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A model for describing the light response of the non-photochemical quenching of chlorophyll fluorescence

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Abstract

The operation of photosynthetic energy-dissipating processes is commonly characterised by measuring the light response of the non-photochemical quenching (NPQ) of chlorophyll fluorescence, or NPQ vs. $E$ curves. This study proposes a mathematical model for the quantitative description of the generic NPQ vs. $E$ curve. The model is an adaptation of the Hill equation and is based on the close dependence of NPQ on the xanthophyll cycle (XC). The model was tested on NPQ vs. $E$ curves measured in the plant *Arabidopsis thaliana* and the diatom *Nitzschia palea*, representing the two main types of XC, the violaxanthin-antheraxanthin-zeaxanthin (VAZ) type and the diadinoxanthin-diatoxanthin (DD-DT) type, respectively. The model was also fitted to a large number of published light curves, covering the widest possible range of XC types, taxa, growth conditions, and experimental protocol of curve generation. The model provided a very good fit to experimental and published data, coping with the large variability in curve characteristics. The model was further used to quantitatively compare the light responses of NPQ and of PSII electron transport rate, ETR, through the use of indices combining parameters of the models describing the two types of light-response curves. Their application to experimental and published data showed a systematic large delay of the build-up of NPQ relatively to the saturation of photochemistry. It was found that when ETR reaches saturation, NPQ is on average still below one fifth of its maximum attainable level, which is only reached at irradiances about three times higher. It was also found that organisms having the DD-DT type of XC appeared to be able to start operating the XC at lower irradiances than those of the VAZ type.

Keyword index: Chlorophyll fluorescence; modelling; non-photochemical quenching; photoacclimation; photoprotection; xanthophyll cycle
Photosynthesis requires a balance between maximizing light absorption and minimizing damages caused by excessively absorbed light energy. Under natural conditions, exposure to sunlight involves unavoidable risks to the photosynthetic apparatus due to the formation of reactive oxygen species of photosynthetic origin produced under excess light (Osmond et al. 1997; Ort 2001; Demming-Adams and Adams 2006). The sustainment of prolonged photosynthetic activity under high solar irradiances is ensured by the regulation of the repartition of absorbed light energy between photochemistry and energy-dissipating pathways. The latter function as photoprotective process against permanent damages to the photosynthetic apparatus, or photoinhibition, and are therefore of considerable interest for understanding the resistance of photoautotrophs to environmental stress (Demming-Adams and Adams 2006; Lavaud 2007; Li et al. 2009).

One of the most important photoprotective processes is the non-photochemical quenching (NPQ). NPQ groups several pathways among which the thermal dissipation of excess energy (or $q_E$, the energy-dependent quenching, Müller et al., 2001) is considered as the most important. $q_E$ takes place in the light-harvesting antenna of the photosystem II (PS II). It relies on the presence of specific PS II antenna proteins (PsbS in plants, Li et al. 2000; LhcSR in green microalgae, Peers et al. 2009; Lhcx in diatoms, Bailleul et al. 2010), the building of a transthylakoid proton gradient ($\Delta$pH), and the subsequent enzyme-mediated operation (de-epoxidation) of the xanthophyll cycle (XC) (Goss and Jakob, 2010). The XC produces de-epoxidised forms of xanthophylls which are essential regulatory partners of $q_E$, especially in some groups of microalgae (like the diatoms) (Lavaud, 2007; Li et al. 2009; Goss and Jakob 2010). In plants, green algae and some Heterokontophyta (brown algae, Chrysophyceae), the XC is based on the reversible conversion of pigment violaxanthin (Vx) to the $q_E$-involved zeaxanthin (Zx), passing through the intermediate form antheraxanthin (Ax) (VAZ type of XC); in other algal groups, like the diatoms, dinoflagellates, Xanthophyceae and Haptophyta, the XC consists in the conversion between only two pigments, diadinoxanthin (DD) and the $q_E$-involved...
diatoxanthin (DT) (DD-DT type of cycle) (Olaizola and Yamamoto 1994; Lavaud et al. 2004; Lavaud 2007). In phycobilisome-containing organisms, there is no XC (cyanobacteria) or its presence is uncertain (red algae) (Goss and Jakob 2010).

The operation of NPQ processes is usually quantified using variable chlorophyll fluorescence (Pulse Amplitude Modulation fluorometry, PAM), by calculating the fluorescence index NPQ, which is based on the relative difference between the maximum fluorescence measured in the dark-adapted state, $F_m$, and upon exposure to light, $F_m'$ (see Table 1 for notation):

$$\text{NPQ} = \frac{F_m - F_m'}{F_m}$$ (1)

This index is a rearrangement of the Stern-Volmer equation and reflects the assumption that the reciprocal of fluorescence yield is proportional to the Zx or DT concentration (Bilger and Björkman 1990; Lavaud et al. 2002; Baker and Oxborough 2004). The NPQ index has been routinely used to quantify the operation of photoprotective processes as well as the extent of photoinhibitory damages. Under experimental conditions allowing to assume that the prevailing processes causing the quenching of fluorescence are of photoprotective nature, NPQ has been used as a measure of the overall photoprotective capacity of the photosynthetic apparatus (e.g. Dimier et al. 2007b; Lavaud et al. 2007).

One common way to characterise the operation of photoprotective processes or the susceptibility to photoinhibition is to quantify the light response of NPQ. This is done by constructing NPQ vs. $E$ curves that record the development of NPQ with increasing incident irradiance. These curves are analogous to the more frequently measured light-response curves of the PSII electron transport rate (ETR) in the sense that they represent the variation of steady state photosynthetic activity between different light levels, thus not informing on the kinetics of NPQ generation, but on the NPQ attainable under each irradiance. The shape of NPQ vs. $E$ curve varies widely, both regarding its overall shape as well as the absolute NPQ values.
Typically, NPQ increases monotonically with irradiance, varying from zero (measured in darkness) to maximum values that vary greatly with taxonomical groups, physiological state, environmental constraints, or light levels applied. The curve presents a variable degree of sigmoidicity, ranging from cases when NPQ starts to increase steeply from the lowest light levels, following a simple saturation-like pattern (no sigmoidicity), to cases when NPQ remains close to zero for a range of low light levels, then showing an abrupt increase only for intermediate irradiances before stabilizing at maximum values (highly sigmoid). Often, although the stabilization of NPQ is evident, a constant value is not reached within the range of irradiances applied and the maximum attainable NPQ cannot be estimated.

Light-response curves of NPQ have been analysed and interpreted on the basis of arbitrarily chosen features, by qualitatively describing its shape (e.g. low or high sigmoidicity), or by selecting NPQ values reached at particular $E$ levels (e.g. NPQ at maximum applied $E$).

The absence of an adequate descriptive model impedes the characterization of the curve along the whole range of irradiances applied. Also, the lack of a commonly-used set of descriptive parameters makes it difficult to compare curves measured in different studies or experimental conditions or taxonomic groups.

