Contrasting Regulation of NO and ROS in Potato Defense-Associated Metabolism in Response to Pathogens of Different Lifestyles

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Abstract

Our research provides new insights into how the low and steady-state levels of nitric oxide (NO) and reactive oxygen species (ROS) in potato leaves are altered after the challenge with the hemibiotroph Phytophthora infestans or the necrotroph Botrytis cinerea, with the subsequent rapid and invader-dependent modification of defense responses with opposite effects. Mainly in the avirulent (avr) P. infestans–potato system, NO well balanced with the superoxide level was tuned with a battery of SA-dependent defense genes, leading to the establishment of the hypersensitive response (HR) successfully arresting the pathogen. Relatively high levels of S-nitrosogluthione and S-nitrosothiols concentrated in the main vein of potato leaves indicated the mobile function of these compounds as a reservoir of NO bioactivity. In contrast, low-level production of NO and ROS during virulent (vr) P. infestans-potato interactions might be crucial in the delayed up-regulation of PR-1 and PR-3 genes and compromised resistance to the hemibiotrophic pathogen. In turn, B. cinerea triggered huge NO overproduction and governed inhibition of superoxide production by blunting NADPH oxidase. Nevertheless, a relatively high level of H₂O₂ was found owing to the germin-like activity in cooperation with NO-mediated HR-like cell death in potato genotypes favorable to the necrotrophic pathogen. Moreover, B. cinerea not only provoked cell death, but also modulated the host redox milieu by boosting protein nitration, which attenuated SA production but not SA-dependent defense gene expression. Finally, based on obtained data the organismal cost of having machinery for HR in plant resistance to biotrophs is also discussed, while emphasizing new efforts to identify other components of the NO/ROS cell death pathway and improve plant protection against pathogens of different lifestyles.

Introduction

Generally, both plants and pathogens of different lifestyles have adopted sophisticated strategies to secure their survival and reproduction. Current knowledge and understanding of the ecological and organismal risk of having the machinery for a hypersensitive response (HR)
have shown that necrotrophic pathogens impart an ecological cost on plant resistance to biotrophic pathogens. Studies on ecological costs, defined as resulting from any plant mechanisms that provide resistance to biotrophs or hemibiotrophs, but increasing their susceptibility to necrotrophs, are based on the concept that the plant immune system possesses an ability to self-regulate programmed cell death with reference to a “threshold for the hypersensitive response” [1]. In view of the observation by the cited authors, if the environment is abundant in biotrophic pathogens, plants may be subjected to evolutionary pressure to activate a hypersensitive response at a lower threshold. In contrast, necrotrophs dominant in an environment kill host cells using toxic compounds by overcoming the threshold for the initiation of the hypersensitive response, thus facilitating infection. Evidence has demonstrated that colonization of plants by necrotrophic B. cinerea may largely be prevented by overexpression of anti-apoptotic genes from animals [2]. Moreover, it has been found that the host defense system against a biotroph or hemibiotroph attack might be targeted by toxins of necrotrophic pathogens to enhance virulence and provide susceptibility [3].

Benefits or risks to plants, connected with having the HR mechanism, are frequently associated with a complex of induced compounds and the signaling network that involve many factors affecting stimulation or repression of defense responses against a diverse set of intruders. In the light of these facts, it is crucial to determine factors controlling and modulating a hypersensitive response in host cells [4–7]. Oxidative burst caused by reactive oxygen species (ROS) is the first strong candidate playing an important role in the rapid activation of the genetic program for cellular suicide under different stress conditions [8]. However, most evidence shows that defense responses via ROS, effective against biotrophic pathogens, increase susceptibility to necrotrophic microorganisms (e.g., [1, 9]). Data supporting this concept showed that gray mold development caused by B. cinerea may be suppressed by spraying the plant with an antioxidant [10, 11].

