provided by Else

Chemistry & Biology 13, 353–364, April 2006 ©2006 Elsevier Ltd All rights reserved DOI 10.1016/j.chembiol.2006.01.009

Involvement of Reactive Oxygen Species and Reactive Nitrogen Species in the Wound Response of *Dasycladus vermicularis*

Cliff Ross,^{1,*} Frithjof C. Küpper,² and Robert S. Jacobs³ ¹ Smithsonian Marine Station at Fort Pierce 701 Seaway Drive Fort Pierce, Florida 34949 ² Scottish Association for Marine Science Dunstaffnage Marine Laboratory Dunbeg Oban, Argyll PA37 1 QA Scotland United Kingdom ³ Interdepartmental Program in Marine Science University of California, Santa Barbara Santa Barbara, California 93106

Summary

We investigated the signaling events involved in the wound response of the marine macroalga *Dasycladus vermicularis*, finding nitric oxide (NO) production in relation to injury. The addition of exogenous H_2O_2 to aliquots of injured algae accelerated the kinetics of NO production in the wounded region. Similarly, the addition of an NO donor caused an increase in detectable H_2O_2 around the site of injury. By wounding or incubating uninjured algae with an NO donor, peroxidase activity was enhanced. Based on the use of selected pharmacological probes, our results indicate that H_2O_2 production involves the upstream activation of signaling events similar to those observed in the physiology of higher plants.

Introduction

As for practically any other organism, the survival of marine algae relies upon chemical defense systems. Aside from the synthesis of secondary metabolites that may act as deterrents of predators, grazers, and pathogens [1–4], the production of reactive oxygen species (ROS) and reactive nitrogen species (RNS) has been shown to have a significant contribution toward the survival of organisms in a variety of contexts [5–9].

The oxidative burst has been defined as the rapid production of ROS in a defensive response to an external stimulus. It was first noted in the human immune system [10] and, much more recently, in higher plants [11, 12]. Such ROS may include superoxide radical (O_2 ·⁻), hydroxyl radical (OH·), hydrogen peroxide (H₂O₂), and singlet oxygen (¹O₂). O₂⁻ and ¹O₂ are relatively short-lived species (with a typical half-life in the microsecond range), depending on the environment [13]. Conversely, H₂O₂ is a longer-lasting species (half-life of hours or days in seawater) that is electrically neutral and able to pass through cell membranes. Due to its longevity, it may reach considerable distances from its site of formation [14].

Recent years have provided a substantial amount of material recognizing the importance of ROS and the emerging role of RNS as defense response compounds, as well as signal transduction agents, in higher plants [5, 15, 16]. Within the last decade, the role of ROS in macroalgal biology, especially in the chemical defense against pathogens, has been emerging. Initially observed upon injury of the red alga Eucheuma [17], oxidative bursts have been reported in both red [18, 19] and brown [20] algae, typically in the defense against bacterial [18, 21] and eukaryotic [21, 22] pathogens. Although substantial genome responses and the activity of many enzymes are known to be affected by ROS and nitric oxide (NO), molecular and biochemical mechanisms of injury responses are poorly understood, and furthermore, many signaling pathways remain elusive.

Within the last decade, the role of NO in plant disease resistance [23], cell death, and DNA fragmentation in Taxus [24], tissue growth [25], stomatal closure [26, 27], and many other aspects of plant life have been described [28, 29]. Despite the wealth of knowledge that is accumulating in terrestrial animals and plants, little is known about the role of NO in marine organisms. Arumugam et al. [30] report NO production in blood cells of the mussel Mytilus galloprovincialis in response to various chemical triggers, including phorbol 12-myristate 13-acetate. Morall et al. [31] used the activity of the NO-generating enzyme, NO synthase (NOS), to develop a biomarker for the health of corals. NOS activity has been associated with thermal stress in populations of zooxanthellae isolated from scleractinian corals [32]. Except for the recent results with the marine diatom Phaeodactylum [33], nothing has been reported about the occurrence or function of NO in free-living marine algae.

Analysis of higher plant models has suggested that the onset of the oxidative burst is regulated by a series of signal transduction events involving an initial elicitation event where putative receptors are activated. In turn, a variety of downstream signaling events occur, involving GTP binding proteins (G proteins), adenylate cyclases, phospholipases, protein kinases, phosphatases, ion channel activation, and the final activation of an ROS enzymatic source.

Most of these findings have been obtained in higher plants, whereas signaling systems are virtually unexplored in marine chlorophytes. We have recently described evidence of a latent oxidative burst involved in the wound repair mechanism in the marine chlorophyte Dasycladus vermicularis [34]. Our previous work emphasized the pharmacological inhibition of a diphenylene iodonium (DPI-)-sensitive, putative NADPH oxidase that is believed to be the major contributor of ROS. We undertook this present study to backtrack systematically and identify the upstream steps involved in ROS production in D. vermicularis once injury is sustained. Furthermore, our additional aims were to investigate whether wounding would result in NO production, to identify the source of NO, to determine if NO had any relationship with ROS production, and, finally, to determine a functional role of NO in relation to injury.

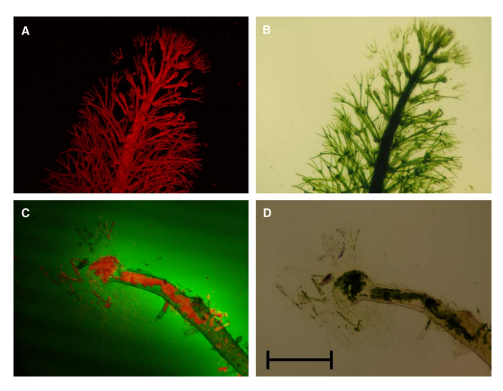


Figure 1. Detection of NO with DAF-FM Diacetate by Confocal Laser Scanning Microscopy

(A) Uninjured specimen with strong visualization of chloroplasts shown in red with no detection of NO (control).

(B) Uninjured specimen displayed under light microscopy (control).

(C) Injured specimen (25 min postinjury) displaying strong NO emission in green.

(D) Injured specimen displayed under light microscopy. Scale bar, 8 mm.

Results

NO Analysis

NO was detected in direct affiliation with injury to *D. vermicularis*. Uninjured specimens displayed no evidence of NO emission. Upon initial mechanical injury, a 5 min lag period occurred before NO could be detected in the wounded region. By 25 min post injury, over 0.15 μ mol NO g⁻¹ FW could be fluorimetrically detected in the surrounding media and visualized by fluorescence microscopy (Figures 1 and 2A). This localized event was more pronounced in adult specimens (25–30 cm) when compared to juveniles (8–10 cm). After 25 min postwounding there was no further increase in the fluorescence emission of the NO signal, suggesting that levels had saturated (~0.20 μ mol NO g⁻¹ FW from 25 to 60 min post injury).

To examine whether NO emission, produced upon injury, originated from NOS, specimens were preincubated with L-NMMA. This NOS-specific inhibitor had a negligible effect on decreasing the NO signal (Figure 2A). Conversely, preincubating healthy algae with the NO scavenger C-PTIO showed a complete elimination of the NO signal once the algae were injured (Figure 2A). In addition, preincubating the injured algae with 100 μ M of the marine natural product pseudopterosin A resulted in complete elimination of the NO signal (data not shown). The elimination of NO had no visual effect on wound plug formation, as described previously [34, 35].

When healthy, noninjured algal specimens were preincubated for 25 min with sodium nitrite, detectible levels of NO were released into the surrounding seawater. Around 0.14 μ mol NO g⁻¹ FW of algae were recorded by 60 min (Figure 2B). By adding KCN prior to nitrite addition, NO production ceased. In addition, when the algae were preincubated with exogenous sodium nitrate, NO was released at concentrations roughly half of that observed when sodium nitrite was used as a substrate (Figure 2B).

Interrelation between NO and H₂O₂

Addition of 100 μ M exogenous H₂O₂ to the injured algae accelerated the initial release of NO from 10 min to 5 min postinjury (0.05 μ mol NO g⁻¹ FW) in the surrounding media (Figure 3A). Not only was the NO signal initially expedited, but NO levels were consistently greater than those levels recorded without the addition of H₂O₂.

The addition of CPTIO (even in addition to the H_2O_2 supplemented experiments) was capable of completely inhibiting the detection of NO (Figure 3A). The preincubation of algae with 5 μ M of the NADPH oxidase inhibitor DPI prior to injury (30 min) resulted in a reduced timeframe of NO production, as shown in Figure 3A.