The present study proposes a simple mathematical model for the quantitative description of the generic NPQ vs. $E$ curve, by means of the estimation of a small number of physiologically-meaningful parameters. The model is based on the close, but not absolute, dependence of NPQ on the operation of the XC, through the light-induced de-epoxidation of PS II antenna pigments Vx or DD. To illustrate the main features and variability of the NPQ vs. $E$ curve, as well as the interpretation of model parameters, light-response curves were measured on a plant (*Arabidopsis thaliana* (L.); VAZ type XC) and a diatom (*Nitzschia palea* (Kütz) W. Smith; DD-DT type XC) grown under experimental conditions expected to induce a large variability in curve shape and NPQ absolute values. The adequacy of the model was further tested by fitting it to published NPQ vs. $E$ curves measured on a large variety of photosynthetic organisms and experimental conditions, and covering the widest possible range of NPQ values and curve shapes. The usefulness of the model was also illustrated by exploring the relationship
between NPQ vs. $E$ curves and the photoacclimation status, characterized by light-response curves of the PSII electron transport rate, ETR.

**Materials and Methods**

**Model rationale**

The model proposed to describe the light-response curve of NPQ is an adaptation of the Hill equation, originally derived in the context of ligand binding to macromolecules with multiple binding sites. This equation describes the variation of the number of filled binding sites with the increase in ligand concentration, and is routinely used to characterise the cooperativity of enzymatic reactions (Voet and Voet 1990). The use of the Hill equation for modelling the NPQ vs. $E$ curve is based on the main assumption that NPQ is mostly due to the operation of the XC coupled to the build-up of the transthylakoid ΔpH, or energy-dependent quenching ($q_E$; Müller et al. 2001), and that the other potential components of NPQ, the quenching due to state transitions ($q_I$) or to photoinhibition ($q_I$) are not expected to significantly affect the NPQ vs. $E$ curves (see Discussion). The analogy underlying this rationale is to consider the epoxidized XC pigments Vx and DD as representing the macromolecule of the Hill model, and the transthylakoidal ΔpH-dependent protonation of specific light-harvesting complex (LHC) antenna sites as corresponding to the ligand concentration that bind to their multiple binding sites. At steady state, each irradiance level corresponds to a stable transthylakoidal ΔpH and resulting Vx or DD de-epoxidation and activation through protonation of LHC sites (the so-called ‘activation’ of Zx and DT; Horton et al. 2000; Goss et al. 2006; Lavaud and Kroth 2006; Horton et al. 2008), and thus to a defined NPQ value. The NPQ vs. $E$ curve is thus proposed to be described by the equation:
\[ \text{NPQ} (E) = \text{NPQ}_{\text{m}} \frac{E^n}{E_{50}^n + E^n} \]  

(2)

where \( \text{NPQ}_{\text{m}} \) is the maximum NPQ value reached during the light curve, \( E_{50} \) is the irradiance level for which NPQ attains 50% of \( \text{NPQ}_{\text{m}} \) and \( n \) is the Hill coefficient, characterizing the sigmoidicity of the curve. Whilst the parameters \( E_{50} \) and \( n \) are directly adopted from the Hill equation (originally having only these two parameters), a third parameter, \( \text{NPQ}_{\text{m}} \), had to be considered to account for the variability in the absolute values of NPQ. Under the analogy with the ligand binding context of the Hill equation, \( \text{NPQ}(E) \) is considered to represent the fraction of \( \text{Vx} \) or \( \text{DD} \) molecules de-epoxidized into \( \text{Zx} \) or \( \text{DT} \) and which have been ‘activated’. As such, \( E_{50} \) represents the irradiance level necessary to de-epoxidize and ‘activate’ 50% of convertible \( \text{Vx} \) or \( \text{DD} \) pool (ligand concentration resulting in half occupation of binding sites) necessary to reach the maximal NPQ level, \( \text{NPQ}_{\text{m}} \). Of potential importance for the characterization of the physiological processes underlying the NPQ vs. \( E \) curve is the sigmoidicity parameter \( n \), which measures the type and extent of the reaction cooperativity. In the case of \( n < 1 \), the curve displays a saturation-like increase asymptotically towards \( \text{NPQ}_{\text{m}} \), indicating the presence of a negatively cooperative reaction; if \( n > 1 \), the curve starts to present a sigmoidal shape, the sigmoidicity increasing with \( n \), indicating a positively cooperative, or allosteric, reaction; if \( n = 1 \), the model is reduced to the Michaelis-Menten equation, indicative of a non-cooperative reaction.

Measurement of light-response curves

To test the adequacy of the model, Eq. 2 was first fit to NPQ vs. \( E \) curves generated for organisms representative of the two types of XC, a plant (\( A. \) thaliana; VAZ type of XC) and a diatom (\( N. \) palea; DD-DT type of XC), grown under light conditions expected to induce large variations in NPQ absolute values and in the shape of the NPQ light-response curve. Plants of \( A. \) thaliana (ecotype Columbia) were grown under controlled growth chamber conditions for 4–
5 weeks under a 16:8 h light/dark cycle and 70% relative humidity. The light/dark temperatures were 22/20 °C. Three irradiance levels were applied: 10 ('low light', LL), 75 ('moderate light', ML) and 150 μmol photons m⁻² s⁻¹ ('high light', HL). The diatom Nitzschia palea (Kütz.) W. Smith (collection of the Department of Biology, University of Aveiro) was grown photoautotrophically in unialgal semi-continuous batch 100 mL cultures in sterile natural seawater enriched with f/2 nutrients (Guillard and Ryther 1962). Cultures were grown at 20 °C, under 20 (LL), 100 (ML) and 400 μmol photons m⁻² s⁻¹ (HL) in a 12:12 h light/dark cycle. Cells were harvested by centrifugation (3000 × g, 10 min) during the exponential phase of growth, and were resuspended in fresh growth medium supplemented with NaHCO₃ to a final concentration > 10 μg Chl a ml⁻¹.

Light-response curves of NPQ and ETR were generated by exposing the samples to 7-12 levels of actinic light, up to 920 μmol m⁻² s⁻¹. Samples were dark-adapted for 30 min before the start of the light curve to allow determination of fluorescence levels $F_o$ (minimum fluorescence) and $F_{m'}$ required for the calculation of NPQ (Eq. 1). Samples were light-activated before the start of each light-response curve, through exposure to low light of 54 μmol photons m⁻² s⁻¹ until a steady state in fluorescence was reached (minimum 15 min). Under each light level, a saturation pulse (0.8 s for A. thaliana and 0.6 s for N. palea) was applied and fluorescence levels $F_s$ (steady state fluorescence) and $F_{m'}$ were recorded and used to calculate NPQ (Eq. 1) and ETR, using (Genty et al. 1989):

$$\text{ETR} = E \frac{F_m - F_s}{F_{m'}}$$  \hspace{1cm} (3)

A different sample (plant leaf or algal culture aliquot) was used for measuring NPQ and ETR under each light level. Three replicated measurements were made for each light level. Light-response curves of ETR were described by fitting the model of Eilers and Peeters (1988), and by estimating the parameters $\alpha$ (the initial slope of the curve), $E_{TR_m}$ (maximum ETR) and $E_k$ (the light-saturation parameter). The parameter $E_k$ is commonly interpreted as a measure of the light
level to which a sample is acclimated to, and commonly used to characterise its photoacclimation status (Behrenfeld et al. 2004).