Especially the necrotrophic fungi kill the host tissue via the secretion of a battery of toxins and degradative enzymes. It was found that secretion of oxalic acid by necrotrophs is the main reason for enormous disturbances in the redox status, which correlated with oxalate-induced cell death in plant tissue [12]. In view of the devastating effects of oxalate-secreting B. cinerea, host plants have evolved mechanisms of removing apoplastic oxalate and producing $H_2O_2$ by expressing the enzyme oxalate oxidase [13], also known as germin-like proteins (GLPs). Germin-like gene expression leading to increased resistance to Sclerotinia spp. has been found in rape [14, 15], peanut [16], sunflower [17], tomato [18] and rice [19]. Because GLPs independent of the NADPH oxidases exhibit superoxide dismutase activity [20], they are probably also involved in R-gene mediated and basal resistance to pathogens of different lifestyles.

Nitric oxide (NO) is the next essential signaling molecule in plant immunity generated as an NO burst, occurring rapidly after pathogen recognition in synergy with ROS (e.g., [21–23]). However, a greater number of investigations focusing on gene expression relevant to the HR have suggested that NO and $H_2O_2$ effects may overlap to a great extent [24], while some observations on the barley response to Blumeria graminis indicated that the NO burst preceded $H_2O_2$ synthesis [25]. Pharmacological, biochemical and genetic approaches provide evidence that early overproduction of NO after challenge by a pathogen or an elicitor functions as a messenger in gene-for-gene defense responses and as a general key factor associated with basal resistance in various plant-biotrophic systems (e.g., [26, 27]). Still, much less is known about the role played by NO in the cross-talk between the plant and the necrotrophic pathogen [28–30].

So far, advances in our understanding of plant defense-signaling pathways have shown that plants are capable of recognizing the type of invaders, and they tightly regulate complex defense mechanisms in order to fend off pathogens with various offensive strategies. Both jasmonic acid (JA) and ethylene (ET) have been identified as crucial signaling molecules, effective
in triggering plant responses against pathogens, which require host cell death to feed on the remains. In the case of biotrophs and hemibiotrophs, salicylic acid (SA) signaling probably results in resistance [5, 31–33]. Apart from the opposite effects, there is evidence of extensive cross-talk between SA and JA/ET-dependent signaling pathways, which allow the host plant to maximize repression of the spreading invader and establish resistance not only to a single pathogen, but also a simultaneous invasion by biotrophic and necrotrophic forms [27, 34]. Although this idea is generally true and of greater explanatory value, further evidence suggests a more puzzling picture of these interactions [35]. In the environment containing mainly biotrophic pathogens, the salicylic signaling pathway would be preferable over the JA/ET signaling pathway, and such antagonistic effects may trigger HR-like cell death, thereby facilitating subsequent tissue colonization by necrotrophs [1].

One of the many issues which need to be answered is whether there is only one comprehensive program of boosted NO or superoxide formation evoking positive defense responses against pathogens of different lifestyles with respect to the same plant genotype. Since physiological effects of these reactive compounds are related to their concentration, probably synthesis of reactive nitrogen species (RNS) and ROS may be reprogrammed in an invader-dependent manner to compromise plant resistance. However, in contrast to ROS, our understanding of how NO participates in controlling host cell death remains rudimentary.

Therefore our interest was focused on the analysis of the intensity and kinetics of NO burst and assessment of the importance of NO in triggering defense responses in potato genotypes challenged by a hemibiotrophic (Phytophthora infestans) or a necrotrophic (Botrytis cinerea) pathogen. We attempted to elucidate how RNS have been linked to ROS generation, SA and ethylene signaling compound sensitization, NO-mediated defense responses, e.g. through protein S-nitrosylation or nitration, and PR gene expression leading to HR and/or the collapse of host resistance.

Materials and Methods

Plant growth
Sterile potato plants (Solanum tuberosum L.) of cv. Bintje—(lacking R genes) highly susceptible to isolate 1.3.4.7.10.11. Phytophthora infestans and cv. Bzura—(carrying the R1 gene [36] and the R2-like gene located in potato chromosome IV [37]) highly resistant to 1.3.4.7.10.11. P. infestans were used in the experiments. Potato plants from in vitro tissue culture were transferred to soil and they were grown in a growth chamber with 16 h of light (180 μmol·m⁻²·s⁻¹) at 18 ± 2°C and 60% humidity for 4 weeks.

