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Nitric oxide, substrate of *Euphorbia characias* peroxidase, switches off the CN\(^-\) inhibitory effect

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**A B S T R A C T**

The oxidation of nitric oxide (NO) by *Euphorbia characias* latex peroxidase (ELP-Fe\(^{III}\)), in the presence or in the absence of added calcium, has been investigated. The addition of hydrogen peroxide to the native enzyme leads to the formation of Compound I and serves to catalyse the NO oxidation. The addition of NO to Compound I leads to the formation of Compound II and, afterwards, to the native enzyme spectrum. Under anaerobic conditions, the incubation of the native enzyme (ELP-Fe\(^{III}\) with NO leads to the formation of the stable complex, showing a characteristic absorption spectrum (ELP-Fe\(^{III}\)-NO\(^+\)). The rate of the formation of this complex is slower in the presence of calcium than in its absence, and the same applies to the rate of the formation of Compound II from Compound I, using NO as substrate. Finally, we demonstrate that NO protects ELP from the inactivation caused by CN\(^-\) via a mechanism presumably requiring the formation of an enzyme-nitrosyl cyanide complex.

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1. Introduction

The free radical nitric oxide (NO) is an ubiquitous signalling molecule involved in a large number of physiological processes. While in animals NO is produced from \(l\)-arginine by nitric oxide synthases or not enzymatically (but possible catalysed by globins) from nitrite, NO synthesis appears more complex in plants and it may be arginine-dependent or nitrite-dependent, this latter by enzymatic or non-enzymatic routes. Several original papers and reviews have been published in the last years concerning NO synthesis in plants and it is therefore difficult to quote them all. As examples, we can cite some of these ([1–4] and references therein).

An important aspect of NO-synthesis is the involvement of polyamines inducing the production of NO [5], and the presence of an unknown enzyme responsible for the direct conversion of polyamines to NO has been hypothesised [6].

Peroxidases (E.C. 1.11.1.x; donor: hydrogen peroxide oxidoreductase) are enzymes utilising hydrogen peroxide or other peroxides to oxidise a second reducing substrate. Heme containing peroxidases (E.C. 1.11.1.7) are grouped in major (super)families one of which is a catalase–peroxidase superfamily [7]. This latter group can be subdivided into three classes (I, II and III) on the basis of sequence similarity. Class III includes the secretory plant peroxidases, monomeric glycosylated proteins distributed throughout the plant kingdom [8,9]. These enzymes are implicated in several plant physiological processes, such as hydrogen peroxide detoxification, auxin metabolism, cell elongation, lignin and suberin formation, salt tolerance and oxidative stress. Moreover, when plants are attacked by pathogens, several defence mechanisms are activated and Class III peroxidases play a very special physiological role [10–12].

Two well studied examples of Class III peroxidases are the enzymes extracted from horseradish (HRP; [13] and references therein) and from latex of the perennial Mediterranean shrub *Euphorbia characias* (ELP; [14]). ELP is present in the latex as unique isoenzymatic form constituted by a single glycosylated polypeptide chain of 347 amino acid residues with a relative molecular mass of 47 kDa and contains a ferric iron–protoporphyrin IX pentacoordinated to a proximal histidine ligand. The ELP sequence ([15]; GenBank accession number AY586601) permits identification of two highly conserved histidine residues (His\(_{50}\) and His\(_{179}\), distal and proximal, respectively). Allike other secreted plant peroxidases, ELP has two calcium binding sites namely "proximal" and "distal" but, unlike to these, the purified protein contains only one mol of endogenous Ca\(^{2+}\)/mol enzyme strongly bound to the proximal site. This proximal Ca\(^{2+}\) ion plays a critical role for retaining the active site geometry. A second Ca\(^{2+}\) ion, necessary for expression of the full activity of the enzyme, appears to be located...
at the distal low affinity binding site [16].

The kinetic mechanism of ELP is well established [14,17] and is briefly summarised here. The reaction of hydrogen peroxide with ELP (PrIXFeIII) generates the green enzyme intermediate Compound I (PrIXFeIV = O), with both of the oxidising equivalents of H2O2 transferred to the enzyme. One of the two oxidising equivalents of peroxide is accounted for by the loss of an electron from the iron atom that is oxidised to a ferryl complex in Compound I (FeO = O2−), whereas the second electron is donated by the porphyrin ring, oxidised to a π-cation radical. Compound I then reverts to the resting state by two successive one-electron reactions with reducing substrate molecules (AH2). The red Compound II (PrIXFeIII = O), a second enzyme intermediate, is produced by the first electron transfer from AH2 to Compound I.