Chlorophyll fluorescence yield was measured using a PAM fluorometer comprising a computer-operated PAM-Control Unit (Walz) and a WATER-EDF-Universal emitter-detector unit (Gademann Instruments GmbH, Germany), using a modulated blue light (LED-lamp peaking at 450 nm, half-bandwidth of 20 nm) as source for measuring, actinic, and saturating light (Cruz and Serôdio 2008). Fluorescence was measured using a 6 mm-diameter Fluid Light Guide fiberoptics. In the case of *N. palea*, the fiberoptics was connected to a fluorescence cuvette (KS-101, Walz, Effeltrich, Germany).

Published light-response curves

The model was also fitted to published NPQ vs. *E* curves, covering a wide range of taxonomic groups (higher plants, mosses, green algae, diatoms, dinoflagellates, others), XC types (VAZ, DD-DT, absence of XC), mutants with variable degrees of impairment of the XC operation, habitats (land and aquatic higher plants; planktonic and benthic microalgae), growth conditions (e.g. low and high light), physiological state (heat stress or XC inhibitors) and fluorescence light-response curve measuring protocols (steady state and rapid light curves, RLCs). Only species with a known functional XC were considered. Studies presenting light curves having less than seven data points were not considered, because of the large errors in the model fitting and parameter estimation. For each study, when three or more light curves were available for each species and experimental treatment, only the two curves presenting the most extreme (higher and lower) NPQ values were used. Detailed information on the data used in the meta-analysis is summarised in Table 3. When available, the corresponding ETR vs. *E* curves were also characterised by fitting the model of Eilers and Peeters (1988) and by estimating its parameters.

Some studies have reported the formation of NPQ in the dark (Perkins et al. 2010). This phenomenon was observed on diatoms (Jakob et al. 1999) and brown algae (Mouget and
and results in NPQ vs. E curves showing a distinctive bi-phasic pattern, with NPQ decreasing from initial high values measured in the dark to a minimum under low to moderate light levels (e.g. Geel et al. 1997; Serôdio et al. 2006). Being relevant only for a limited number of taxonomic groups and physiological conditions, this type of NPQ light-response curve was not considered in the present study.

Model fitting and parameter estimation

The light-response curve models were fitted using a procedure written in Microsoft Visual Basic and based on Microsoft Excel Solver. Model parameters were estimated iteratively by minimizing a least-squares function, forward differencing, and the default quasi-Newton search method. The model can be easily fitted using commonly available software packages. On preliminary tests, this method was compared with the fitting procedures implemented in Sigmaplot 9.0 (Systat Software, Inc., San Jose, USA) and Statistica 8.0 (Statsoft, Inc., Tulsa, USA) and no significant differences were found between the estimates of model parameters.

The standard errors of the parameter estimates were calculated following Ritchie (2008) and are asymptotic standard errors. For nonlinear models such as the one here tested, asymptotic standard errors may not be adequate, as they possibly underestimate the actual parameter uncertainty and cannot evaluate the eventual asymmetry of the confidence regions of estimated parameters (Johnson 2008). A number of alternative approaches exist, the best being based on Monte Carlo methods, such as the Bootstrap (Press et al. 1996; Johnson 2008). However, the benefits of pursuing alternative methods, requiring substantially more computation time, depend on the magnitude of the uncertainty associated to parameter estimation. In this study, the use of asymptotic standard errors was justified by the finding that they were on average relatively low, in a majority of cases below 10% of parameter estimates (see below, Results), in which case they can be considered acceptable (Tellinghuisen 2008).

Results
Model fitting to NPQ vs. $E$ curves

The light-response curves of NPQ measured in the plant *A. thaliana* and in the diatom *N. palea* acclimated to different growth light conditions showed a wide range of curve shapes, varying between two main types: curves with low sigmoidicity, presenting a simple saturation-like pattern (e.g. Fig. 1a, HL), and curves with high sigmoidicity, presenting an initial period of low values, followed by a phase of rapid increase leading to a stage of plateau of maximum values (e.g. Fig. 2a, HL). In all cases, the model provided an excellent fitting to the experimental data throughout the whole range of light levels explaining always more than 99.1% of the data variability (Table 2). Although the residuals (Figs. 1b, 2b) showed in some cases a clearly non-random pattern of variation, values were typically low, between ± 5%. The fitting of the model resulted in a large variation of model parameters with species and growth light conditions, following some consistent trends. Both for *A. thaliana* and *N. palea*, the increase in growth light resulted in an increase in NPQ$_m$ (Fig. 1a, 2a; Table 2). This increase in NPQ$_m$ could be detected despite the fact that in most cases the curves did not reach their maximum values within the range of irradiances applied (e.g. Fig. 1a). Another effect of increasing growth light was the increase in the irradiance level for which the curves reached maximum values. For LL-grown samples, maximum NPQ was reached for lower irradiances, with the curves showing an overall lower sigmoidicity, while for HL-grown samples, curves reached saturation at much higher light levels. This trend could be well described by model parameter $E_{50}$, which increased with growth light in both species (Table 2).

The most noticeable difference between the NPQ light-response curves of *A. thaliana* and *N. palea* was the apparent curve sigmoidicity, with less sigmoid curves being found for the former and more sigmoid curves (and a larger variation) for the latter. This variation in curve sigmoidicity was well characterized by the model parameter $n$, which averaged 1.2 for *A. thaliana* (varying from 0.98 to 1.44), whilst reaching an average value of 2.11 for *N. palea*
(varying from 1.61 to 2.61). However, a different pattern of variation was found for each species, with $n$ decreasing with growth light in *A. thaliana*, and increasing in *N. palea* (Table 2).

Model fitting to published NPQ vs. *E* curves

The compilation of published light-response curves of NPQ allowed gathering a large number of datasets covering a wide range of NPQ absolute values and curve shapes. The model provided a very good fit to the published data, coping with the large diversity in curve characteristics resulting from different combinations of maximum NPQ values attained ($\text{NPQ}_{\text{m}}$), irradiance range of NPQ build-up ($E_{50}$), and curve sigmoidicity ($n$) (Table 3). Considering all cases, the model explained more than 96% of data variability, independently of type of XC (and XC impaired mutants), species, growth conditions, and experimental protocol used for the generation of light curves. A summary of the meta-analysis carried out on the data set of published NPQ vs. *E* curves is presented in Table 4. $\text{NPQ}_{\text{m}}$ was the most variable parameter (c.v. 112.5%), mostly responding to growth conditions, averaging 2.89 but reaching maximum values above 9 for diatoms and even higher for some mosses (Table 3). $E_{50}$ ranged from 30-40 μmol photons m$^{-2}$ s$^{-1}$ to above 3500 μmol m$^{-2}$ s$^{-1}$, in all cases attaining values well above growth irradiances (see below). However, with few exception (the diatoms *Skeletonema costatum* and *Phaeocystis antarctica*, and the high light-grown *A. thaliana*), all values of $E_{50}$ above 1200 μmol m$^{-2}$ s$^{-1}$ were obtained for *A. thaliana* mutants with impaired operation of the XC (Table 3).

$n$ was the least variable parameter (c.v. 50.8%), meaning that the curve shape remained relatively unaltered, and that the variability in the NPQ vs. *E* curves was mainly due to changes in NPQ maximum values and in NPQ onset along the irradiance range. $n$ averaged 1.7, corresponding to a noticeable degree of sigmoidicity, although values around 1.0 (no sigmoidicity) or slightly lower (mostly mutants with impaired XC) as well as above 3.0 (very high sigmoidicity) were also found (Table 3).