Pathogen culture
Botrytis cinerea Pers.- isolate 1072 (The Bank of Plant Pathogens, Poznań, Poland) was cultured in the dark at a temperature of 23 ± 2°C, on a potato-agar medium pH 6.3 with a 2% addition of glucose. Fungus culture was restored monthly by mycelium passage onto fresh medium.

Phytophthora infestans—an isolate 1.3.4.7.10.11. was kindly obtained from the Plant Breeding and Acclimatization Institute, Research Division at Młochów, Poland. The oomycete was grown on a cereal-potato medium with an addition of dextrose.

Method of inoculation
For P. infestans inoculation, detached compound leaves of both cultivars were sprayed with a zoospore suspension in water (conc. 2.0 × 10⁵ per ml) on the abaxial leaf surface and kept at 100% humidity in a growth chamber.

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For *B. cinerea* inoculation, potato compound leaves were sprayed with a conidial suspension (conc. $2.0 \times 10^5$ per ml), after which leaves were transferred to the growth chamber at 100% humidity.

The material for further analysis was taken directly or 0.5 h after inoculation (due to technical limitation).

**NO quantification by electrochemical method**

All electrochemical measurements were performed by using an universal electrochemical analyser PGSTAT 30 (EcoChemie, Utrecht, the Netherlands). NO generation in leaf tissue was monitored by differential pulse amperometry with an NO-selective needle-type electrode. The electrode was prepared by electropolymerizing a thin film of poly-eugenol on a cleaned Pt needle. This was done by repetitive scanning of the electrode potential between -0.2 and 0.6 V in 10 mM solution of eugenol (Fluka) in 0.1 M NaOH. The modified electrode was then conditioned by applying a constant potential of 0.9 V in a phosphate buffer (pH 7.4) until a stable constant background current was reached. Electrochemical monitoring of NO generation in leaf tissue after pathogen inoculation was performed as described earlier [29]. The current was recalculated into concentration units on the basis of a calibration curve.

**TUNEL assay**

The TUNEL assay measures DNA fragmentation using the terminal deoxynucleotidyl transferase (TdT)-mediated deoxyuridine triphosphate (dUTP) nick end labelling method, which involves the TdT-mediated addition of fluorescein-12-dUTP to the 3'-OH ends of fragmented DNA (TUNEL fluorescein; Roche, Indianapolis, IN, USA). Thin slices of the surface of leaf tissue were removed with a bistoury and immediately immersed for 1 h in 4% formaldehyde in phosphate buffered saline (PBS). After being rinsed in PBS, the samples were treated with liquid nitrogen. The samples were rehydrated and incubated for 2 min in Triton X-100 solution (0.1% Triton X-100 in 0.1% sodium citrate) on ice. After rinsing the samples in PBS, the TUNEL reaction was performed according to the manufacturer’s protocol (Roche, http://www.roche.com). Negative controls were conducted in the absence of the TUNEL enzyme. In positive controls, tissue was incubated with DNase I (Roche) for 10 min at 25°C, prior to labelling. The samples were examined using a fluorescence microscope (Axiostar plus, Carl Zeiss, Jena, Germany) equipped with a digital camera, with excitation at 488 nm and emission at 515 nm. Experiments were repeated at least four times with ten slices per treatment. A region of 100 epidermal cells from at least 5 randomly selected slides in each treatment was counted and statistically analysed (results not presented).

**Measurement of ROS**

The level of $O_2•^−$ was assayed spectrophotometrically based on the capacity of the superoxide anion-radical to reduce NBT to diformazan [38]. The concentration of $H_2O_2$ was assayed spectrophotometrically using the titanium (Ti$^{4+}$) method [39].