We have previously reported the relationship between injury and H_2O_2 production in *D. vermicularis* [34]. Just as the addition of exogenous H_2O_2 had an intensifying effect on NO release, the addition of diethylamine NO (DEANO) donor amplified the H_2O_2 release (Figure 3B). Addition of 100 μ M DEANO promoted the initial detection of H_2O_2 from ~25 min to 15 min postinjury. By 45 min postinjury, the DEANO-supplemented experiments displayed H_2O_2 concentrations greater than

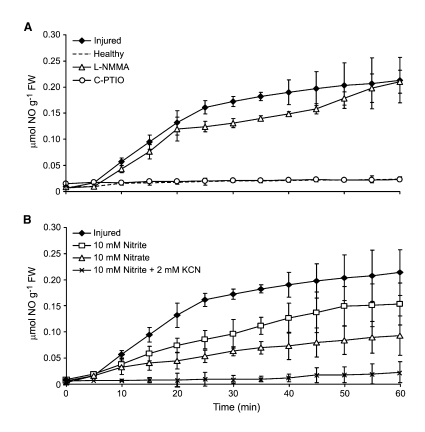


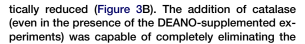
Figure 2. Fluorimetric Quantification of NO Generated Following Injury

The concentration of NO emitted into solution was calculated as µmol NO/g fresh weight as shown in the ordinate. Data points represent the mean ± 1 SEM.

(A) The NO scavenger 2-(4-carboxyphenyl)-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide, potassium salt (carboxy-PTIO) showed complete scavenging of detectible NO, while the NOS inhibitor NG-methyl-L-arginine (L-NMMA) had a negligible inhibitory effect upon NO production (n = 10 individuals).

(B) The additions of 10 mM sodium nitrite and sodium nitrate were both capable as serving as substrates for the nitrate reductase-catalyzed production of NO. Preincubating the algae for 25 min with 2 mM potassium cvanide prior to the introduction of sodium nitrite inhibited the production of NO.

80 μ mol H₂O₂ g⁻¹ FW (Figure 3B). By preincubating the algae for 30 min with 5 μ M DPI prior to injury and subsequently adding DEANO, the H₂O₂ production was dras-



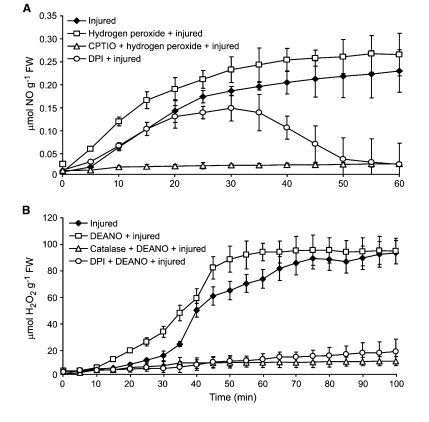
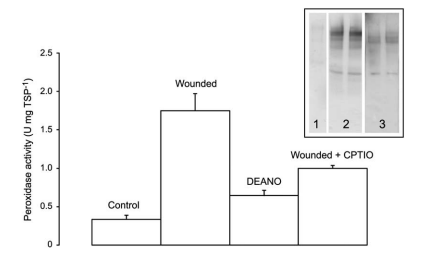


Figure 3. Interrelation between NO and H₂O₂ (A) Upregulation of NO generated by the addition of 100 µM exogenous H₂O₂. The concentration of NO emitted into solution was calculated as µmol NO/g fresh weight. Injured algae supplemented with exogenous H₂O₂ showed an increase in NO production. The addition of CPTIO was capable of completely inhibiting the reaction. Preincubating the algae with DPI prior to injury reduced the time frame of NO production.

(B) Fluorimetric quantification of H₂O₂ generated following the addition of the NO donor DEANO. The concentration of H₂O₂ emitted into solution was calculated as μmol H₂O₂/g fresh weight. DEANO was capable of increasing the signal, yet the addition of catalase or DPI completely eliminated the H₂O₂ signal.

Data points represent the mean ± 1 SEM (n = 10 individuals).



 $\rm H_2O_2$ signal as well (Figure 3B), yet had no effect on NO synthesis.

Activation of Peroxidase Activity by Wounding or NO Supplementation

By wounding the algae or preincubating specimens with DEANO, an increase in peroxidase activity was observed (Figure 4). In spectrophotometric assays by the peroxidase-catalyzed oxidation of o-dianisidine to bis (3,3'-dimethoxy-4-amino) azobiphenyl, wounding caused a 10-fold intensification of detectible peroxidase activity by 300 min, when compared to uninjured algae. The addition of 100 μ M DEANO caused a doubling of peroxidase activity when compared to uninjured algae. Gel activity assays showed the activation of several peroxidase-reactive bands when algae were wounded or when pretreated with DEANO (Figure 4).

Signal Transduction Events Leading to H₂O₂ Release

In this study, we carried out a more detailed investigation of the signaling events that ultimately give rise to an oxidative burst at 40 min postinjury. The protein kinase-c activator phorbol 12-myristate 13-acetate (PMA) was ineffective at enhancing H_2O_2 levels (at the level of accuracy that could be achieved in these experiments) when algae were preincubated in concentrations ranging from 10 to 20 μ M (Figure 5A). However, by preincubating the algae with 50 μ M PMA (prior to wounding), a 25% increase in H_2O_2 production could be detected. Incubating the algae with concentrations of 100 μ M or more resulted in a saturation of detectible H_2O_2 levels (~25% increase above standard levels normally detected after wounding).

To study the possible role of protein phosphorylation in the process of injury recognition and subsequent signal transduction in *D. vermicularis*, we applied staurosporine, calyculin A, and cantharadin. Staurosporine is a potent inhibitor of serine and threonine kinases in animal cells [36]. In *D. vermicularis*, staurosporine was capable of blocking H₂O₂ production with an IC₅₀ of 15.5 μ M (Figure 5B). The protein phosphatase inhibitors calyculin A and cantharidin [37, 38] had no effect on H₂O₂ production. Figure 4. Addition of the NO Donor DEANO Leads to the Upregulation of Peroxidase Activity, Exhibiting a Similar Effect in this Respect as Wounding

Specimens of D. vermicularis were untouched (control), crushed with forceps (wounded), incubated with DEANO for 2 hr, or preincubated with 100 μM carboxy-PTIO for 30 min prior to injury (wounded + CPTIO). Partially purified extracts were assayed for peroxidase activity using the oxidation of o-dianisidine. Data points represent the mean ± 1 SEM (n = 3 groups). Units of peroxidase activity are expressed as U/mg TSP (total soluble protein). The same extracts are shown on an activity gel stained with o-dianisidine. Lane 1 represents uninjured algae. Group 2 represents crushed algae (run in duplicate). Group 3 represents uninjured algae incubated with DEANO (in duplicate).

The anion channel blocker NPPB was capable of partially inhibiting the generation of ROS. Preincubation with 100 μ M NPPB was needed to cause a 15% reduction in ROS detection (Figure 5C).

To determine the relationship between Ca²⁺ channel blocking and H_2O_2 generation, *D. vermicularis* cells were incubated with methoxyverapamil prior to injury. A 20% decrease in H_2O_2 levels occurred in the presence of 100 μ M methoxyverapamil as noted in Figure 5D. Increasing concentrations of methoxyverapamil did not cause an additional decrease in H_2O_2 levels. Concentrations less than 100 μ M had negligible effects.

Mastoparan is an amphiphilic wasp venom tetradecapeptide capable of directly activating pertussis toxinsensitive G proteins by a mechanism analogous to that of G protein-coupled receptors. Mastoparan had no effect on H₂O₂ generation. However, pertussis toxin had a significant inhibitory effect on the oxidative burst, showing an IC₅₀ of 0.17 μ g ml⁻¹ (data not shown).

Antioxidants were capable of completely quenching the detectible H_2O_2 levels. Ascorbic acid was capable of 100% inhibition of the ROS signal, with an IC₅₀ of 20 μ M. Two coumarins with known antioxidant effects were also capable of eliminating the H_2O_2 signal [39, 40]. Umbelliferone (7-hydroxycoumarin) had an IC₅₀ of 50 μ M and esculetin (6,7-dihydroxycoumarin) had an IC₅₀ of 65 μ M (data not shown). Both compounds showed a significant decrease in H_2O_2 production when compared to injured algae incubated without any added compound.

lon flux regulation, in response to injury, was investigated by using a series of specific ion channel blockers, as well as ionophores, to examine their impact on ROS generation. To determine the relationship between Ca²⁺ and H₂O₂ generation, *D. vermicularis* cells were incubated with the Ca²⁺ ionophore A23187 prior to injury. A23187 was capable of enhancing H₂O₂ levels (postinjury) by 35%, as seen in Figure 6. Concentrations ranging from 5 to 50 μ M showed a steady percent increase in H₂O₂ levels. By 50–75 μ M A23187 the effects were clearly saturated.