Despite the low number of data points forming most NPQ vs. *E* curves, standard errors of parameter estimates were on average relatively low (15.6, 27.3 and 12.2% of estimates for
NPQ_m, E_50 and n, respectively). However, standard errors were typically much higher in the cases when XC was impaired (24.5, 43.9 and 13.5% for NPQ_m, E_50 and n, respectively), indicating that a high level of precision in the estimation of model parameters can be expected for samples under natural conditions.

When comparing different XC types, no clear differences were found between model parameters, due to the large overlap of their range of variation. Model parameters appeared to depend mostly on growth conditions, as larger variations could be found within the same species than amongst different taxonomic groups. The model also described very well the light-response curves of NPQ for composite samples (phytoplankton, microphytobenthos). Model parameters varied independently from each other, as no significant correlations were found between NPQ_m, E_50 or n, either when considering all data pooled together, or when considering for separate for each XC type.

Considering the published data, the effects of growth light levels on model parameters were identical to those described for the experimental data of this study. When compared to lower light conditions, NPQ light-response curves of samples acclimated to high light showed higher values of NPQ_m (Bilger and Bjorkman 1990; Burritt and MacKenzie 2003; Müller et al. 2004; Ralph and Gademann 2005; Rodríguez-Calcerrada et al. 2007; Cruz and Serôdio 2008) and, with the exception of Ralph and Gademann (2005), all these studies showed a similar increase in E_50 with growth light. The model also fitted very well the light-response curves of NPQ with impaired functioning of the XC, illustrating the usefulness of estimating model parameters to characterize quantitatively the effects on NPQ of inhibitors (e.g. Bilger and Bjorkman 1990), mutations (e.g. Niyogi et al. 1998), or changes in temperature (Ralph et al. 2001).

Finally, the model was also found to be adequate to describe NPQ vs. E curves generated under non-steady state conditions, i.e. derived from RLCs. The comparison of the model parameters estimated for light curves of different light steps showed the same pattern on VAZ and DD-DT samples, with shorter light steps resulting in lower NPQ_m and higher E_50, whilst n not showing any consistent trend (Ralph and Gademann 2005; Perkins et al. 2006).
Relationship between NPQ and ETR light-response curves

To illustrate the application of the model to evaluate to what extent the NPQ response to light is related to the photoacclimation status, the light levels required for induction of NPQ and for photochemistry or ETR saturation were compared. A simple form of achieving this is by the direct comparison of model parameters $E_{50}$ and $E_k$ estimated for the two types of light-response curves. These parameters have the advantage of being measured in the same scale (PAR irradiance, $\mu$mol photons m$^{-2}$ s$^{-1}$) and of being independent of the units used to measure photosynthetic rates, ETR or NPQ, which can vary with the method in use (the case of photosynthesis) or instrument settings (the case of ETR).

For the experimental data obtained in this study, the increase in growth irradiance induced a substantial change in the photoacclimation status, noticeable by a clear change in the ETR vs $E$ curves. As a result, the increase observed for $E_{50}$ was followed by a similarly large increase in $E_k$, (due to the decrease of $\alpha$ and the increase of $ETR_m$), both in *A. thaliana* (Fig. 3) and in *N. palea* (Fig. 4). However, $E_{50}$ was found to be in all cases higher than $E_k$, by 3.7 and 2.5 times, on average, for *A. thaliana* and *N. palea*, respectively (2.90 for the whole data). This result indicates that the light-induced build-up of substantial values of NPQ started only after linear electron transport reached near saturation. Furthermore, it shows that a large fraction of $E_{50}$ variability was related to variations in the photoacclimation status due to different growth light conditions, as $E_{50}$ increased linearly with $E_k$, both for the plant as for the diatom (although the correlation was not statistical significant, but a clear linear trend is obvious; Fig. 5). This analysis also showed that the slope of the regression line of $E_{50}$ on $E_k$ was higher for *A. thaliana* than for *N. palea*, indicating that, for the same photoacclimation status (the same $E_k$) the plant requires a higher light level for NPQ to reach 50% of its maximum capacity. Or, in other words, that NPQ is induced latter in the range of irradiances in the plant than in the diatom.

Interestingly, the analysis of the subset of published data presenting both NPQ and ETR light-response curves, showed the same generic trend. When pooling together the published and
experimental data for samples with fully operational XC (i.e. excluding mutants or samples treated with XC inhibitors) and $E_k$ estimates based only on steady state light curves (i.e. excluding RLCs as $E_k$ estimation is largely dependent on light step duration; Serôdio et al. 2006), the ratio $E_{50}/E_k$ was found to be higher on organisms having a VAZ type of XC and to differ significantly from those having a DD-DT type (3.32 and 2.39, respectively; $t$-test $P = 0.038$; Fig. 6a).

Another way to compare the light responses of NPQ and ETR is to calculate the fraction of NPQ that is formed when ETR approaches saturation (i.e. when $E = E_k$), or:

$$\text{NPQ}_{E_k} = \frac{\text{NPQ}(E_k)}{\text{NPQ}_{\text{max}}} \quad (4)$$

Low values of NPQ$_{E_k}$ indicate that when photochemistry saturates, NPQ is still not significantly developed (corresponding to high $E_{50}/E_k$), whilst high values indicate that NPQ responds more promptly to light increase and the approach of ETR saturation (corresponding to low $E_{50}/E_k$).

Considering both published and experimental data (the data subset used above for comparing $E_{50}/E_k$), NPQ$_{E_k}$ averaged 0.16, and remained below 0.35, indicating that when $E = E_k$, NPQ hardly reached over one third of the maximum attainable level. Confirming the expected inverted relationship between NPQ$_{E_k}$ and $E_{50}/E_k$, significantly lower values were found for organisms with a VAZ type of XC than for those with the DD-DT type (0.14 and 0.21, respectively; $t$-test, $P = 0.023$; Fig. 6b).

Model fitting and parameter estimation

Despite the usually reduced number of data points in the analyzed NPQ light-response curves (average 9.7, maximum 13), the model proposed in this study was found to be easily fitted to experimental curves, in most cases yielding parameter estimates virtually independently from the start values used in the iterative fitting procedure (although realistic start values allows a
more rapid and precise parameter estimation). The main problem found when trying to fit Eq. 2
to NPQ vs. E curves occurred when the curves were very close to linear, showing no clear
features like sigmoidicity or saturation. This resulted in that a similarly good fit could be
reached with different combinations of model parameters values. Still, this occurred only in
relatively small number of cases, like the case of the diatom *Skeletonema costatum* (Lavaud et
al. 2007) and some *A. thaliana* mutants with severe impairment of XC operation, for which
some degree of uncertainty remain associated to the presented parameter estimates.

This problem seems to result from the common situation of having light-response
curves constructed with the main purpose of characterizing the ETR vs. E curves, NPQ having
only a secondary interest. As it often happens that the NPQ approaches its maximum for light
levels much above the range used for measuring ETR, the NPQ vs. E curve may result truncated
and its complete shape may not be available for model fitting. Yet, the relationship established
between NPQ and ETR light curves in similar data may be used to impose boundaries to the
model parameter estimates, and thus help to reach a single set of meaningful values.