**Detection of peroxynitrite**

The level of peroxynitrite was assayed using folic acid as the peroxynitrite scavenger, giving high fluorescent emission products [40]. Leaf discs (1 g) were immersed in the incubation mixture, containing barbital buffer solution (pH 9.4) and folic acid ($1.0 \times 10^{-5}$ mol L$^{-1}$). Fluorescence intensity of the solution was recorded at 460 nm with the excitation wavelength set at
380 nm. The standard curve was prepared for 3-Morpholinosydnonimine (SIN-1) from Calbiochem as a donor of peroxynitrite at the range of concentration from 1 to 14 nM.

**NADPH oxidase and germin-like oxidase activities assay**

The NADPH dependent \( \text{O}_2^{\cdot-} \) generating activity was determined by a modified assay based on a reduction of XTT by \( \text{O}_2^{\cdot-} \) anions [41]. Fresh leaves (0.5 g) were homogenized in 50 mM potassium-phosphate buffer, pH 7.0 (1:4; w/v), containing 0.1% Triton X-100 (v/v), 1% PVP, 0.04% \( \text{Na}_2\text{S}_2\text{O}_5 \), 1 mM EDTA and centrifuged at 18,000 × g for 20 min. Supernatants were passed through Sephadex G-25 gel filtration columns (Illustra NAP-10, GE Healthcare) and served as the enzyme extract. The volume of 1 ml assay reaction mixture contained 0.5 mM XTT, 0.1 mM NADPH and 30 μl enzyme extract in 50 mM Tris–HCl buffer, pH 7.5. XTT reduction was determined at 470 nm and rates of \( \text{O}_2^{\cdot-} \) generation were calculated using an extinction coefficient for XTT of 2.16 × 10⁴ M⁻¹ cm⁻¹ and the enzyme activity was expressed as \( \text{O}_2^{\cdot-} \) per 1 min per 1 mg protein.

Germin-like oxidase activity was assayed using the method as described [42]. Fresh leaves (0.5 g) were homogenized with cold distilled water in 1:3 ratio (w/v) and centrifuged at 15,000 × g for 30 min. The reaction mixture contained 0.05 M sodium succinate buffer (pH 4.0), 10 mM CuSO₄ and enzyme extract (100 μl). The reaction was started by adding 100 mM oxalate. After incubating at 37°C for 10 min, the color reagent (50 mg 4-aminophenazone, 100 mg solid phenol and 1 mg horseradish peroxidase per 100 ml of 0.4 M sodium phosphate buffer, pH 7.0) was added and incubated at room temperature for 15 min. Amount of \( \text{H}_2\text{O}_2 \) generated in the assay was calculated using standard curve at 520 nm. One unit of enzyme was defined as the amount of enzyme required to catalyze the generation of 1 nmol of \( \text{H}_2\text{O}_2 \) from oxalate per 1 min per 1 mg protein.

**Quantification of total SNOs**

Total SNO content was determined by chemiluminescence using a Sievers ® Nitric Oxide Analyzer NOA 280i (GE Analytical Instruments, Boulder, CO, USA) according to the procedure proposed earlier [43, 44].

**Immunohistochemical studies of SNOs, GSNO and nitrotyrosine**

Detection of SNOs by fluorescence microscopy was performed using the fluorescent Alexa Fluor 405 Hg–Link reagent phenylmercury [45] with some modifications. Potato leaf segments were incubated at 25°C for 1h in darkness with 10 mM NEM prepared in ethanol, and then were washed three times in 10mM Tris–HCl buffer, pH 7.4, for 15 min each. Then they were incubated in the dark with 10μM Alexa Fluor 405 Hg-link phenylmercury (Molecular Probes, USA) for 1h at 25°C. After being washed three times in 10mM Tris–HCl buffer the sections were analyzed with an epifluorescence microscope (Axiostar plus, Zeiss) equipped with an AxioCam MRc 5 camera (Zeiss) using standard filters for Alexa Fluor 405 blue fluorescence (excitation 401nm; emission 421nm). For background staining control sections were incubated with \( \beta \)-mercaptoethanol plus Alexa Fluor 405 and without NEM.