The K⁺ ionophore, valinomycin, was capable of enhancing H_2O_2 levels (postinjury) from 5 to 75 μ M (Figure 6). By applying 100 μ M, a 10% increase in H_2O_2

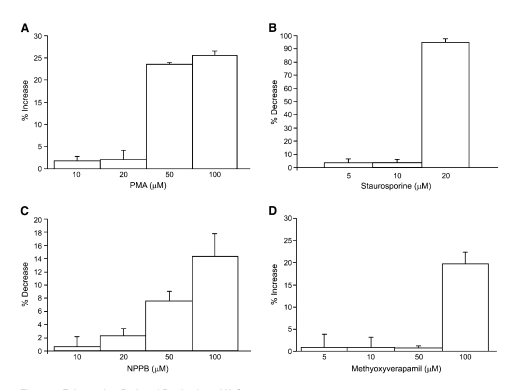


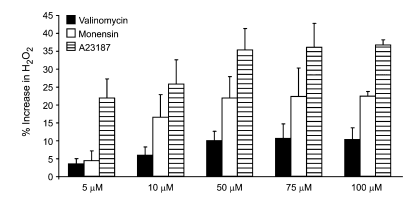
Figure 5. Enhanced or Reduced Production of H_2O_2 Relative percent change of H_2O_2 generated when injured algae were preincubated with (A) the protein kinase-c activator Phorbol 12-myristate 13-acetate (PMA); (B) the protein kinase inhibitor staurosporine; (C) the anion channel blocker NPPB; and (D) the calcium channel blocker methoxyverapamil (n = 10 individuals). Data points represent the mean \pm 1 SEM.

production could be observed. No further percent increase in ROS production was noted with increasing valinomycin concentrations greater than 100 μ M.

The Na⁺ ionophore monensin was capable of enhancing H₂O₂ levels (postinjury) from 5 to 50 μ M (Figure 6). By 50 μ M, a 20% increase in H₂O₂ production could be observed, with the increase leveling off at monensin concentrations around 50 μ M.

O₂ Measurements

Oxygen consumption was monitored for 50 min for healthy (basal respiration) and injured cells. As shown in Figure 7, the increase in oxygen consumption was linear as a function of time postinjury. By 20 min postwounding, the injured algae consumed twice as much O_2 as uninjured cells. By 50 min postwounding, injured



cells showed a relative O_2 consumption rate four times that of uninjured cells.

Discussion

Despite the paramount role of NO in medicine and animal and higher plant physiology, reports of its occurrence in marine algae are scarce and its role remains poorly understood. To our knowledge, this study represents the first finding of NO production in a macroalga and is, together with the recent findings of NO in the marine diatom *Phaeodactylum* [33], among the first in any marine algal lineage.

Fluorescent probes, such as 4,5-diaminofluorescein diacetate (DAF-2 DA) or 2'7'-difluorofluorescein diacetate (DAF-FM DA), have become strong analytical tools for the detection and quantification of NO in higher

Figure 6. Relative Percent Increase of H_2O_2 Generated when Injured Algae Were Preincubated with the K⁺ lonophore Valinomycin, the Na⁺ lonophore Monensin, and the Ca²⁺ lonophore A23187

Data points represent the mean \pm 1 SEM (n = 10 individuals).

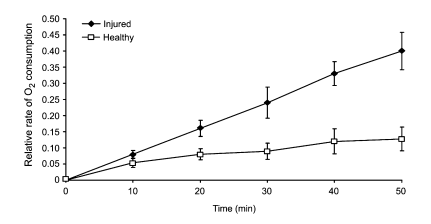


Figure 7. The Relative Rate of Oxygen Consumption per Cell in Healthy and Injured Specimens

Relative rate of oxygen consumption per cell was compared between healthy (uninjured) and injured specimens over a 50 min time course. Data points represent the mean \pm 1 SEM (n = 10 individuals per series).

plants and mammals [41, 42]. This study highlights their usefulness in marine models as well. In mammalian systems, the generation of NO plays a key role in inflammation and tissue repair [43]. It is capable of acting as a cytostatic, chemotactic, and vasodilatory agent during the early wound response; and it mediates cell proliferation, collagen deposition, and angiogenesis, leading to wound closure [44]. The multifunctional roles of NO are ascertained by a series of biochemically diverse redox and additive interactions [45]. We believe that there are strong parallels between the function of NO in algal lineages and that in animals and terrestrial plants.

More and more evidence is supporting the multisignaling functions of H₂O₂ and NO in higher plants. NO can be synthesized during wound responses concomitantly with H₂O₂ production [46, 47]. For example, NO has been shown to act synergistically with reactive oxygen intermediates to increase oxidative cell death in soybean cells [28]. In addition, NO has been shown to inhibit wound-inducible H2O2 generation as well as the downregulation of wound-inducible defense genes in tomato [48, 49]. NO synthesized at the same time as H₂O₂ in response to pathogen attack was found to mediate defense responses similar to those seen following H₂O₂ generation [46]. We used D. vermicularis as a model organism to investigate whether wound-induced NO formation is a conserved response in lower chlorophytes. Additionally, it was of interest to determine whether the production of NO was involved in any synergistic crosstalk with H₂O₂ or had a pertinent functional role with respect to mechanical injury.

In this report, we demonstrate that NO and H_2O_2 are indeed both emitted upon injury to *D. vermicularis*, and that their respective activation appears to be based upon partly coregulated processes. NO emission precedes the oxidative burst (~25 and 45 min, respectively), and furthermore, the two reactive species seem to have an inductive relationship with each other. Reactive oxygen intermediates and NO have been previously shown to have mutually potentiating effects in higher plant cell systems [23]. The addition of the NO donor, DEANO, to incubations of injured *D. vermicularis* caused an increase in H_2O_2 levels and, conversely, adding exogenous H_2O_2 to an analogous experimental setup increased the NO signal detected in wounded *D. vermicularis*.

The detection of low micromolar concentrations of NO is in agreement with relative concentrations detected in soybean suspensions [23] and in tobacco leaves [47]. We stress the term relative, since NO is short-lived due to its high reactivity with many biomolecules. In addition, the ability of a fluorescent probe to bind to released NO does not take into account the NO that is actually trapped within cellular organelles. While the addition of an NO donor may expedite the wound healing process (based upon the promotion of H_2O_2 generation and the upregulation of peroxidase activities), the inhibition of NO formation appears to have no deleterious effect on the immediate wound sealing process; yet we can not exclude that long-term responses may be impacted.

Cytofluorometric analysis of NO and H₂O₂ in oat (Avena sativa) noted an inhibitory effect between the two compounds [48]. Other higher plant studies have noted the mediating role of NO in relation to decrease in catalase and ascorbate peroxidase activity [49]. Our results would support the existing evidence suggesting that a signaling relationship probably exists between NO and H₂O₂ [50, 51]. Our data show that the addition of exogenous NO donor, DEANO, was capable of mimicking the wound response by causing an upregulation in peroxidase activity (Figure 4), suggesting that NO may contribute to coordinating the secondary hardening of the wound plug and other, relatively late, events. The exogenous addition of CPTIO (even when algal samples were wounded) was capable of reducing peroxidase activity when compared to DEANO-treated or injured algae samples, suggesting the involvement of NO in peroxidase upregulation. Heme-based enzymes, such as peroxidase, catalase, dioxygenase, and cyclooxygenases, have metal- and thiol-containing prosthetic groups capable of serving as reaction sites for NO [45, 52]. In addition, nitrogenous compounds, including ammonium ion, nitrosonium ion, and NO free radicals, have been reported to increase heme-based peroxidase activity several-fold in radish [53, 54], human leukocytes [55], and human thyrocytes [56].

In *D. vermicularis*, it is doubtful whether NO synthesis can be attributed to NOS activity, due to the lack of inhibition by L-NMMA. A similar response was noted in tobacco, where the partial inhibition of DAF-2DA fluorescence, by an NOS inhibitor, could only be attributed to the high specificity for an individual NOS isoform [57]. The individual forms of animal and plant NOS activity differ in their degree of sensitivity to NOS inhibitors [58, 59].

Carboxy-PTIO is a highly specific scavenger for NO that is not only membrane-permeable, but capable of

reacting in a stoichiometric manner [60]. Since carboxy-PTIO does not react with ROS, it has been specifically utilized as an inhibitor for NO-related studies. For example, it has been reported to block NO production as well as NO-dependent cell death and defense gene activation in tobacco, soybean, *Arabidopsis*, and barley [23, 57, 58]. While carboxy-PTIO is a useful control to specify NO accumulation, it does not give any indication as to its origin.