Discussion

Model assumptions

The model proposed for describing the NPQ vs. E curve is based on the main assumption that
NPQ is mostly due to $q_E$, the ‘energy-quenching’ associated to the build-up of the transthylakoid
ΔpH and the operation of the XC. This assumption is generally supported by the finding of
strong linear relationships between NPQ and de-epoxidized xanthophylls in all groups
investigated (plants, green algae, diatoms, Chrysophyceae) and under a wide range of
experimental conditions.

However, besides $q_E$, NPQ also quantifies the fluorescence quenching caused by state
transitions ($q_T$) or by photoinhibition ($q_I$) which may potentially significantly affect the NPQ vs.
$E$ curves. $q_T$ is relevant in phycobilisome-containing organisms (cyanobacteria and red algae) (Campbell et al. 1998; Mullineaux and Emlyn-Jones 2004), although not as much in higher plants and green algae (Pfannschmidt 2005; Eberhard et al. 2008; Ruban and Johnson 2009), and it does not occur for the Heterokontophyta (Lavaud 2007). When occurring, $q_T$ is not significant in light conditions triggering $q_E$-related NPQ (Mullineaux and Emlyn-Jones 2004; Tikkanen et al. 2006; Ruban and Johnson 2009). With the exception of some land plants (overwintering conifers and tropical evergreen species), $q_I$ origin is not clearly defined and it requires particular conditions such as prolonged environmental stress (Müller et al. 2001; Demmig-Adams and Adams 2006; Horton et al. 2008).

The effects of $q_T$ or $q_I$ on the light-response curve of NPQ would likely consist in a general change in curve shape, through the increase of NPQ at low light levels (for $q_T$) and of $NPQ_m$ (for $q_I$). However, these components of NPQ may be expected not to significantly affect the NPQ vs. $E$ curves measured under the usually applied experimental protocols. This is because the duration of the light steps commonly applied ($\leq 2$-3 min, and much shorter in the case of RLCs) correspond to light doses clearly different than those required to induce both $q_T$ or $q_I$ (Campbell et al. 1998; Müller et al. 2001; Demmig-Adams and Adams 2006; Horton et al. 2008; Ruban and Johnson 2009).

The same reasoning applies to another implicit assumption of the model, that during the generation of a light curve $q_E$ is due solely to the development of the $\Delta p$ and the subsequent de-epoxidation of the present (and susceptible of being de-epoxidized) pool of xanthophylls. In fact, it is well established that exposure to high light can induce the de novo synthesis of XC pigments (e.g. Olaizola and Yamamoto 1994; D’Haese et al. 2004; Lavaud et al. 2004; Demmig-Adams and Adams 2006). This process causes the accumulation of de-epoxidized forms, which could result in the linear increase of NPQ in the high-light part of the curve.

Again, the de novo synthesis requires the prolonged exposure to very high irradiances (e.g. Lavaud et al. 2004; Demmig-Adams and Adams 2006; Lavaud 2007), well above the light doses applied during the construction of NPQ light-response curves.
Nevertheless, even if significant $q_T$, $q_I$ or \textit{de novo} pigment synthesis do occur, the resulting curve may still be of the same general shape and describable by the model. In fact, this study showed that the model fits very well to data obtained in a wide range of experimental conditions, in which the occurrence of any of these processes cannot be completely ruled out (Table 3). In this case, however, care should be taken in interpreting and comparing the model parameters estimated for different species or condition, as they may be affected differently by processes other than the build-up of the $\Delta p$H and the XC. In the case of organisms with impaired operation of the XC (mutants or inhibitor-treated samples), the model was nevertheless shown to be of value, providing a useful quantitative characterization of the light response of NPQ.

Another major assumption of the model is that the formation of $q_E$, through the de-epoxidation of Vx or DD generated by the acidification of the lumen relates to incident irradiance following the generic conditions of application of the Hill equation, i.e. that NPQ can be considered as allosterically regulated by irradiance through the build-up of the transthylakoidal $\Delta p$H. This assumption is partially supported by the finding that, in isolated plant chloroplasts, the regulation of $q_E$ is of allosteric nature, resulting in response curves of $q_E$ as a function of $\Delta p$H of sigmoidal shape, shown to be adequately described by the Hill equation (Pfündel and Dilley 1993; Ruban et al. 2001; Takizawa et al. 2007; Pérez-Bueno et al. 2008). This sigmoidicity is attributed to the protonation of binding sites of the LHC protein Psbs induced by lumen acidification, and subsequent conformational changes within the LHC, switching the de-epoxidized xanthophylls to an ‘activated’ state. It is taken as an indication of allosteric regulation of $q_E$, resulting from proton binding showing positive cooperativity, and the ‘activated’ de-epoxidized xanthophylls Zx (Horton et al. 2000; Horton et al. 2008; Li et al. 2009) or DT (Goss et al. 2006; Lavaud and Kroth 2006) acting as a positive effector.

The strong physiological basis of the proposed model is likely to explain its success to cope with a large diversity of light-response curve characteristics. Other mathematical models have also been used to successfully describe NPQ vs. $E$ curves, namely an exponential saturation function, the Michaelis-Menten function, and a hyperbolic tangent function (Ritchie
2008). However, these models have only been tested on a reduced dataset, and on light curves of low sigmoidicity. Interestingly, the model that yielded the better fit to experimental data was the Michaelis-Menten equation (Ritchie 2008), a particular case of the model proposed in the present study.

Model fitting to NPQ vs. $E$ curves

The usefulness of the model was demonstrated by the possibility to estimate NPQ$_m$ despite the fact that in most cases the curves did not reach a saturation plateau within the range of irradiances applied (e.g. Fig. 1a). When assuming NPQ to represent mostly $q_E$, the maximum NPQ reached by a sample quantifies its photoprotective potential via the dissipation of excessive light energy. By applying the model, curves measured under different experimental conditions can be readily compared, and changes in NPQ$_m$ (e.g. following changes in growth light conditions) can be quantified, which would be otherwise impossible. The accurate estimation of NPQ$_m$ is also required for quantifying $E_{50}$, which, in some cases of strict relationship (e.g. for most Heterokontophyta; Lavaud et al. 2004; Lavaud 2007), is interpretable as the light level corresponding to the de-epoxidation and ‘activation’ of the xanthophyll convertible pool necessary to induce half of the maximal NPQ. The size of the xanthophyll convertible pool can be very different in VAZ and DD-DT organisms varying with species and growth conditions, ranging between 50 and 70% (Lavaud 2007; Goss and Jakob 2010). Again, the fit of Eq. 2 to NPQ vs. $E$ curves provided an adequate form of characterizing how promptly the XC is ΔpH-activated when responding to an increase in incident irradiance.

However, while NPQ$_m$ and $E_{50}$ can be grossly estimated by visual inspection of the curves (at least when saturation is reached for the light levels applied), the curve sigmoidicity can hardly be characterized quantitatively without the fit of the model and estimation of $n$. The sigmoidicity coefficient was found to vary around 1.5-2.0, values corresponding to positive cooperativity and thus indicating the allosteric regulation of NPQ by light. Minimum $n$ values averaged about 1 (reaching significantly lower values only for cases of XC impairment),
indicate that negative cooperativity is not involved in the regulation of processes underlying NPQ. These results generally support that the allosteric nature known for the more fundamental regulation of $q_E$ by $\Delta$pH in plants (Ruban et al. 2001; Li et al. 2009) holds for the relationship of NPQ to irradiance in a large diversity of photosynthetic organisms.