We previously reported that calcium ions enhance the catalytic efficiency of ELP toward substrates during the peroxidative cycle [14], reduce ELP activity in the so called catalase-like cycle [18], and regulate the activity of the enzyme to execute different metabolic pathways toward the same substrate, as in the oxidation of tyramine [19] and thiocyanate [20]. In this study we have investigated the binding and the oxidation of NO by Euphorbia peroxidase in the presence and absence of calcium ions. We show that native ELP forms, both in the absence or in the presence of calcium ions, a stable complex with nitric oxide but, in the presence of these ions, the affinity of ELP for NO is lowered by a factor of 3. Moreover, native ELP utilises NO as second substrate by an unusual kinetic model to form Compound II from Compound I and the native enzyme from Compound II. In the presence of calcium added, after addition of NO to Compound I, the Compound II is formed but evolves to a slow process that populates ELP-FeIV-NO− complex. Finally, probably the most interesting and new result, we show that NO protects ELP from the inactivation caused by CN− by a novel mechanism requiring the formation of an enzyme-nitrosyl cyanide complex.

2. Materials and methods

2.1. Materials

2.2.2’-Azinobis(3-ethylbenzthiazoline-6-sulphonic) acid (ABTS), guaiacol, ascorbic acid and diethylenetriamine–NONOate (DEA/NO) were purchased from Sigma (St. Louis, MO). We used DEA/NO for the reliable generation of nitric oxide and we considered 16 min the half-life of DEA/NO (25 °C and pH 7.0) and 1.5 the efficiency of NO release (molecule of NO per molecule dissociates) [21]. Hydrogen peroxide was from Merck (Darmstadt, Germany) and an ε420 = 43.6 M−1 cm−1 was used to determine its concentration. All reagents were obtained as pure commercial products and used without further purification.

2.2. Enzyme

 Peroxidase from E. characias latex (Reinheitszahl (RZ) value A510/A273 = 2.7 in 100 mM Tris–HCl buffer, pH 7.0) was purified as previously described [14]. The enzyme concentration was determined spectrophotometrically using an ε401 = 130.7 mM−1 cm−1.

2.3. Spectrophotometry

Absorption spectra and data from all activity assays were obtained with an Ultraspec 2100 spectrophotometer (Biochrom Ltd., Cambridge, England) using cells with a 1 cm path length.

Anaerobic experiments were made in a Thunberg-type spectrophotometer cuvette (Soffieria Vetro, Sassari, Italy) after several cycles of evacuation followed by flushing with O2-free argon. Anaerobic additions of various reagents to the cuvette were made through a rubber cap with a syringe.

2.4. Peroxidase activity

Activity measurements were performed in 100 mM Tris–HCl buffer, pH 7.0, in both the absence and the presence of 10 mM Ca2+ ions, using hydrogen peroxide and the reducing substrate by the below reported procedures:

(i) ABTS oxidation following the increase in absorbance at 415 nm resulting from the formation of the ABTS cation radical product (ε415 = 36 M−1 cm−1).

(ii) Guaiacol following its oxidation by the increase in absorbance at 470 nm (ε470 = 26.6 M−1 cm−1 [22]).

(iii) NO oxidation by NO electrode (see below).

2.5. NO electrode

Stock solutions of NO (Air Liquide, Paris, France) were prepared by equilibrating degassed water at room temperature with the pure gas at 1 atm. NO measurements were carried out at 25 °C using a NO selective electrode (ISO-NO MarkII World Precision Instruments, Florida, USA). O2 concentration in the assay was also monitored, in parallel, using a high resolution respirometer (Oxygraph-2k, Oroboros Instruments, Innsbruck, Austria). In a typical experiment, NO–equilibrated water was added to the reaction chamber of the respirometer (V = 1.5 mL) containing deoxygenated and N2-equilibrated 100 mM Tris–HCl buffer, pH 7.0. Afterwards, 1.5 μL of ELP (0.1 μM final) and H2O2 were added and the rise and fall in NO concentration was continuously monitored.