Several sources of NO have been described in higher plants. NO may arise from NOS, nitrate reductase, or nonenzymatic sources. NOS catalyzes the NADPHand O_2 -dependent oxidation of L-arginine to NO and L-citrulline [61, 62]. The L-arginine analog, L-NMMA, inhibits NOS in a variety of cells, including neurons, endothelial cells, and macrophages [63]. Several plant proteins have been shown to cross-react with antibodies directed against mammalian NOS [64]; however, only one NOS plant gene (*AtNOS1*) has been identified to date [65].

Studies describing a nitrate reductase-based production of NO have been reported in a diverse set of photosynthetic organisms, including *Helianthus annuus* (sunflower), *Spinacia oleracea* (spinach) [66], and the freshwater microalga *Chlamydomonas reinhardtii* [67]. These reports prompted our investigation to see if this process was conserved in marine green algae.

Studies using C. reinhardtii previously demonstrated that the exogenous addition of sodium nitrite in the presence of nitrate reductase could promote the one-electron reduction to NO [67]. Our results might suggest that a similar process is occurring in D. vermicularis. The enzymatic production of NO was inhibited with potassium cyanide. Although this inhibitor is fairly unspecific, it has been used as a nitrate reductase inhibitor in other biological systems [67, 68]. Interestingly, C. reinhardtii was not capable of using sodium nitrate as a substrate, unlike D. vermicularis. The two-step electron reduction may be more feasible in D. vermicularis, given the high concentration of coumarins that may serve as reducing equivalents. Phenolic metabolites in Hordeum vulgare (barley) have been shown to serve as reducing agents capable of accelerating NO formation [69]. The nonenzymatic production of NO in D. vermicularis should not be completely overlooked. While this type of NO production requires acidic conditions, physical injury results in a mixing of cellular components that may result in conducive physiological conditions (nitritecytosol, acidic pH-vacuole, and reductants-chloroplasts) that may contribute to NO formation [70].

While experimental evidence is increasingly identifying the components of higher plant signaling pathways, the reports of signal transduction systems in macrochlorophytes presently remain scarce. To add to the complexity, the activation of different signaling systems may vary based upon the origin of the sensor and on the nature of the stress event [71–74]. As described in a recent review of abiotic plant stressors, it is currently accepted that there are two groups of membranedependent stress-sensing systems in higher plant cells [74], which we would consider as potential sensors for injury in *Dasycladus*. One group consists of the redox/ H_2O_2 -dependent systems localized in mitochondria, chloroplasts, or peroxisomes, and the other group is based on cell wall-plasma membrane interactions. In regard to the latter, mechanotransduction (physical injury) has a profound effect on the signaling relationships between the cell wall and plasma membrane.

The most widely recognized model for plasma membrane-based ROS production is the NADPH oxidase system. The use of specific inhibitors of NADPH oxidase in plant and animal systems, especially DPI [37, 75], has confirmed the presence of an NADPH oxidase system in various algal lineages [20, 34]. According to the NADPH oxidase model, an elicitor molecule or mechanical pressure is detected by a receptor located on the plasma membrane. In the case of mechanical injury, it appears that a mechanical transducer detects cell wall-plasma membrane perturbations and, in turn, a signal transduction cascade becomes activated. Data available from higher plants support the involvement of GTP binding proteins, ion channels (especially calcium), protein kinases, protein phosphatases, phospholipases A and C, and cyclic AMP. This pathway ultimately gives rise to the activation of an NADPH oxidase complex, which produces superoxide anion. The basis for this reaction is the reduction of molecular oxygen to superoxide anion. This molecule is rapidly dismutated to H₂O₂ in the presence of superoxide dismutase. Our data show that a variety of identifiable signal transduction steps (homologous to what has been reported in higher plants) are believed to be involved in ROS production in D. vermicularis once injury is sustained [14].

Our O_2 consumption data provide evidence that there is a notable uptake of O_2 from the ambient seawater, which would clearly provide adequate levels of oxygen as a substrate for the enzymatically based turnover of O_2 to H_2O_2 . DPI was shown to be very potent in the inhibition of H_2O_2 formation in *D. vermicularis*, as evidenced by the IC₅₀ of 2.4 μ M [34]. Taken together, this inhibition and the finding of increased O_2 uptake during the oxidative burst suggest that *D. vermicularis* may possess a flavoprotein-containing NADPH oxidase complex that is homologous to those involved in the production of AOS in animals and higher plants, and that would reduce molecular oxygen to generate ROS [76, 38].

We have previously reported that quinacrine, an inhibitor of flavin-dependent redox enzymes [77], yielded an IC_{50} value of 5 μ M when H_2O_2 levels were measured in injured *D. vermicularis* [34]. This may indicate specificity toward the flavonoid group of the membrane-associated gp91^{PHOX} subunit of the NADPH-oxidase complex.

The reversible phosphorylation of moieties by kinases and phosphatases is considered to be an integral part of the basic regulatory mechanisms. The general inhibitor of protein kinases, staurosporine, was reported to block the protein phosphorylation events involved in induction of defense responses in tobacco cells upon elicitation by cryptogein [78] or oligogalacturonides [79]. Similar inhibition observations were reported with the inhibition of oligoguluronate-induced oxidative bursts in the brown algal kelp Laminaria digitata [20]. Staurosporine did prove to be an effective inhibitor in the D. vermicularis system, suggesting that serine and threonine kinases are involved in an upstream stage of signal transduction in D. vermicularis. The protein phosphatase inhibitors calyculin A and cantharidin [37, 38] had no effect on the release of H_2O_2 in this system.

Heterotrimeric G proteins play a key role in signal transduction by integrating cell surface receptors to effector systems. Bacterial toxins have been proven to be extremely useful tools for identifying and studying the role of G proteins [80, 81]. The participation of G proteins has been analyzed using cholera toxin in French beans [82]. This same toxin, as well as mastoparan and PMA, was shown (in the bloodroot flower Sanguinaria canadensis) to involve the participation of G proteins and protein kinases, respectively, in the signal transduction for the synthesis of alkaloids induced by fungal elicitors [83]. Pertussis toxin covalently modifies the α subunits of numerous G proteins by ADP-ribosvlating specific amino acid residues [84]. As in L. digitata [20], mastoparan (G protein activator) failed to influence ROS production in D. vermicularis. However, a potent G protein inhibitor, pertussis toxin, proved to strongly inhibit **ROS** production.

Changes in membrane permeability and the resulting ion fluxes, mainly Ca^{2+} and H⁺ influx and K⁺ and Cl⁻ efflux, are among the most rapid responses of plant cells to elicitation [85, 86]. Ion flux regulation in response to stress changes involves the activation of K⁺ and Cl⁻ channels in motor cells in leaves of higher plants [87, 88], as well as the light-dependent ionic movements dictating turgor adaptation involving stomatal function. The multifunctional roles of ion fluxes in plant systems prompted our investigation into the utilization of ion channels as part of the signal transduction process in response to injury in *D. vermicularis*.

An increasing amount of evidence is demonstrating the interrelationship between dependence of ROS production on ion fluxes and the activation of plasma membrane bound Ca^{2+} channels by H_2O_2 [89, 90]. The calcium ionophore A23187 has been used to establish the participation of calcium in the abscisic acid (ABA) transduction system [91], as well as in the hypersensitive response developed in lemon seedlings [92]. Elicitors promote both cytosolic Ca^{2+} increases and ROS production. In some cases, and depending on the species investigated, Ca^{2+} elevations have been reported both upstream and downstream of ROS production [93], indicating complex spatiotemporal Ca^{2+} elevation mechanisms.

The importance of the presence of Ca²⁺ fluxes has been demonstrated in isolated plasma membrane-rich fractions of potato tuber [94]. In this report, activation of the NADPH-dependent O2⁻-generating reaction was strictly dependent upon the presence of Ca2+ ions. Similarly, the emission of H₂O₂ required the increased cytosolic presence of calcium ions in injured D. vermicularis. This was demonstrated by the enhancing effect of the Ca²⁺ ionophore A23187 and the inhibitory effects of the antagonists of Ca2+ channels, such as methoxyverapamil. In Fucus rhizoid cells, DPI inhibited both tip growth and the tip-localized Ca2+ gradient [95]. This infers that the ROS-Ca2+ relationship may represent a more widely used, signal codependence, aside from what has been observed in higher plant physiology. Our findings support the notion that the Ca2+-ROS relationship has a strong signaling role in the wound response in D. vermicularis.

Current knowledge emphasizes the multifunctional roles of ROS and RNS in higher plants and mammalian

systems. Marine algae cover evolutionary lineages that diverged approximately one billion years ago [96]. We have demonstrated that wounding in *D. vermicularis* elicits homologous signal transduction events, suggesting that the cellular machinery to respond to wounding developed early on in the course of evolution.