The Hill coefficient has been interpreted as representing the number of allosteric regulators (D’Haese et al. 2004). Considering the diversity of factors likely to contribute to the NPQ vs. $E$ curve (Demmig-Adams and Adams 2006; Eberhard et al. 2008; Horton et al. 2008; Li et al. 2009), some of them not fully identified in some photosynthetic taxa (Lavaud 2007), $n$ should be considered simply as an empirical coefficient informing on the degree of allostery and serving as a practical descriptor of the curve shape.

Relationship of NPQ light response to photoacclimation status

The modelling of the NPQ vs. $E$ curves opens the new possibility of quantitatively comparing the light responses of NPQ and of photochemistry or ETR. Photosynthesis and ETR light-response curves are very commonly used as a form of characterizing the photoacclimation status of photosynthetic organisms, through the estimation of the parameters of a number of available models (Henley 1993; Behrenfeld et al. 2004; Ralph and Gademann 2005; Guarini and Moritz 2009; Perkins et al. 2010). By describing the NPQ vs. $E$ curve by a small set of meaningful parameters, the light response of NPQ can be characterized relatively to the one of ETR or photosynthetic rate.

This possibility is useful for the definition of the light levels corresponding to the onset of NPQ (e.g. activation of the XC) relatively to the saturation of photochemistry. As the light response of NPQ may depend greatly on the photoacclimation status, the characterization and comparison of NPQ vs. $E$ curves should be preferably normalized to parameters indicative of photochemistry saturation. Furthermore, the comparison of the NPQ and ETR light responses is also of interest because it provides insight on the way an organism combines the responses of photochemistry and of photoprotective processes to changes in ambient light. However, without
adequate parameterization of the NPQ light-response, this question can be answered only
approximately.

Despite its simplicity, the indices \( E_{50}/E_k \) and NPQ_{\text{Ek}} here proposed provide an efficient
form of comparing the light levels for which NPQ reaches significant values and
photochemistry reaches saturation. The application of these indices allowed quantifying the
delay of the light response of NPQ relatively to the saturation of photochemistry. Under the
assumption that NPQ mostly represents \( q_{E_k} \), it showed that it takes about three times the
irradiance at \( E_k \) for half of \( q_{E_k} \) to develop and that, at \( E_k \), the formation of \( q_{E_k} \) is still at relatively
low levels, providing typically less than one fifth of maximum photoprotective capacity. For the
cases when \( q_{E_k} \) is strictly related to the XC operation (e.g. in most of Heterokontophyta; Lavaud
et al. 2004; Lavaud 2007), the values of the indices \( E_{50}/E_k \) and NPQ_{\text{Ek}} can be interpreted in
terms of the de-epoxidation state of the convertible pool of xanthophylls and their ‘activation’
upon LHC protonation. Yet, the modelling of the NPQ vs. \( E \) curve may allow for the
development of other indices that better characterize this relationship.

Interestingly, the meta-analysis carried out in this study showed consistent differences
between the relationships of NPQ light response to photoacclimation status in organisms with
VAZ and DD-DT types of XC. The fact that these differences were found despite the diversity
in taxa and growth conditions may indicate that they are due to fundamental differences in the
way the two types of organisms cope with high light, especially regarding the underlying \( q_{E_k} \)
mechanism and regulation of the XC, illustrating the advantages provided by the quantitative
description of the NPQ vs \( E \) curve.

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Table 1. Notation

\( \alpha \) – Initial slope of the ETR vs. \( E \) curve

Ax – Antheraxanthin

DD – Diadinoxanthin

DT – Diatoxanthin

\( E \) – PAR irradiance (\( \mu \text{mol photons m}^{-2} \text{s}^{-1} \))

\( E_{50} \) – Irradiance level corresponding to 50% of \( \text{NPQ}_m \) in a NPQ vs. \( E \) curve

\( E_k \) – Light-saturation parameter of the ETR vs. \( E \) curve

ETR – PSII relative electron transport rate

\( \text{ETR}_m \) – Maximum ETR in a ETR vs. \( E \) curve

\( F_o, F_m \) – Minimum and maximum fluorescence of a dark-adapted sample

\( F_o', F_m' \) – Steady state and maximum fluorescence of a light-adapted sample

NPQ – Non-photochemical quenching

\( \text{NPQ}_m \) – Maximum NPQ value reached in a NPQ vs. \( E \) curve

\( \text{NPQ}_{Ek} \) – Fraction of \( \text{NPQ}_m \) reached when \( E = E_k \)

\( n \) – Sigmoidicity coefficient of the NPQ vs. \( E \) curve

PSII – Photosystem II

RLC – Rapid light-response curve

VAZ – Vx-Ax-Zx XC

Vx – Violaxanthin

XC – Xanthophyll cycle

Zx – Zeaxanthin
Table 2. Summary of the results of the fitting of the model to NPQ vs. \( E \) curves measured in \( A. \) \textit{thaliana} and \( N. \) \textit{palea} grown under low (LL), moderate (ML) and high light (HL). Parameter estimates ± one standard error.

<table>
<thead>
<tr>
<th>Model parameters</th>
<th>NPQ(_{m})</th>
<th>( E_{50} )</th>
<th>n</th>
<th>( r^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( A. ) \textit{thaliana}</td>
<td>LL</td>
<td>3.91 ± 0.82</td>
<td>209.9 ± 8.31</td>
<td>1.44 ± 0.52</td>
</tr>
<tr>
<td></td>
<td>ML</td>
<td>4.63 ± 0.24</td>
<td>542.0 ± 49.2</td>
<td>1.21 ± 0.04</td>
</tr>
<tr>
<td></td>
<td>HL</td>
<td>5.90 ± 1.04</td>
<td>1141.0 ± 372.1</td>
<td>0.98 ± 0.06</td>
</tr>
<tr>
<td>( N. ) \textit{palea}</td>
<td>LL</td>
<td>2.70 ± 0.10</td>
<td>91.7 ± 7.01</td>
<td>1.61 ± 0.19</td>
</tr>
<tr>
<td></td>
<td>ML</td>
<td>3.65 ± 0.05</td>
<td>312.6 ± 5.59</td>
<td>2.16 ± 0.05</td>
</tr>
<tr>
<td></td>
<td>HL</td>
<td>3.86 ± 0.11</td>
<td>551.4 ± 15.8</td>
<td>2.55 ± 0.08</td>
</tr>
</tbody>
</table>
Table 3. Results of fitting of Eq. 2 to published NPQ vs. \( E \) curves. Composite: samples containing organisms with different XC types. LC: light curve protocol (SS: steady state LL; RLCx: rapid light curve, x seconds light step; N-SLC: non-sequential light curve). Treatment: conditions applied before the measurement of the light curves (PAR, temperature); growth conditions when more than one curve available for the same taxon (HL: high light; LL: low light; DTT: XC inhibitor dithiothreitol; LL/HL: transfer from LL to HL; Dense: plants from dense forest stand; Edge: plants from forest edge; IL: intermittent light; CL: continuous light; wt: wild type; aba1, crr6, CRR6, lut1, lut2, npq1, npq2, npq4, npq5, vtc2: mutants; LI-1, Sf-2, Col-0, Ws-2: Arabidopsis accessions; Mixing: water column mixing). Parameter estimates ± one standard error.