The value of Kcat for ELP using varying NO concentrations (1–10 μM) at a fixed concentration of hydrogen peroxide (10 μM), or varying concentrations of hydrogen peroxide (0.1–5 μM) at a fixed concentration of NO (5 μM), was calculated from data fitted to the Michaelis–Menten equation by nonlinear regression and by double reciprocal plots by Michaelis–Menten analysis. The kinetic parameter kcat/Km was defined as (mol of substrate consumed)/(mol of enzyme active sites) × s−1. The kcat/Km value was also used as a measure of substrate specificity. The effects of Ca2+ ions on ELP activity were examined in buffers with or without CaCl2. All kinetic parameters were the mean of three different determinations.

2.6. Stopped-flow experiments

Stopped-flow experiments were carried out using an Applied Photophysics MV 17 apparatus (Leatherhead, UK), equipped with an observation chamber with a 1 cm path length and either a monochromator and a photomultiplier tube (for single wavelength measurements) or a spectrometer and a photodiode array detector (for rapid acquisition of absorbance spectra over the range 250–800 nm). Single wavelength measurements were analysed by a least squares minimisation routine (developed using the Borland Pascal 7.0 compiler) capable of fitting any desired theoretical model to the experimental data, using either analytical or numerical integration.

2.7. Laser photolysis experiments

Laser photolysis experiments were carried out using the instrument previously described ([23,24] and references therein). The instrument allowed different setups of which only one was used for the experiments described below. A pulsed Nd-YAG solid-state laser (model HIL 101, Quanta System, Milano, Italy) capable of 5 ns pulses of 80 mJ at λ = 532 nm, was focused onto an optical Thunberg tube containing the desired solution. The transmittance of the sample was monitored using the stationary light, orthogonal to the laser beam, emitted by a 50-W lamp. The observation wavelength (425 nm)
was obtained using a Spex 1681 monochromator, and a Hamamatsu H6780-01 photosensor module was used to measure the transmitted light. The current emitted by the photosensor was amplified using a Hamamatsu C1053 current to voltage amplifier and read using a digital Tektronix TDS 360 oscilloscope.

3. Results

3.1. NO is a substrate for Euphorbia peroxidase

The addition of NO under anaerobic conditions to 100 mM Tris–HCl buffer, pH 7.0 (5 μM NO final), in a chamber equipped with a NO-selective electrode, showed that the NO signal increased rapidly (Fig. 1, solid line). Addition of hydrogen peroxide (500 μM) to the reaction mixture had no significant effect on the rate of NO consumption. The addition of ELP (0.1 μM) induces a rapid disappearance of NO. Dotted line: ELP (1.5 μM) binds NO causing a decrement in NO concentration. The addition of H2O2 results in the rapid consumption of NO. Buffer 100 mM Tris–HCl, pH 7.0.

3.2. Competitive inhibition mechanism by NO and hydrogen peroxide on the active site of ELP

It is well known that NO inhibits the peroxidase activity [25]. Indeed, as described below, NO binds to the ELP-FeII forming a stable ferrous nitrosyl complex. Thus, NO competes with hydrogen peroxide. To determine the constant of inhibition (Ki) we calculated the inhibition varying the amount of hydrogen peroxide at fixed concentrations of NO, in the presence and absence of 10 mM Ca2+ ions. Since ABTS spontaneously reacts with NO (the oxidation of ABTS occurring at 1:1 stoichiometry with NO [26]), guaiacol was used as second substrate and the initial velocity was determined monitoring the absorption change at 470 nm. We obtained a competitive inhibition with a Ki value, determined by Dixon’s plot, of 15.7 (± 1.1) μM. In the presence of calcium ions a higher Ki value was obtained (300 (± 21) μM; not shown).

3.3. Euphorbia peroxidase (ELP-FeII)-NO complex: spectrophotometric features and stopped flow spectroscopy

The electronic absorption spectrum of native ferric ELP, in 100 mM Tris–HCl buffer, pH 7.0, in the presence or in the absence of 10 mM Ca2+ ions, showed maxima at 278, 401, 498, and 637 nm (Fig. 2). Both in the absence and in the presence of calcium added and in anaerobic conditions, NO induced an increase and a red shift of the Soret band from 401 to 418 nm. Moreover, two well defined peaks at 530 and 565 nm appeared. These spectral features are very similar to those previously reported for HRP [27], and the shape and positions of these bands are typical of low spin ferrous derivative (ELP-FeII–NO+). No further spectral changes were observed after 30 min under anaerobic conditions. After readmission of oxygen, the formation of the original native ELP was observed.