Significance

These results provide evidence for the presence of nitric oxide (NO) in a marine macroalga, and suggest a role in the rapid wound-healing response in controlling peroxidase activities likely involved in the woundhealing process. Real-time measurements of these compounds can be acquired with fluorescent probes for reactive oxygen species (ROS) and NO. Additionally, pharmacological agonists and specific metabolic inhibitors have been utilized to demonstrate the signal transduction and molecular crosstalk amongst signaling pathways involved in the production of ROS and NO. This experimental evidence clearly demonstrates that Dasycladus vermicularis produces NO in response to wounding, and that there is a potential for feedback amplification between ROS and NO levels that may lead to an accelerated wound-healing response. The initial production of ROS and NO, and potentially the crosstalk between them, rely on signal transduction mechanisms that involve GTP binding proteins, protein kinases, and ion channels. These mechanisms share similarities with signaling pathways in higher plant systems, but they have never before been demonstrated in a marine chlorophyte or other macroalgae, suggesting that key elements in wounding and signal transduction responses are evolutionarily conserved.

Experimental Procedures

Algal Material

Juvenile and adult specimens of *D. vermicularis* (Scropoli) Krasser were collected live off Indian Key, Florida, and Cay Sal Bank, The Bahamas. For laboratory studies, a unialgal culture (strain LB 2685 *D. vermicularis*) was ascertained from the Culture Collection of Algae at the University of Texas at Austin. Cultures were maintained with a photoperiod of 12 hr, with 12 hr of darkness at a photon flux density of 50–60 μ mol m⁻² s⁻¹ and at a constant temperature of 25°C. Muller's medium was utilized in all experiments, and was filtered through 0.22 μ m membranes (no. SCGPT10RE; Millipore, Bedford, MA) prior to introduction of the algae.

Light Microscopy

Wounded and noninjured specimens were visualized with a Leica DMLB compound microscope (Leica Microsystems, Bannockburn, IL) outfitted with a Nikon Cool Pix 8700 camera (Nikon, Melville, NY).

NO Detection

Confocal Laser Scanning Microscopy

DAF-FM DA (Molecular Probes, Eugene, OR) is a cell-permeable probe that passively diffuses across cellular membranes. Once inside the cell, it is deacetylated by intracellular esterases to become DAF-FM. Oxidation of DAF-FM by NO yields a fluorescent benzotriazole derivative [83]. The protocols used in this study were kindly provided by C. Bowler and A. Vardi (Ecole Normale Superieure, Paris): DAF-FM DA was dissolved in DMSO in 5 mM aliquot stocks (stored at -80° C). Individual specimens of *D. vermicularis* were transferred into an incubation mixture of 2 ml filtered seawater and 20 µl DAF-FM DA (final concentration of 20 µM DAF-FM DA). The algae were incubated in the dark for 20 min. In order to detect NO production via

physical wounding, postincubation, the algae were subsequently injured by a single transverse cut with a scalpel and crushed with forceps. The specimens were then washed in 5 ml fresh seawater and subsequently imaged. Confocal laser scanning microscopy (CLSM) was performed with a Nikon Eclipse E800 compound microscope (Nikon Instruments, Kanagawa, Japan) equipped with a Bio-Rad Radiance 2000 laser system (Bio-Rad, Hercules, CA). Laser power was set at 20%, with an excitation of 488 nm and an emission of 525 nm (channel 1, green) or 580 nm (channel 2, red). Series of 0.2 µm optical sections with maximum intensity projection along the z axis were made into one 2D image with greater focal depth. Bio-Rad images were imported into Confocal Assistant 4.02 and converted into TIF files.

Quantitative NO Measurements

The NO release from injured cells was calculated by measuring the oxidation of DAF-FM DA in the presence of esterase (E.C 3.1.1.1; Sigma, St. Louis, MO; 40 U ml⁻¹). Reaction mixtures included 1000 μ l algal extract, 0.40 U esterase, and 20 μ M DAF-FM, for a total volume of 2000 μ l. For NO inhibition or promotion studies, algal samples were preincubated for 30 min with either 100 μ M carboxy-PTIO, L-NMMA, 2 mM potassium cyanide, or 100 μ M DEANO (all stored at -80°C as 100 mM stock solutions dissolved in H₂O). For nitrate reductase detection, solutions of 10 mM sodium nitrate or 10 mM sodium nitrite were preincubated with the algae for 25 min prior to NO analysis. For calculating the concentration of NO present in the samples, calibration with a standard curve was carried out at least once during any series of experiments. Standard curves were composed with known amounts of DEANO sodium salt in addition to 0.40 U esterase and 20 μM DAF-FM DA, for a total reaction volume of 2000 $\mu\text{I}.$ The fluorimetric quantification of NO was analyzed for a 120 min time interval on a Bio-Rad VersaFluor fluorometer.

NO Upregulates Peroxidase Activity

A total of 100 g fresh algae was divided into four treatments: 25 g was crushed with forceps and left undisturbed for 1 hr; 25 g of uninjured algae was incubated with 1 mM of the NO donor, DEANO, for 1 hr; an additional 25 g was preincubated for 30 min with 100 μ M carboxy-PTIO and subsequently crushed with forceps; and, as a control, 25 g of algae was left undisturbed for the same 30 min time frame.

Partial Peroxidase Purification

A partially purified extract of peroxidase was obtained as reported by Ross et al. [34] with slight modification. After wounding, or incubation with an NO donor, 25 g of algae was pulverized into a homogeneous powder with liquid N₂, mortar, and pestle. The solid powder was subsequently mixed (4°C cold room) into 400 ml of 100 mM Tris HCl (pH 8.0) with the addition of 0.1% Triton X-100 and PVPP (6% w/v buffer).

Extracts were centrifuged at 7000 rpm for 20 min at 4°C on a Beckman TJ-6 centrifuge. Supernatants were collected and mixed with 100 mM BaCl₂ for 10 min. Upon centrifugation, under the prior conditions, the supernatant was collected and taken to 30% ammonium sulfate saturation. Upon centrifugation, the supernatant was collected and taken to 90% ammonium sulfate saturation. After the final centrifugation step, the peroxidase-active pellet was collected, resuspended in a minimal amount of Milli-Q water, and dialyzed overnight against several high-volume changes of 1 mM Tris HCI (pH 8.0) at 4°C.

The dialyzed extract was batch loaded onto a High Trap DEAE Sepharose FF 1 ml anion exchanger (Amersham Biosciences) using a Bio-Rad Econopump (Bio-Rad). A step gradient of elution buffers was used, ranging from 100 mM Tris-HCl (pH 8.0) to 100 mM Tris-HCl (pH 8.0) + 1 M NaCl (200 mM increments). Active fractions were pooled, subsequently desalted via dialysis, and then equilibrated against 100 mM acetate buffer (pH 5.5). Samples were concentrated using Centricon Plus-20 5000 NMWL polyethersulfone centrifugal filters (Millipore). Total soluble protein was quantified with the Quick Start Bradford Protein Assay kit (Bio-Rad) according to the manufacturer's instructions.

Peroxidase Activity

Peroxidase extracts of healthy, wounded, and DEANO-supplemented (1 mM) *D. vermicularis* were normalized by protein concentration and assayed for peroxidase activity. Peroxidase extracts (1.45 mg ml⁻¹) were mixed with 5 mM o-dianisidine and 0.5 mM H_2O_2 in acetate buffer (100 mM, pH 5.5), for a total volume of 2 ml. Samples were immediately added to a cuvette and spectrophotometrically assayed for the peroxidase-catalyzed oxidation of o-dianisidine at 460 nm (ϵ_{460} = 11.3 mM⁻¹ cm⁻¹) with a Shimadzu UV-265 spectrophotometer (Shimadzu Biotech, Columbia, MD).

Thirty microliters of normalized samples $(1.032 \text{ mg ml}^{-1})$ were run under nondenaturing SDS-PAGE conditions (10% Tris-HCI Ready Gels; Bio-Rad). Peroxidase activity was detected with a staining solution of 5 mM o-dianisidine and 0.5 mM H₂O₂ in 100 mM acetate buffer (pH 5.5).