<table>
<thead>
<tr>
<th>XC type</th>
<th>Taxon</th>
<th>LC</th>
<th>Treatment</th>
<th>( \text{NPQ}_m )</th>
<th>( E_{50} )</th>
<th>( n )</th>
<th>( r^2 )</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>VAZ</td>
<td>Magnoliophyta (flowering plants)</td>
<td>Arabidopsis thaliana</td>
<td>SS</td>
<td>wt 10 °C</td>
<td>2.07 ± 0.04</td>
<td>113.6 ± 6.0</td>
<td>1.11 ± 0.08</td>
<td>0.998</td>
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<td></td>
<td>2.98 ± 0.40</td>
<td>1118.0 ± 293.8</td>
<td>1.09 ± 0.11</td>
<td>0.996</td>
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<td></td>
<td>npq1 10 °C</td>
<td>1.68 ± 0.25</td>
<td>636.5 ± 587.0</td>
<td>0.40 ± 0.06</td>
<td>0.998</td>
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<td>npq1 25 °C</td>
<td>1.16 ± 0.22</td>
<td>867.1 ± 557.5</td>
<td>0.63 ± 0.10</td>
<td>0.995</td>
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<tr>
<td>SS</td>
<td></td>
<td></td>
<td></td>
<td>Li-1</td>
<td>3.76 ± 0.18</td>
<td>563.8 ± 40.6</td>
<td>1.83 ± 0.16</td>
<td>0.998</td>
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<td>Sf-2</td>
<td>4.12 ± 0.16</td>
<td>661.9 ± 33.7</td>
<td>2.03 ± 0.13</td>
<td>0.999</td>
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<td></td>
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<td></td>
<td>Col-0</td>
<td>3.29 ± 0.19</td>
<td>683.7 ± 55.0</td>
<td>1.84 ± 0.16</td>
<td>0.998</td>
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<td>Ws-2</td>
<td>3.32 ± 0.19</td>
<td>599.8 ± 49.1</td>
<td>1.96 ± 0.21</td>
<td>0.997</td>
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<tr>
<td>SS</td>
<td></td>
<td></td>
<td></td>
<td>wt</td>
<td>2.06 ± 0.08</td>
<td>184.1 ± 13.4</td>
<td>2.21 ± 0.32</td>
<td>0.993</td>
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<td>lut1</td>
<td>1.52 ± 0.06</td>
<td>195.5 ± 13.7</td>
<td>2.28 ± 0.33</td>
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<td>lut2</td>
<td>1.76 ± 0.10</td>
<td>231.0 ± 22.2</td>
<td>1.72 ± 0.22</td>
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<td>aba1</td>
<td>2.36 ± 0.19</td>
<td>251.1 ± 43.2</td>
<td>1.14 ± 0.13</td>
<td>0.996</td>
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<td>lut2aba1</td>
<td>1.17 ± 0.07</td>
<td>214.7 ± 30.6</td>
<td>1.08 ± 0.10</td>
<td>0.997</td>
</tr>
<tr>
<td>Species</td>
<td>Treatment</td>
<td>X (μmol/m²/s)</td>
<td>Y (μmol/m²/s)</td>
<td>R²</td>
<td>Source</td>
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<tr>
<td><em>Begonia erythrophylla</em></td>
<td>LL</td>
<td>1.27 ± 0.04</td>
<td>388.3 ± 14.1</td>
<td>3.51 ± 0.36</td>
<td>Burritt and Mackenzie (2003)</td>
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<tr>
<td></td>
<td>LL/HL</td>
<td>1.77 ± 0.05</td>
<td>467.3 ± 15.8</td>
<td>2.95 ± 0.23</td>
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<tr>
<td></td>
<td>HL</td>
<td>2.83 ± 0.17</td>
<td>618.3 ± 38.6</td>
<td>2.82 ± 0.32</td>
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<tr>
<td><em>Hedera canariensis</em></td>
<td>LL</td>
<td>3.61 ± 0.04</td>
<td>382.6 ± 6.3</td>
<td>2.90 ± 0.13</td>
<td>Bilger and Björkman (1990)</td>
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<tr>
<td></td>
<td>LL+DTT</td>
<td>1.14 ± 0.18</td>
<td>656.2 ± 218.9</td>
<td>1.09 ± 0.20</td>
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<td></td>
<td>HL</td>
<td>5.34 ± 0.33</td>
<td>761.1 ± 59.6</td>
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<td>HL+DTT</td>
<td>1.48 ± 0.05</td>
<td>437.0 ± 27.8</td>
<td>2.20 ± 0.26</td>
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<tr>
<td><em>Quercus petraea</em></td>
<td>Dense</td>
<td>2.23 ± 0.03</td>
<td>158.2 ± 6.3</td>
<td>1.83 ± 0.13</td>
<td>Rodríguez-Calcerrada et al. (2007)</td>
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<td>Edge</td>
<td>5.55 ± 0.11</td>
<td>423.2 ± 13.3</td>
<td>2.04 ± 0.10</td>
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<tr>
<td><strong>Zostera marina</strong> SS</td>
<td>LL</td>
<td>0.87 ± 0.15</td>
<td>797.3 ± 302.0</td>
<td>1.01 ± 0.14</td>
<td>0.995</td>
<td>Ralph and Gademann (2005)</td>
<td></td>
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<tr>
<td>SS HL</td>
<td>3.12 ± 0.25</td>
<td>400.5 ± 65.3</td>
<td>1.30 ± 0.16</td>
<td>0.995</td>
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<tr>
<td>RLC5 HL</td>
<td>1.36 ± 0.06</td>
<td>276.9 ± 26.9</td>
<td>1.39 ± 0.17</td>
<td>0.997</td>
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<td>RLC10 HL</td>
<td>2.34 ± 0.17</td>
<td>399.0 ± 60.7</td>
<td>1.02 ± 0.11</td>
<td>0.997</td>
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<td>RLC40 HL</td>
<td>3.37 ± 0.15</td>
<td>360.7 ± 30.4</td>
<td>1.45 ± 0.12</td>
<td>0.998</td>
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<tr>
<td><strong>Bryophyta</strong> (mosses)**</td>
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<tr>
<td><em>Eurhynchium crassinervium</em> SS</td>
<td>5.79 ± 0.58</td>
<td>237.9 ± 34.7</td>
<td>1.53 ± 0.12</td>
<td>0.998</td>
<td>Marschall and Proctor (2004)</td>
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<tr>
<td><em>Pogonatum urnigerum</em></td>
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<tr>
<td><em>Polytrichum juniperinum</em></td>
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<tr>
<td><em>Tortula (Syntrichia) ruralis</em></td>
<td>12.23 ± 0.42</td>
<td>458.6 ± 16.5</td>
<td>2.61 ± 0.15</td>
<td>0.999</td>
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<tr>
<td><em>Racomitrium aquaticum</em></td>
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<td><em>Trichocolea tomentella</em></td>
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<tr>
<td><strong>Chlorophyta</strong> (green algae)**</td>
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<tr>
<td><em>Chara corallina</em> SS</td>
<td>1.67 ± 0.04</td>
<td>34.0 ± 0.5</td>
<td>6.80 ± 0.73</td>
<td>0.993</td>
<td>Krupenina and Bulychev (2007)</td>
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<tr>
<td><em>Chlamydomonas reinhardtii</em> wt</td>
<td>1.83 ± 1.46</td>
<td>1010.0 ± 1660.0</td>
<td>0.81 ± 0.15</td>
<td>0.992</td>
<td>Elrad et al. (2002)</td>
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<tr>
<td><em>picocelullarum</em> npq5</td>
<td>0.19 ± 0.01</td>
<td>56.2 ± 3.58</td>
<td>3.66 ± 0.74</td>
<td>0.986</td>
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<td><em>Picochlorum sp.</em> SS</td>
<td>1.88 ± 0.29</td>
<td>410.2 ± 142.0</td>
<td>1.06 ± 0.20</td>
<td>0.975</td>
<td>Dimier et al. (2007a)</td>
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<td><strong>Eustigmatophyceae</strong> <em>Nannochloropsis oculata</em> RLC10</td>
<td>1.01 ± 0.06</td>
<td>224.5 ± 31.7</td>
<td>1.01 ± 0.08</td>
<td>0.998</td>
<td>Cosgrove and Borowitzka (2006)</td>
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<tr>
<td><strong>DD-DT</strong> <strong>Bacillariophyceae</strong> (diatoms)**</td>
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<tr>
<td><em>Chaetoceros socialis</em> SS</td>
<td>5.71 ± 0.08</td>
<td>149.4 ± 4.8</td>
<td>1.88 ± 0.10</td>
<td>0.999</td>
<td>Dimier et al. (2007b)</td>
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<td><em>Skeletonema marinoi</em></td>
<td>1.27 ± 0.03</td>
<td>233.0 ± 9.3</td>
<td>1.95 ± 0.13</td>
<td>0.998</td>
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<tr>
<td><em>Thalassiosira rotula</em></td>
<td>1.14 ± 0.18</td>
<td>198.9 ± 89.8</td>
<td>0.89 ± 0.23</td>
<td>0.977</td>
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<tr>
<td><strong>Fragilariopsis cylindrus</strong> SS</td>
<td>1.00 ± 0.18</td>
<td>637.5 ± 266.1</td>
<td>1.08 ± 0.23</td>
<td>0.966</td>
<td>Kropuenske et al. (2009)</td>
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<tr>
<td>65 μmol m⁻² s⁻¹</td>
<td>0.74 ± 0.08</td>
<td>1023.1 ± 233.6</td>
<td>1.05 ± 0.9</td>
<td>0.996</td>
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<td>125 μmol m⁻² s⁻¹</td>
<td>0.59 ± 0.12</td>
<td>1024.5 ± 540.6</td>
<td>0.94 ± 0.13</td>
<td>0.986</td>
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<td><strong>Phaeodactylum tricornutum</strong> SS</td>
<td>2.30 ± 0.10</td>
<td>367.6 ± 24.1</td>
<td>2.59 ± 0.45</td>
<td>0.984</td>
<td>Lavaud et al.</td>
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<td>Organism/Phytoplankton</td>
<td>Temp/C</td>
<td>Initial Stock</td>
<td>Chlorophyll</td>
<td>Biomass</td>
<td>Growth Rate</td>
<td>P/B Ratio</td>
<td>Genus</td>
<td>Year</td>
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<td>Skeletonema costatum</td>
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<td>Navicula phylepta</td>
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<td>Nitzschia palea</td>
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<td>Symbiodinium sp.</td>
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<td>Phaeocystis antarctica</td>
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**Table 1:** Growth performance and字样 representation of various phytoplankton species and microphytobenthos under different conditions. Growth rates are given as mean ± standard deviation, and chlorophyll and biomass data are presented in micrograms per liter (μg L⁻¹) for chlorophyll and milligrams per liter (mg L⁻¹) for biomass. Growth rates are expressed as daily growth rates (μg C cell⁻¹ day⁻¹) and initial stock concentrations are in milligrams per liter (mg L⁻¹).
Table 4. Summary of the meta-analysis of the results of the fitting of the model to published NPQ vs. \( E \) curves (listed in Table 3). Mean values ± one standard error.

