surrounds the RC, that is the LH1/RC ratio is 1. LH2 is located further away from the RC (but still close enough to facilitate energy transfer) and consists of $27$ BChl and $9$ carotenoid molecules. LH1 and LH2 are differently regulated in purple non-sulfur bacteria, thus providing an additional degree of freedom to adapt to varying light conditions (LH2 proportion increases under low-light conditions). As already mentioned, _R. rubrum_—used as model organism herein—does not possess antennae of type LH2. The advantage for modeling is that the total amount B of BChl can be directly assigned to LH1 ($32/36$ B) and RC ($4/36$ B). The LHC absorbs light energy and transfer it to the RC where the energy is used for charge separation processes.

A detailed description of the processes occurring in the RCs of purple non-sulfur bacteria can be found in Nicholls and Ferguson (2002). Here, we consider a thermodynamic description based on the overall reaction scheme according to equation (6). In this way, the photosynthetic reaction cycle is modeled at the same level of detail as the electron transfer processes described above while retaining thermodynamic consistency. The change in Gibbs energy associated with this reaction scheme is given by

$$\Delta G_{\text{RC}} = -\mu_p - 2(\Delta E_{\text{RC}} - \Delta \Psi)$$

where we have taken into account that two electrons are transferred against the membrane potential (Nicholls and Ferguson, 2002) and $\mu_p = \mu_v \cdot (1 - T_j/T_r)$ is the Gibbs energy change associated with photon absorption (Meszna and Westerhoff, 1999). Here, $\mu_v$ and $h$ denote Avogadro’s and Planck’s constant, respectively, while $T$ corresponds to the environmental temperature of the RCs and $T_r$ is the temperature of the radiation field, for example, the surface temperature of the sun $T_r$=5000 K. The absorption frequency $v$ for purple non-sulfur bacteria corresponds to a wavelength of approximately 870 nm. For the reaction rate of the photosynthetic cycle, we use an adaption of the expression given in Meszna and Westerhoff (1999)

$$r_{\text{RC}} = k_{\text{RC}} I(e^{\alpha h/RT}e^{(\beta/RT)(|c_2|^2 + |QH_2|^2)} - 2|QH_2|^2 |c_2|^2)$$

(15)

with $x = \frac{\alpha}{h} (\Delta G_{\text{bc}_1} - \Delta E_{\text{bc}_1}/2) + \Delta \Psi$ and where $l$ depends on the number of photons collected (which in turn depends on the number of LHCs) and on the number of RCs (for a detailed expression, see Table III). Equation (15) may be rewritten as

$$r_{\text{RC}} = k_{\text{RC}} I(e^{-2\beta/RT}(|c_2|^2 + |QH_2|^2) - e^{-\beta/RT}|QH_2|^2 |c_2|^2)$$

(15')
Modeling regulatory pathways

The composition of the ETC and the formation of photosynthetic membranes in purple bacteria are mainly governed by the external signals oxygen and, to a lower extent, light intensity and the supply of reducing equivalents by the carbon substrate (Oh and Kaplan, 2001; Bauer, 2004). Although our model focuses on the operation of the ETC, we need to include basic regulatory events such that the photosynthetic apparatus consisting of LHC and RC is synthesized under anaerobic conditions. The molecular basis of the regulation of the photosynthetic gene cluster containing the genes for BChl (bch) and carotenoid (crt) biosynthesis, and for LHC polypeptides (puc and puf) and the RC subunits (puf and pufj) (Alberti et al, 1995), has been studied in detail in Rhodobacter capsulatus and Rhodobacter sphaeroides (Oh and Kaplan, 2001; Bauer, 2004) and appears to be conserved, including in R. rubrum (Addlessee and Hunter, 2002). Three major regulators have been identified that control the expression of genes in the PGC and those involved in the synthesis of other ETC components such as ubiquinone, cytochrome cbb3, as well as of the ubiquinol and the cytochrome (cbb3) oxidase.

(i) The RegB/RegA (PrrB/PrrA) two-component system with the global regulator RegA activates the expression of genes involved in respiration (e.g. oxidases), photosynthesis (PGC), CO2 fixation, nitrogen fixation and hydrogen release (Swem et al, 2001; Elsen et al, 2004). It is still under debate what the initial signal for auto phosphorylation of RegB is. The cbb3 oxidase has been demonstrated to generate an inhibiting signal for RegB autophosphorylation (Oh and Kaplan, 2000). Recent works indicate that RegB activity is triggered by the redox state of the ubiquinone pool (Swem et al, 2006), similar to the mechanism of the global ArcB sensor kinase in Escherichia coli (Georgellis et al, 2001). Under reducing conditions, the RegB autophosphorylase and subsequently phosphorylates and thereby activates the transcription activator RegA.

(ii) Another regulator is FnrL (Fnr-like protein), a homolog of the global regulator Fnr known in many other bacteria, including E. coli. Under anaerobic conditions, with an unknown mechanism, FnrL dimerizes to its active form and enhances, among others, the expression of cytochrome c, cbb3, and some photosynthetic genes including those required for LH2 formation (Zeilstra-Rylls et al, 1997; Zeilstra-Rylls and Kaplan, 1998; Oh and Kaplan, 2001).