Oxidative Burst Measurements

The concentration of H₂O₂ present in the medium around the algal specimens was determined from a protocol previously reported by Ross et al. [34]. Briefly, the H₂O₂ release was calculated by measuring the oxidation of DCFH-DA in the presence of esterase. Stock solutions of H₂DCF-DA (Molecular Probes; 10 mM) and esterase (E.C 3.1.1.1; Sigma; 41 U ml⁻¹) were prepared in DMSO and Muller's medium, respectively. A total of 350 mg (n = 25 cells) of cells was towelblotted dry and weighed. Each cell was then transversely cut in half to make 50 fragments (or 50 exposed injured surfaces). The 50 fragments were placed in a beaker of 50 ml Muller's medium and slowly mixed for a continuous exchange of reagents across the algal surface. For inhibitory experiments, catalase (E.C 1.11.1.6; Sigma; 25 U ml⁻¹) was preincubated with the algae for 30 min prior to injury. Excitation and emission wavelengths were 488 and 525 nm, respectively. Reaction mixtures included 1000 µl algal extract, 0.82 U esterase, and 25 μ M DCFH-DA, for a total volume of 2000 μ l. For calculating the concentration of H2O2 present in the samples, calibration with a standard curve was carried out at least once during any series of experiments. Standard curves were composed with known amounts of H2O2 in addition to 0.82 U esterase, and 25 µM DCFH-DA, for a total reaction volume of 2000 $\mu l.$ The fluorimetric quantification of H₂O₂ was analyzed for a 120 min time interval on a Perkin Elmer LS50B luminescence spectrometer (Perkin-Elmer, Norwalk, CT).

Inhibitors and Activators of H₂O₂ and NO Production

In order to identify upstream signal transduction mechanisms involved in the oxidative burst, selected inhibitors and activators were assayed for their physiological activity. Drugs were preincubated with healthy algae for 30-45 min prior to injury. Diphenyleneiodonium (Sigma) is a suicide inhibitor of NADPH oxidases that binds irreversibly to the flavonoid group of the membrane-associated gp91^{phox} subunit. A 1 mM stock solution was prepared in dimethyl sulfoxide (DMSO). Methoxyverapamil (a Ca2+ channel blocker widely used in medicine, which has also been utilized in plant studies), staurosporine (a protein kinase inhibitor) and NPPB (5-nitro-2-[3phenylpropylamino]-benzoic acid; an anion channel blocker) were dissolved in ethanol (all from Sigma). The ionophores valinomycin (K⁺), monensin (Na⁺), and A23187 (Ca²⁺) (Sigma) were prepared from stocks in ethanol as well. Ascorbic acid, sodium nitrate, sodium nitrite, and potassium cyanide (Sigma) were dissolved in Milli-Q water. The coumarins esculetin (6,7-dihydroxycoumarin; Indofine, Hillsborough, NJ) and umbelliferone (7-hydroxycoumarin; Indofine) were diluted from stocks dissolved in ethanol. Calyculin A, cantharidin (both target protein phosphatases; Sigma), and phorbol 12-myristate 13-acetate (protein kinase c activator; Sigma) were from stocks dissolved in DMSO. Pertussis toxin (a G protein inhibitor, specifically used in the study of adenylate cyclase regulation and the role of G_i proteins; Sigma) was prepared from a water soluble (2 mg/ml) stock solution. Pseudopterosin A (G protein inhibitor) was obtained from our laboratory [67]. The NO scavenger, 2-(4-carboxyphenyl)-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide, potassium salt (carboxy-PTIO), the NO generator (diethylamine NO), and the NOS inhibitor N^G-methyl-L-arginine (L-NMMA) were purchased from Molecular Probes.

O₂ Measurements

Oxygen consumption was measured in healthy and injured specimens with a YSI 5300 Biological Oxygen Monitor (Scientific Division, Yellow Springs Instrument Co., Inc.). Analyzer chambers were filled with 10 ml filtered seawater. A total of 25 uninjured algal cells (~ 0.35 g) were placed in the chamber. O₂ consumption was monitored for 50 min. The same procedure was repeated for injured cells

(same number, with a transverse cut completely through the middle of the siphon). The rate of O_2 utilization was measured as the relative O_2 consumption per cell.

Acknowledgments

We are grateful to George K. Tofaris (University of Cambridge, Cambridge, UK), Assaf Vardi and Chris Bowler (Ecole Normale Superieure, Paris, France), Claudia Moya (University of California, Santa Barbara, California), and Jared Lucas for stimulating discussions, inciting us to carry out this study. We kindly thank Peter Collins and Peter Chaille for assistance with initial fluorescent microscopy experiments. We also gratefully acknowledge funding from the Department of the Army, Grant Award Number PR054175 (R.S.J.) and funding from the Boehringer Ingelheim Fonds to F.C.K., supporting his visit to UC Santa Barbara. This represents Smithsonian Marine Station at Ft. Pierce contribution #647.

Received: July 11, 2005 Revised: January 17, 2006 Accepted: January 20, 2006 Published: April 21, 2006

References

- Lumbang, W.A., and Paul, V.J. (1996). Chemical defenses of the tropical green seaweed *Neomeris annulata* Dickie: effects of multiple compounds on feeding by herbivores. J. Exp. Mar. Biol. Ecol. 201, 185–195.
- Schnitzler, I., Pohnert, G., Hay, M., and Boland, W. (2001). Chemical defense of brown algae (*Dictyopteris* spp.) against the herbivorous amphipod *Ampithoe longimana*. Oecologia *126*, 515– 521.
- Pohnert, G., and Boland, W. (2002). The oxylipin chemistry of attraction and defense in brown algae and diatoms. Nat. Prod. Rep. 19, 108–122.
- Pohnert, G., Lumineau, O., Cueff, A., Adolph, S., Cordevant, C., Lange, M., and Poulet, S. (2002). Are volatile unsaturated aldehydes from diatoms the main line of chemical defence against copepods? Mar. Ecol. Prog. Ser. 245, 33–45.
- Brisson, L.F., Tenhaken, R., and Lamb, C. (1994). Function of oxidative cross-linking of cell wall structural proteins in plant disease resistance. Plant Cell 6, 1703–1712.
- Bolwell, G.P., Butt, V.S., Davies, D.R., and Zimmerlin, A. (1995). The origin of the oxidative burst in plants. Free Radic. Res. 23, 517–532.
- Yahraus, T., Chandra, S., Legendre, L., and Low, P.S. (1995). Evidence for a mechanically induced oxidative burst. Plant Physiol. *109*, 1259–1266.
- Otte, O., and Barz, W. (1996). The elicitor-induced oxidative burst in cultured chickpea cells drives the rapid insolubilization of two cell wall structural proteins. Planta 200, 238–246.
- Razem, F.A., and Bernards, M.A. (2003). Reactive oxygen species production in association with suberization: evidence for an NADPH-dependent oxidase. J. Exp. Bot. 54, 935–941.
- 10. Baldridge, C.W., and Gerard, R.W. (1933). The extra respiration of phagocytosis. Am. J. Physiol. *103*, 235–236.
- Doke, N. (1983). Involvement of superoxide anion generation in the hypersensitive response of potato tuber tissues to infection with an incompatible race of *Phytophthora infestans* and to the hyphal wall components. Physiol. Plant Pathol. 23, 345–357.
- Doke, N. (1983). Generation of superoxide anion by potato tuber protoplasts during the hypersensitive response of potato tuber tissues to infection with an incompatible race of *Phytophthora infestans* and to the hyphal wall components. Physiol. Plant Pathol. 23, 345–357.
- Vranova, E., Inze, D., and Breusegem, F.V. (2002). Signal transduction during oxidative stress. J. Exp. Bot. 53, 1227–1236.
- Wojtaszek, P. (1997). Oxidative burst: an early plant response to pathogen infection. Biochem. J. 322, 681–692.
- Lamb, C., and Dixon, R.A. (1997). The oxidative burst in plant disease resistance. Annu. Rev. Plant Physiol. Plant Mol. Biol. 48, 251–275.