Stopped-flow experiments were used to determine the association (kcat) and the dissociation (kdiss) rates of NO binding to the native ELP in the absence or in the presence of 10 mM Ca2+ ions. Experiments were carried out under anaerobic conditions by rapid mixing of a native ELP solution with 100 mM Tris–HCl buffer, pH 7.0, containing different NO concentrations. The formation of the ELP-FeII–NO complex was monitored following the increase in absorbance at 418 nm. When ferric ELP was mixed with NO, in the absence of calcium, a relatively fast optical change was recorded, leading to a new species identified

![Fig. 1. Effect of ELP and hydrogen peroxide on NO consumption. Solid line: a typical trace recording in a chamber equipped with an NO-selective electrode showing the rapidly NO signal increase. Addition of hydrogen peroxide (500 μM) to the reaction mixture has no significant effect on the rate of NO consumption. The addition of ELP (0.1 μM) induces a rapid disappearance of NO. Dotted line: ELP (1.5 μM) binds NO causing a decrement in NO concentration. The addition of H2O2 results in the rapid consumption of NO. Buffer 100 mM Tris–HCl, pH 7.0.](image-url)
This dissociation constant is much lower than the $K_i$ observed, consistent with the higher reactivity of ELP with $\text{H}_2\text{O}_2$ that is induced by calcium ions [14].

Since the ELP–NO complex is photolabile, the time course of re-binding could also be recorded by flash photolysis using a pulsed laser source for photoexcitation (Fig. 4). The apparent second order rate constants, recorded by this technique in the presence and in the absence of calcium ions, are very similar ($1.15 (\pm 0.2) \times 10^6 \text{M}^{-1} \text{s}^{-1}$ in the absence of calcium and $0.9 (\pm 0.15) \times 10^6 \text{M}^{-1} \text{s}^{-1}$ in its presence).

However, it is interesting to compare the stopped flow data with the empirical and semi-quantitative observation that $\text{O}_2$ converts ELP–NO to ELP at a slower rate if calcium is present. Indeed, if the reaction was simply due to the consumption of free NO by $\text{O}_2$, it should be slower in the absence of calcium ions and faster in its presence, given that these ions lower the affinity of ELP for NO and speed up the dissociation of the complex. Thus, we take this observation as an indication that NO bound to ELP is more prone to react with $\text{O}_2$ and other substances than free NO in solution. Further observations in this direction are reported below.

### 3.4. Reaction of ELP with hydrogen peroxide and nitric oxide: spectrophotometric features and stopped flow spectroscopy

The addition of equimolar amount of hydrogen peroxide to native ELP, both in the absence and in the presence of 10 mM $\text{Ca}^{2+}$ ions, led to the formation of the Compound I, with characteristic absorption maxima at 278, 398, and 651 nm. In the absence of these ions the addition of DEA/NO to the reaction mixture reduced Compound I to Compound II with characteristic absorption maxima at 278, 417, 522 and 555 nm (Fig. 5). Afterwards, Compound II was reduced to the native enzyme after addition of equimolar amount of DEA/NO.

Although it is not easy to determine the relevant concentration of NO when this gas has been released from DEA/NO, repeated experiments, varying concentrations of ELP (2–4 $\mu$M) and/or varying concentrations of DEA/NO, showed that two mol of NO were required to reduce Compound I to the native enzyme (considering 16 min the half-life of DEA/NO and 1.5 the efficiency of NO release; see Section 2). Thus, NO reacted with the oxidised states of the enzyme (Compounds I and II) as an usual substrate (Fig. 6). In the presence of calcium added, after addition of NO to Compound I, we observed a progressive disappearance of the Compound I absorption spectrum, and the gradual formation of the characteristic spectrum of ELP–Fe$^{III}$–NO$^+$ complex (not shown). It may be indicative of a slow process.
the native enzyme. Buffer 100 mM Tris–HCl, pH 7.0.