(iii) A repressor of photosynthetic genes is PpsR (CrtJ). Under aerobic conditions, PpsR binds to its target promoters in the PGC, thereby blocking their expression (Elsen et al, 2005; Kovacs et al, 2005). Regarding the triggered signal, it was considered that PpsR might have the capability of sensing redox changes through the formation of disulfide bond(s) (Masuda and Bauer, 2002), but it remains still unknown what the actual input signal is. PpsR signaling becomes more complicated by the fact that AppA (known in Rh. capsulatus, not in Rh. sphaeroides) behaves as an antirepressor of PpsR. AppA itself is influenced by redox signals and may reduce its antirepressor activity when exposed to blue light (Masuda and Bauer, 2002).

In the differential equations for LHC, RC and other ETC components, we included kinetic terms expressing the dependency of the synthesis of the ETC components on the oxygen tension and the redox state of the ubiquinone pool mimicking the combined effects of the regulators AppA/PpsR, FnrL and RegB/RegA (compare with Table III):

- Expression of LHC and RC depends negatively on oxygen and positively on the reduction degree of ubiquinone. Note that the latter is rather a hypothesis than an ascertained fact and our model aims at verifying whether the reduction degree of ubiquinone is a suitable signal for regulating LHC and RC expression or not (i.e. whether we can reproduce the macroscopic behavior of LHC/RC synthesis).
- Since we focus here on R. rubrum, LH2 is not considered and the ratio of LH1/RC is constant.

With
\[
\beta = \frac{2F}{RT} \left( \Delta G_{m,\text{QH2}}^{0} - \Delta G_{m,\text{O2}}^{0} \right) + \frac{N_{\text{H}_2}}{F} (1 - T/T_{c}) \\
\approx \frac{2F}{RT} \left( \Delta G_{m,\text{QH2}}^{0} - \Delta G_{m,\text{O2}}^{0} \right) + 1340 \text{ mV}
\]

Decomposing the fluxes in the ETC into EMs

Having the kinetic model simulated, we can use the calculated steady-state flux values in the ETC to determine the relative contribution of the elementary modes (Table I and Figures 2 and 4) for the simulated scenario. A reaction flux vector in a stoichiometric network can always be decomposed into a weighted sum of EMs but this decomposition is usually not unique and several different decomposition algorithms have been proposed (Schwartz and Kanehisa, 2006). However, in the case of the nine EMs computed for the ETC, with the following reasonable assumptions the decomposition of a given flux vector is straightforward:

- As in the dynamic simulations, we do not consider combined activity of photosynthesis and respiration.
- If ubiquinol oxidase (Ubioxidase_flux > 0) and cytochrome oxidase (Oxi_flux > 0) operate concurrently, we assume that their relative contributions in respiration with succinate (EM5 and EM6) and with NADH (EM7 and EM8) as electron donor as well as in aerobic reverse electron flow (EM3 and EM4) are distributed as the ratio of these two fluxes.
- EMs are arbitrarily scaleable. Here, we assume EM1 to be normalized on RC_flux, EM2, EM5, EM6 and EM9 on Succ_DH_flux and EM3, EM4, EM7 and EM8 on NADH_DH_flux and then compute the respective coefficients.
- Proton leak (considered in the dynamic model but not explicitly in the stoichiometric model) is subsumed as equivalent proton efflux in the ATP synthase flux.

With these assumptions, for a given steady-state flux vector of the dynamic model we can assign the relative activity for each EM with the following simple rules (for variable names, see Table III):

*EM1 = EM2 = EM3 = EM4 = EM5 = EM6 = EM7 = EM8 = EM9 = 0;*

*if(Succ_DH_flux < 0) % Fumarate Reduction EM9 = Succ_DH_flux*

*end*

*respflux = Oxidase_flux + Ubioxidase_flux*

*if(respflux = 0)*

*if(RC_s1s2_flux > 0) % photosynthesis EM1 = RC_flux*

*end*

*if(Succ_DH_flux > 0) % photosynthetic reverse electron flow EM2 = Succ_DH_flux*

*EM1 = EM1-Succ_DH_flux % subtract photosynthesis flux for reverse electron flow end end*

*else % Respiration*

*if(NADH_DH_flux < 0) % aerobic reverse electron flow EM3 = Ubioxidase_flux*NADH_DH_flux/respflux*

*EM4 = Oxidase_flux*NADH_DH_flux/respflux*

*end if(Succ_DH_flux > 0) % respiration with electron from succinate EM5 = Ubioxidase_flux*(NADH_DH_flux-EM9)/respflux*

*EM6 = Oxidase_flux*(Succ_DH_flux-EM4)/respflux; end end*
Model implementation and availability
The full model comprises 33 time-dependent variables of which 10 are state-variables (mathematically expressed by differential equations); the rest depends algebraically upon the state variables (Table III). The model has 55 parameters; 17 of them could be directly taken (or estimated) from literature values. The other 38 parameters were estimated by fitting an available set of measurements given in Table II.

We implemented and simulated the ETC model with self-written MATLAB scripts and implemented it additionally with the Systems Biology Toolbox (Schmidt and Jirstrand, 2006; http://www.sbttoolbox.org/) for MATLAB. The input file for the Systems Biology Toolbox can be downloaded from http://www.mpi-magdeburg.mpg.de/projects/cna/etcmodel.html.

Supplementary information
Supplementary information is available at the Molecular Systems Biology website (www.nature.com/msb).

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