- Orozco-Cardenas, M., and Ryan, C.A. (1999). Hydrogen peroxide is generated systemically in plant leaves by wounding and systemin via the octadecanoid pathway. Proc. Natl. Acad. Sci. USA 96, 6553–6557.
- Collen, J., and Pedersen, M. (1994). A stress-induced burst in Euchema platycladum (Rhodophyta). Physiol. Plant. 92, 417– 422.
- Weinberger, F., and Friedlander, M. (1999). Oligoagars elicit a physiological response in *Gracilaria conferta* (Rhodophyta). J. Phycol. 35, 747–755.
- Bouarab, K., Potin, P., Correa, J., and Kloareg, B. (1999). Sulfated oligosaccharides mediate the interaction between a marine red algae and its green algal pathogenic endophyte. Plant Cell *11*, 1635–1650.
- Küpper, F.C., Kloareg, B., Guern, J., and Potin, P. (2001). Oligoguluronates elicit an oxidative burst in the brown algal kelp *Laminaria digitata*. Plant Physiol. 125, 278–291.
- Küpper, F.C., Müller, D.G., Peters, A.F., Kloareg, B., and Potin, P. (2002). Oligoalginate recognition and oxidative burst play a key role in natural and induced resistance of the sporophytes of Laminariales. J. Chem. Ecol. 28, 2057–2081.
- Bouarab, K., Potin, P., Weinberger, F., Correa, J., and Kloareg, B. (2001). The *Chondrus crispus-Acrochaete operculata* hostpathogen association, a novel model in glycobiology and applied phycopathology. J. Appl. Phycol. *13*, 185–193.
- Delledonne, M., Xia, Y.J., Dixon, R.A., and Lamb, C. (1998). Nitric oxide functions as a signal in plant disease resistance. Nature 394, 585–588.
- Pedroso, M.C., Magalhaes, J.R., and Durzan, D. (2000). A nitric oxide burst precedes apoptosis in angiosperm and gymnosperm callus cells and foliar tissues. J. Exp. Bot. 51, 1027–1036.
- Beligni, M.V., and Lamattina, L. (2001). Nitric oxide in plants: the history is just the beginning. Plant Cell Environ. 24, 267–278.
- Desikan, R., Griffiths, R., Hancock, J., and Neill, S. (2002). A new role for an old enzyme: nitrate reductase-mediated nitric oxide generation is required for abscisic acid-induced stomatal closure in *Arabidopsis thaliana*. Proc. Natl. Acad. Sci. USA 99, 16314–16318.
- Garcia-Mata, C., and Lamattina, L. (2003). Abscisic acid, nitric oxide and stomatal closure: is nitrate reductase one of the missing links? Trends Plant Sci. 8, 20–26.
- Wendehenne, D., Durner, J., and Klessig, D.F. (2004). Nitrix oxide: a new player in plant signaling and defense responses. Curr. Opin. Plant Biol. 7, 449–455.
- Romero-Puertas, M.C., Perazzolli, M., Zago, E.D., and Delledonne, M. (2004). Nitric oxide signaling functions in plant-pathogen interactions. Cell. Microbiol. 6, 795–803.
- Arumugan, M., Romestand, B., Torreilles, J., and Roch, P. (2000). In vitro production of superoxide and nitric oxide (as nitrite and nitrate) by *Mytilus galloprovincialis* haemocytes upon incubation with PMA or laminarin or during yeast phagocytosis. Eur. J. Cell Biol. 79, 513–519.
- Morrall, C.E., Trapido-Rosenthal, H.G., Knap, A.H., and Depledge, M.H. (1998). Development of nitric oxide and nitric oxide synthase as ecotoxological markers in the tropical marine environment. Mar. Environ. Res. 46, 429–432.
- Trapido-Rosenthal, H., Zielke, S., Owen, R., Buxton, L., Boeing, B., Bhagooli, R., and Archer, J. (2005). Increased zooxanthellae nitric oxide synthase activity is associated with coral bleaching. Biol. Bull. 208, 3–6.
- Vardi, A., Formiggini, F., Casotti, R., de Martino, A., Ribalet, F., Miralto, A., and Bowler, C. (2006). A stress surveillance system based on calcium and nitric oxide in marine diatoms. PloS Biology 4 (3), e60.
- Ross, C., Küpper, F.C., Vreeland, V.J., Waite, J.H., and Jacobs, R.S. (2005). Evidence of a latent oxidative burst in relation to wound repair in the giant unicellular chlorophyte *Dasycladus vermicularis*. J. Phycol. 41, 531–541.
- Ross, C., Vreeland, V.J., Waite, J.H., and Jacobs, R.S. (2005). Rapid assembly of a wound plug: stage one of a two-stage wound repair mechanism in the giant unicellular chlorophyte Dasycladus vermicularis (Chlorophyceae). J. Phycol. 41, 46–54.
- Mackintosh, C., and Mackintosh, J. (1994). Inhibitors of protein kinases and phosphatases. Trends Biochem. Sci. 19, 444–448.

- Levine, A., Tenhaken, R., Dixon, R.A., and Lamb, C.J. (1994). H₂O₂ from the oxidative burst orchestrates the plant hypersensitive disease resistance response as a local trigger of programmed cell death and a diffusible inducer of cellular protectant genes. Cell 79, 583–593.
- Jabs, T., Tschöpe, M., Colling, C., Halbrock, K., and Scheel, D. (1997). Elicitor-stimulated ion fluxes and O²⁻ from the oxidative burst are essential components in triggering defense gene activation and phytoalexin synthesis in parsley. Proc. Natl. Acad. Sci. USA 94, 4800–4805.
- Foti, M., Piattelli, M., Baratta, M.T., and Ruberto, G. (1996). Flavenoids, coumarins, and cinnamic acids as antioxidants in a micellar system: structure-activity relationship. J. Agric. Food Chem. 44, 497–501.
- Zai-Qun, L., Yu, W., and Zhong-Li, L. (1999). Antioxidative and prooxidative effects of coumarin derivatives on free radical initiated and photosensitized peroxidation of human low-density lipoprotein. Chem. Phys. Lipids *103*, 125–135.
- Nagano, T., and Yoshimura, T. (2002). Bioimaging of nitric oxide. Chem. Rev. 102, 1235–1269.
- Corpas, F.J., Barroso, J.B., Carreras, A., Quiros, M., Leon, A.M., Romero-Puertas, M.C., Esteban, F.J., Valderrama, R., Palma, J.M., Sandalio, L.M., et al. (2004). Cellular and subcellular localization of endogenous nitric oxide in young and senescent pea plants. Plant Physiol. *136*, 2722–2733.
- Nathan, C., and Shiloh, M.U. (2000). Reactive oxygen and nitrogen intermediates in the relationship between mammalian hosts and microbial pathogens. Proc. Natl. Acad. Sci. USA 97, 8841– 8848.
- Poonawala, T., Levay-Young, B., Hebbel, R.P., and Gupta, K. (2005). Opioids heal ischemic wounds in the rat. Wound Repair Regen. 13, 165–174.
- Stamler, J.S. (1994). Redox signaling: nitrosylation and related target interactions of nitric oxide. Cell 78, 931–936.
- Jih, P., Chen, Y., and Jeng, S. (2003). Involvement of hydrogen peroxide and nitric oxide in expression of the ipomoelin gene from sweet potato. Plant Physiol. *132*, 381–389.
- Huang, X., Stettmaier, K., Michel, C., Hutzler, P., Mueller, M.J., and Durner, J. (2004). Nitric oxide is induced by wounding and influences jasmonic acid signaling in *Arabidopsis thaliana*. Planta 218, 938–946.
- Orozco-Cardenas, M., Narvaez-Vasquez, J., and Ryan, C.A. (2001). Hydrogen peroxide acts as a second messenger for the induction of defense genes in tomato plants in response to wounding, systemin, and methyl jasmonate. Plant Cell *13*, 179–191.
- Orozco-Cardenas, M., and Ryan, C.A. (2002). Nitrix oxide negatively modulates wound signaling in tomato plants. Plant Physiol. *130*, 487–493.
- Neill, S.J., Desikan, R., Clarke, A., Hurst, R.D., and Hancock, J.T. (2002). Hydrogen peroxide and nitric oxide as signaling molecules in plants. J. Exp. Bot. 54, 1237–1247.
- Planchet, E., Gupta, K.J., Sonoda, M., and Kaiser, W.M. (2005). Nitric oxide emission from tobacco and cell suspensions: rate limiting factors and evidence for the involvement of mitochondrial electron transport. Plant J. 41, 732–743.
- Maccarrone, M., Putti, S., and Agro, A.F. (1997). Nitric oxide donors activate the cyclo-oxygenase and peroxidase activities of prostaglandin H synthase. FEBS Lett. 410, 470–476.
- Fridovich, I. (1963). The stimulations of horseradish peroxidase by nitrogenous ligands. J. Biol. Chem. 238, 3921–3927.
- Lee, D.J., Kim, S.S., and Lee, M.Y. (2000). Peroxidase activity boosting by various nitrogenous compounds. J. Biochem. Mol. Biol. 33, 312–316.
- Abu-Soud, H.M., and Hazen, S. (2000). Nitric oxide modulates the catalytic activity of myeloperoxidase. J. Biol. Chem. 275, 5425–5430.
- Millatt, L.J., Johnstone, A.P., and Whitley, G. (1998). Nitric oxide enhances thyroid peroxidase activity in primary human thyrocytes. Life Sci. 63, 373–380.
- Foissner, I., Wendehenne, D., Langebartels, C., and Durner, J. (2000). In vivo imaging of an elicitor-induced nitric oxide burst in tobacco. Plant J. 23, 817–824.