(b) Compound I is formed after addition of equimolar amount of hydrogen peroxide to the native enzyme. Compound II is formed after addition of approximately 3.7 nmol NO to Compound I. Addition of another amount of NO reduced Compound II to the native enzyme. Buffer 100 mM Tris–HCl, pH 7.0.

c) Compound III (Fig. 6) is formed after addition of equimolar amount of NO to the native enzyme. Compound II is formed after addition of equimolar amount of NO to Compound I. Addition of another amount of NO reduced Compound II to the native enzyme. Buffer 100 mM Tris–HCl, pH 7.0.

that populates contemporarily several intermediates (Compound I, Compound II, native ELP, and ELP-FeII–NO+ complex).

The catalytic cycle of ELP with NO and H2O2 was also followed by stopped flow experiments. However, due to the properties of the reducing substrate, the experiment had some inherent complexities demanding consideration. A straightforward experiment would be realised by mixing ELP-FeIII with a solution containing NO and H2O2 in the desired amounts. However, the long incubation of the two substrates in the driving syringe of the instrument seemed unsafe given the reactivity of both of them. Thus we resorted to two different designs: we either mixed a solution of ELP-FeIII and NO with H2O2 or we mixed a solution of Compound I with NO in the presence of minimal excess of H2O2. In the former case the actual starting material was the ELP-FeIII–NO+ complex, and dissociation of NO rate limited the onset of the steady state condition. The observed time courses were compatible with the rate constant for NO dissociation from ELP-FeIII measured in the absence of H2O2 (see above). The time courses recorded in the absence of calcium (Fig. 7(A)) strikingly differed from those recorded in the presence of this ion (Fig. 7(C)). This result was expected and fully explained by the fact that the reaction of ferric ELP with H2O2 is much faster than the Compound II and ELP-FeII–NO+ complex.

We determined the second order rate constant for the transfer of the first electron from NO to the oxidised enzyme, which resulted 2.0 (± 0.2) × 106 M−1s−1. The reaction product, the Compound II, evolved to the ELP-FeIII–NO+ complex in a much slower process that, under our experimental conditions, appeared to be first order with a rate constant of approximately 1 s−1. Only at concentrations of NO lower than its equilibrium constant the reaction rate slowed down and the absorbance change diminished, due to the incomplete saturation of ferric ELP with the gas. In this case the absorbance spectrum at the end of the reaction was consistent with a mixture of ELP-FeIII and ELP-FeII–NO+ complex. We cannot at present provide a complete explanation of this behaviour but we remark that the observed kinetic process must involve as least two bimolecular processes (one requiring a molecule of NO converting Compound II to the ferric enzyme, the other requiring a further molecule of NO to convert the ferric enzyme to its NO complex) plus one not yet identified monomolecular one.

In the presence of calcium ions (Fig. 8(C) and (D)) the experiment yielded a somewhat more complex time course, since the Compound I present before mixing was not completely converted to Compound II and then to ELP-FeII–NO+, as we expected. Probably, under these experimental conditions, a side reaction occurred, accounting for the (temporary) loss of a fraction of the molecules of the enzyme, that also explains why, under steady state conditions, the Compound I was more populated in the presence of calcium than in its absence. In the latter type of experiment, the starting material was Compound I, which, contrary to the ELP-FeIII–NO+ complex, is an intermediate of the ELP catalytic cycle. In single turnover experiments, realised by rapidly mixing Compound I with excess NO, the presence or absence of calcium ions induced striking differences in the time courses. The case of the experiment carried out in the absence of calcium ions was relatively straightforward (Figs. 8(A) and 7(B)). The Compound I had the lowest absorbance at 418 nm and evolved in two subsequent exponential steps, assigned to the formation of Compound II and ELP-FeIII–NO+. We determined the second order rate constant for the transfer of the first electron from NO to the oxidised enzyme, which resulted 2.0 (± 0.2) × 106 M−1s−1. The reaction product, the Compound II, evolved to the ELP-FeIII–NO+ complex in a much slower process that, under our experimental conditions, appeared to be first order with a rate constant of approximately 1 s−1. Only at concentrations of NO lower than its equilibrium constant the reaction rate slowed down and the absorbance change diminished, due to the incomplete saturation of ferric ELP with the gas. In this case the absorbance spectrum at the end of the reaction was consistent with a mixture of ELP-FeIII and ELP-FeII–NO+ complex. We cannot at present provide a complete explanation of this behaviour but we remark that the observed kinetic process must involve at least two bimolecular processes (one requiring a molecule of NO converting Compound II to the ferric enzyme, the other requiring a further molecule of NO to convert the ferric enzyme to its NO complex) plus one not yet identified monomolecular one.
was recovered over a time regime longer than that explored in our stopped flow measurements.