<table>
<thead>
<tr>
<th>XC type</th>
<th>( \text{NPQ}_m ) ( \pm ) one std error</th>
<th>( E_{50} ) ( \pm ) one std error</th>
<th>( n ) ( \pm ) one std error</th>
</tr>
</thead>
<tbody>
<tr>
<td>VAZ</td>
<td>3.30 ± 0.49</td>
<td>926.7 ± 135.1</td>
<td>1.77 ± 0.13</td>
</tr>
<tr>
<td>DD-DT</td>
<td>1.88 ± 0.41</td>
<td>657.9 ± 145.9</td>
<td>1.60 ± 0.14</td>
</tr>
<tr>
<td>Mixed</td>
<td>3.68 ± 0.67</td>
<td>472.1 ± 47.3</td>
<td>2.01 ± 0.33</td>
</tr>
<tr>
<td>All</td>
<td>2.89 ± 0.35</td>
<td>824.3 ± 99.2</td>
<td>1.73 ± 0.09</td>
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</tbody>
</table>
Figure legends

Fig. 1. Fitting of Eq. 2 to light-response curves of NPQ measured in the plant *A. thaliana* grown under low (LL), moderate (ML) and high light (HL). a. Light-response curves (data points), fitted model (lines), and estimates of model parameters. b. Residuals of the model fitting.

Fig. 2. Fitting of Eq. 2 to light-response curves of NPQ measured in the diatom *N. palea* grown under low (LL), moderate (ML) and high light (HL). a. Light-response curves (data points), fitted model (lines), and estimates of model parameters. b. Residuals of the model fitting.

Fig. 3. Comparison between the light-response curves of NPQ and of ETR of *A. thaliana* grown under low (LL), moderate (ML) and high light (HL). Estimates of the parameters of the model of Eilers and Peeters (1988) fitted to the ETR vs. *E* curves are presented. Estimates of the parameters of the curves fitted to the NPQ vs. *E* curves are presented in Table 2.

Fig. 4. Comparison between the light-response curves of NPQ and ETR of *N. palea* grown under low (LL), moderate (ML) and high light (HL). Estimates of the parameters of the model of Eilers and Peeters (1988) fitted to the ETR vs. *E* curves are presented. Estimates of the parameters of the curves fitted to the NPQ vs. *E* curves are presented in Table 2.

Fig. 5. Relationship between the parameter *E*$_{50}$ of Eq. 2 fitted to the NPQ vs. *E* curves and the parameter *E*$_{k}$ of the Eilers and Peeters (1988) model fitted to the ETR vs. *E* curves, for the plant *A. thaliana* (circles) and the diatom *N. palea* (squares) grown under high (white), moderate (gray) and low light (black).

Fig. 6. Comparison between the ratios *E*$_{50}$/*E*$_{k}$ (a) and NPQ(*E*$_{k}$)/NPQ$_{m}$ (b) as calculated for the published (excluding cases with impaired XC operation) and experimental (this study) light-
response curves of NPQ and ETR in VAZ and DD-DT types of XC. Bars represent one standard error, boxes represent the lower (25%) and upper (75%) quartiles, and the thin and thick horizontal lines inside the boxes represent the mean and the median, respectively.
Figure 1
Figure 2
Figure 6