- Durner, J., and Klessig, D.F. (1999). Nitric oxide as a signal in plants. Curr. Opin. Plant Biol. 2, 369–372.
- Ogden, J.E., and Moore, P.K. (1995). Inhibition of nitric oxide synthase: potential for a novel class of herapeutic agent? Trends Biotechnol. 13, 70–78.
- Yoshida, M., Akaike, T., Wada, Y., Sato, K., Ikeda, K., Ueda, S., and Maeda, H. (1994). Therapeutic effects of imidazolineoxyl N-oxide against endotoxin shock through its direct nitric oxide-scavenging activity. Biochem. Biophys. Res. Commun. 202, 923–930.
- Marletta, M.A. (1994). Approaches toward selective inhibition of nitric oxide synthase. J. Med. Chem. 37, 1899–1907.
- Alderton, W.K., Cooper, C.E., and Knowles, R.G. (2001). Nitric oxide sythases: structure, function, and inhibition. Biochem. J. 357, 593–615.
- Abu-Soud, H.M., Feldmen, P.L., Clark, P., and Stuehr, D.J. (1994). Electron transfer in the nitric oxide synthases: characterization of L-arginine analogs that block heme iron reduction. J. Biol. Chem. 269, 32318–32326.
- Ribeiro, E.A., Jr., Cunha, F.Q., Tamashiro, W.M.S.C., and Martins, I.S. (1999). Growth phase-dependent subcellular localization of nitric oxide synthase in maize cells. FEBS Lett. 445, 283–286.
- Zeidler, D., Zahringer, U., Gerber, I., Dubery, I., Hartung, T., Bors, W., Hutzler, P., and Durner, J. (2004). Innate immunity in *Arabidopsis thaliana*: lipopolysaccharides activate nitric oxide synthase (NOS) and induce defense genes. Proc. Natl. Acad. Sci. USA *101*, 15811–15816.
- Rockel, P., Strube, F., Rockel, A., Wildt, J., and Kaiser, W.M. (2002). Regulation of nitric oxide (NO) production by plant nitrate reductase in vivo and in vitro. J. Exp. Bot. 53, 103–110.
- Sakihama, Y., Nakamura, S., and Yamasaki, H. (2002). Nitric oxide production mediated by nitrate reductase in the green alga *Chlamydomonas reinhardtii*: an alternative NO production pathway in photosynthetic organisms. Plant Cell Physiol. 43, 290– 297.
- Forget, P. (1974). The bacterial nitrate reductases: solubilization, purification and properties of the enzyme A of *Escherichia coli* K 12. Eur. J. Biochem. 42, 325–332.
- Bethke, P.C., Badger, M.R., and Jones, R.L. (2004). Apoplastic synthesis of nitric oxide by plant tissues. Plant Cell 16, 332–341.
- Yamasaki, H. (2004). The NO world for plants: achieving balance in an open system. Plant Cell Environ. 28, 78–84.
- Bohnert, H.J., Nelson, D.E., and Jensen, R.G. (1995). Adaptations to environmental stresses. Plant Cell 7, 1099–1111.
- Desikan, R., Mackerness, A.H., Hancock, J.T., and Neil, S.J. (2001). Regulation of the *Arabidopsis* transcriptome by oxidative stress. Plant Physiol. *127*, 159–172.
- Pastori, G.M., and Foyer, C.H. (2002). Common components, networks, and pathways of cross-tolerance to stress: the central role of 'redox' and abscisic acid-mediated controls. Plant Physiol. 129, 7460–7468.
- Kacperska, A. (2004). Sensor types in signal transduction pathways in plant cells responding to abiotic stressors: do they depend on stress intensity? Physiol. Plant. *122*, 159–168.
- Desikan, R., Hancock, J.T., Coffey, M.J., and Neill, S.J. (1996). Generation of active oxygen in elicited cells of *Arabidopsis thaliana* is mediated by a NADPH oxidase-like enzyme. FEBS Lett. 382, 213–217.
- Groom, Q.J., Torres, M.A., Fordham-Skelton, A.P., Hammond-Kosack, K.E., Robinson, N.J., and Jones, J.D.G. (1996). *rbohA*, a rice homolog of the mammalian *gp91phox* respiratory burst oxidase gene. Plant J. *10*, 515–522.
- Auh, C.K., and Murphy, T.M. (1995). Plasma membrane redox enzymes is involved in the synthesis of O₂⁻ and H₂O₂ by *Phytophthora* elicitor-stimulated rose cells. Plant Physiol. 84, 1276–1280.
- Viard, M.P., Martin, F., Pugin, A., Ricci, P., and Blein, J.P. (1994). Protein phosphorylation is induced in tobacco cells by the elicitor cryptogein. Plant Physiol. *104*, 1245–1249.
- Bellincampi, D., Dipierro, N., Salvi, G., Cervone, F., and Lorenzo, G. (2000). Extracellular H₂O₂ induced by oligogalacturonides is not involved in the inhibition of the auxin-regulated *rolB* gene expression in tobacco leaf plants. Plant Physiol. *122*, 1379–1386.

- Moya, C. (2004). *Tetrahymena thermophila* used as a pharmacological model to study the cellular mechanism of action of pseudopterosin A. PhD thesis, University of California, Santa Barbara.
- Mydlarz, L.D., and Jacobs, R.S. (2004). Comparison of an inducible oxidative burst in free-living and symbiotic dinoflagellates reveals properties of the pseudopterosins. Phytochemistry 65, 3231–3241.
- Bolwell, G.P., Coulson, V., Rodgers, M.W., Murphey, D.L., and Jones, D. (1991). Modulation of the elicitation response in French bean cells and its implication for the mechanism of signal transduction. Phytochemistry 30, 399–405.
- Mahady, T.C., Liu, C., and Beecher, C.W. (1998). Involvement of protein kinase and G proteins in the signal transduction of benzophenathridine alkaloid biosynthesis. Phytochemistry 48, 93– 102.
- Moss, J., and Vaughan, M. (1984). ADP-ribosyltransferase that act on adenylate cyclase systems. Meth. Enzymol. 106, 411–418.
- Noh, B., and Spalding, E.P. (1998). Anion channels and the stimulation of anthocyanin accumulation by blue light in *Arabidopsis* seedlings. Plant Physiol. *116*, 503–509.
- Dennison, K.L., and Spalding, E.P. (2000). Glutamate-gated calcium fluxes in Arabidopsis. Plant Physiol. 124, 1511–1514.
- Lee, Y. (1990). Ion movements that control pulvinar curvature in nyctinastic legumes. In The Pulvinus: Motor Organ for Leaf Movement, R.L. Sattler, H.L. Gorton, and T.C. Vogelmann, eds. (Rockville, MD: American Society of Plant Physiology), pp. 130–141.
- Antkowiak, B., and Engelmann, W. (1995). Oscillations of apoplasmic K⁺ and H⁺ activities in *Desmodinium motorium* (Houtt.) Merril. Pulvini in relation to the membrane potential of motor cells and leaflet movements. Planta *196*, 350–356.
- Pearson, G., and Brawley, S.H. (1998). A model for signal transduction during gamete release in the fucoid algae *Pelvetia compressa*. Plant Physiol. *118*, 305–313.
- 90. Murata, Y., Pei, Z.M., Mori, I.C., and Schroeder, J. (2001). Abscisic acid activation of plasma membrane Ca²⁺ channels in guard cells requires cytosolic NAD(P)H and is differentially disrupted upstream and downstream of reactive oxygen species production in *abi1-1* and *abi2-1* protein phosphatase 2C mutants. Plant Cell *13*, 2513–2523.
- Chrispeels, M.J., Holuigue, L., Latorre, R., Luan, S., Orellana, A., Pena-Cortes, H., Raikhel, N.V., Ronald, P.C., and Trewavas, A. (1999). Signal transduction networks and the biology of plant cells. Biol. Res. *32*, 35–60.
- Castaneda, P., and Perez, L.M. (1996). Calcium ions promote the response of *Citrus limon* against fungal elicitors or wounding. Phytochemistry 42, 595–598.
- Bowler, C., and Fluhr, R. (2000). The role of calcium and activated oxygens as signals for controlling cross-tolerance. Trends Plant Sci. 5, 241–246.
- 94. Doke, N., and Miura, Y. (1995). In vitro activation of NADPHdependent O_{2^-} generating system in a plasma membrane-rich fraction of potato tuber tissues by treatment with an elicitor from *Phytophthora infestans* or with digitorin. Physiol. Mol. Plant Pathol. 46, 17–28.
- Coelho, S.A., Taylor, A.R., Ryan, K.P., Sousa-Pinto, I., Brown, M.T., and Brownlee, C. (2002). Spatiotemporal patterning of reactive oxygen production and Ca²⁺ wave propagation in *Fucus* rhizoid cells. Plant Cell *14*, 2369–2381.
- Yoon, H.S., Hackett, J.D., Ciniglia, C., Pinto, G., and Bhattacharya, D. (2004). A molecular timeline for the origin of photosynthetic eukaryotes. Mol. Biol. Evol. 21, 809–818.