3.5. Formation of the cyanide derivative: spectrophotometric features of ELP in the presence of CN⁻ and NO

CN⁻ behaves as a competitive inhibitor of ELP since it combines with the ferric heme iron. In 100 mM Tris–HCl buffer, pH 7.0, a Kᵢ value of 1.7 (± 0.2) × 10⁻³ M was determined by Dixon’s plot, using ABTS and H₂O₂ as substrates. When CN⁻ was added to ELP, the absorption band at 401 nm disappeared in parallel with formation of a band at 418 nm, and a new peak at 535 nm appeared. The final spectrum obtained is typical of low-spin derivative six-coordinate inactive complex (Fig. 9).

When an excess of NO was added, in anaerobic conditions, to the ELP–CN⁻ complex, its spectrum immediately changed and the characteristic spectrum of ELP-Fe²⁺–NO⁺ complex appeared. No further spectral changes were seen after 3 h in anaerobic conditions indicating that this complex is very stable. Readmission of oxygen caused the gradual formation of the native enzyme (Fig. 9) instead of the expected ELP–CN⁻ complex, suggesting that cyanide had been consumed in a reaction that involves ELP and NO.

In a successive experiment, even in anaerobic conditions, we added NO to the ELP–CN⁻ complex in a stoichiometric ratio NO:CN 1:5. Again, the spectrum of the complex immediately changed and the characteristic spectrum of ELP-Fe²⁺–NO⁺ complex appeared. This spectrum gradually disappeared after readmission of oxygen with the contemporary formation of the ELP–CN⁻ spectrum (not shown).

4. Discussion

Studying the extent of NO binding and its oxidation by Euphorbia peroxidase, the most interesting and new result is the competition between NO and CN⁻ for ELP–Fe³⁺, the NO switching off the CN⁻ from Fe³⁺ of ELP in anaerobic condition and the ELP–Fe³⁺–NO⁺ complex degraded by cyanide in aerobic conditions.

When considering this reaction mechanism, two important questions arise: (i) how cyanide can react with NO complexed with ELP (since free CN⁻ does not react in any case with free NO), and (ii) what could be the reaction products and the role of oxygen. The first question can be answered suggesting that the NO complexed with ELP is more liable to degradation by CN⁻ than free NO. This is not surprising because, as reported in stopped-flow experiments, we have indications that NO bound to ELP is more prone to react with O₂ and other substances than free NO in solution. For the explanation of the second question, the above reported results induced us to hypothesise a reaction mechanism that operates when ELP is incubated with CN⁻ and NO. We hypothesise that CN⁻ reacts with NO forming an unstable transient complex nitrosyl cyanide (ONCN). The formation of this complex has been well described [28]. The solvolysis of nitrosyl cyanide was expected to form, as the hydrolysis products, CO₂ and NH₃, effectively scavenging equimolar amounts of NO and CN⁻. Due to the difficulty to detect CO₂ in aqueous solution of this complex environment, we searched ammonia as one of the reaction product which was determined from the amount of NADH consumed in the presence of glutamate dehydrogenase and oxoglutaric acid. At the end of the reaction, 1 mol ammonia/mol CN⁻ was detected, thus supporting our hypothesis.

Thus, the formation of ONCN from cyanide and NO is a redox reaction that is expected to release an electron. Thus, it cannot proceed in the absence of an oxidant (i.e. under strictly anaerobic conditions), and the role played by oxygen would be that of an electron acceptor.

The above reported mechanism between ELP, NO and CN⁻ leads us to take in account in vivo for the in vitro observed modulation of ELP activity. As reported [20], CN⁻, formed from the oxidation of the pseudohalide thiocyanate (SCN⁻) by ELP in the presence of H₂O₂ and Ca²⁺ ions, reacts with ELP and blocks the enzyme activity. It is plausible that NO levels, increasing in particular conditions, can protect ELP from the inactivation caused by CN⁻ finely modulating the release of the reaction products.

Besides the main conclusion above reported, several other conclusions are immediately obvious from the experimental findings. NO behaves a substrate for ELP with a very high affinity as detected by experiments with a NO-selective electrode.

The kinetic model for ELP catalysis when NO is used as a second substrate, described in Fig. 6, is compatible with stopped flow and flash photolysis data, but is probably insufficient to describe the complete reaction cycle. Indeed, as one may easily remark, the model predicts inhibition by one of the two substrates (NO). This is compatible with kinetic data but has never been observed in steady state measurements. It is therefore likely that some hitherto uncharacterised
Fig. 10. A variety of biotic and abiotic stresses may elicit the transient increase of NO by both nitrite or arginine pathways. Polyamines, derived from arginine by arginase and arginine decarboxylase (1 in the figure), can increase the NO amount (2 in the figure) [6]. NO can regulate calcium homeostasis by activation of plasma membrane Ca2+-permeable channels [34,35]. The cellular metabolism of polyamines catalysed by Euphorbia amine oxidase (ELAO) generates diacidaldehydes and H2O2 [30]. Scavenging of H2O2 is mediated by a complex network of enzymes, including ELP which can utilise NO as a second substrate generating H2O2, NO− and NO2. NO2 also activates plasma membrane Ca2+ channels. Calcium ions, normally necessary for expression of the full activity of the enzyme, inhibit ELP activity in the presence of NO or SCN− as second substrates. CN−, obtained by thiocyanate oxidation [20], acts as a reversible inhibitor of ELP and NO switches off the CN− inhibitory effect (present paper). NO is an irreversible inhibitor of ELAO [36]. Dashed lines mark positive regulations, dashed lines blocked negative ones. Neither all the reaction products nor the reaction stoichiometries are showed in the figure.

chemical reaction may occur, and should be added to the model, causing degradation of bound NO (e.g. ELP-FeII−NO2 + H2O2 → ELP-FeII + NO2 + H2O2). The hypothesis that coordination of NO to the heme iron increases its reactivity towards an oxidant such as H2O2 is not unlikely and a similar effect has been observed in cytochrome oxidase and its model compounds for Cu-bound NO [29].

The experiments carried out by stopped flow apparatus do not allow a full description of the steady state oxidation of NO, since the rate constant of conversion of Compound II to ELP could not be precisely determined. However, the rate constants measured in this and preceding works [14], compatible with the Michaelis plots recorded potentiometrically, provided that the second order constant for the reaction of Compound II with NO (k3 in Fig. 6) is not lower than 5–10 × 104 M−1 s−1; e.g. simulations carried out with k3 = 3 × 109 M−1 s−1 yields km H2O2 = 1.5 μM (to be compared with the experimental value of 1.6 μM reported in Table 1). The agreement between steady state and stopped flow experiments is therefore reasonable, the main discrepancy being that in the simulated plots some substrate inhibition is predicted, which is not observed experimentally. Further work is required to clarify this point.

E. characias latex could represent an interesting experimental model to study the complexity of plant latex biochemistry considering the scheme of multi-enzymatic interactions taking place in this system. Very little information is available about Ca2+ signalling, NO production and its metabolism, and H2O2 cycling in Euphorbia latex and on the associated enzymes. There are evidences clearly indicating the coexistence of multiple enzymatic activities within the latex-driven system of E. characias involved in NO and H2O2 regulation. We have characterised two main players in this experimental model, namely the H2O2-producing amine oxidase (ELAO) [30] – a copper/quinone-containing enzyme that catalyse the oxidative deamination of diamines and polyamines to aldehyde and ammonia – and a Ca2+-regulated Class III secreted peroxidase (ELP), both probably involved in the activation of plant defence responses and in the homeostasis of H2O2. Moreover, the two enzymes are believed to be implicated in the oxidative burst through the production of H2O2 [31–33]. Thus, based on the results reported here, we propose a model of the interplay between NO, Ca2+ and H2O2 signalling pathways in E. characias latex (Fig. 10). Beyond the references cited in this paper on the production and regulation of NO in plants, two papers deserve attention, the polyamine-induced NO synthesis in plant [6] and the production and cross-talk of NO with calcium signalling [34,35]. As depicted in Fig. 10, these findings further extend our view of latex biochemistry and permit to sketch a more detailed map of some of the multi-enzymatic interactions potentially taking place in this unusual environment.

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References