Immunohistochemical localization of GSNO and 3-nitrotyrosine was carried out by fluorescence microscopy of immunofluorescence-stained potato leaf segments using an anti-GSNO rat polyclonal antibody (1:1500) and a rabbit polyclonal antibody against nitrotyrosine (1:300), respectively.

**Ethylene production assay**

Leaf discs (diameter = 1.0 cm) were mock inoculated (control) or inoculated with pathogens as described earlier. After incubation, 10 leaf discs were placed in a 20 ml glass vial closed with a
crimp top cap with a Teflon/silicon septum. Ethylene was extracted by the SPME (solid phase microextraction) technique with Carboxen/PDMS SPME fibre (Supelco, Bellefonte, PA, USA). The vial septum was pierced with an SPME needle and the fibre was exposed to the headspace over analysed leaves for 10 min. After that time the fibre was retracted into the SPME needle and transferred to the GC injection port, where it was exposed to desorbed volatiles adsorbed on the SPME fibre. The desorption process lasted 10 min, and after that time the fibre was used to sample the next vial. Sampling was performed at 20°C. The gas chromatograph (GC) used for analyses was a Hewlett Packard HP6890 equipped with a split/splitless injector, a flame ionization detector, and a PLOT Al₂O₃, KCl (50 m × 0.32 mm × 8 μm) column (19091P-M15, Agilent Technologies, Palo Alto, CA, USA). Helium was used as a carrier gas at a constant pressure of 20 psi. To facilitate sample comparison, amounts of emitted ethylene expressed in nl per 1h per 1 g of FW. All samples were run in triplicate.

**Determination of salicylic acid level**

Measurement of SA content essentially followed the protocol described earlier [46]. HPLC analysis was performed on the Agilent HP 1100 HPLC system, equipped with a photodiode array detector (DAD). For all separations a Lichrospher 100 RP18 column (250.0×4.0 mm, 5.0 μm) from Merck was used. The mobile phase consisted of water, methanol, acetic acid (69:28:3, v/v/v), applied in the isocratic elution. The flow rate was adjusted to 1.0 ml/min, the detection wavelength set to DAD at λ = 302.0 nm and 40.0 μl of samples was injected. All separations were performed at the temperature of 45°C. Peaks were assigned by spiking the samples with the standard compound and a comparison of the UV-spectra and retention times (tₚ = 8 min). The analytical method was validated according to ICH guidelines. The relationship between the peak area and the concentration of salicylic acid injected was found to be highly linear with a regression coefficient R² 0.9999. Intra-day repeatability of the HPLC-DAD method was evaluated by performing five repetitive analyses of the standard, which gave an RSD 2 of 15%, showing good precision.

**Defense gene expression**

RNA was isolated from 0.2 g of frozen leaf tissues using TriReagent (Sigma). The obtained RNA was purified with the use of a Deoxyribonuclease Kit (Sigma). For the reverse transcription 1 μg of RNA from every experimental variant was processed with a Reverse Transcription Kit (Thermo Scientific Fermentas) according to the manufacturer’s instructions. Real-time PCR was performed on a RotorGene 6000 Thermocycler. The reaction mixture contained 0.1 μM of each primer, 1 μl of 5 x diluted cDNA, 10 μl of the Power SYBR Green PCR Master mix (Applied Biosystems) and DEPC treated water to the total volume of 20 μl. The real-time PCR reaction conditions included an initial 5-minute denaturation at 95°C, followed by 55 cycles consisting of 10 s at 95°C, 20 s at 55°C and 30 s at 72°C. The reaction was finalized by denaturation at a temperature rising from 72°C to 95°C by 1°/5 seconds. Reaction specificity was confirmed by the occurrence of one peak in the melting curve analysis. The data were normalized to two reference genes encoding the elongation factor (ef1α, AB061263) and 18S RNA (X67238). All used primers are presented in Table 1. The Ct values were determined with the use of a Real-time PCR Miner [47] and the relative gene expression was calculated with the use of efficiency corrected calculation models [48].

**Assessment of disease index**

The area affected by disease symptoms was assessed on potato leaves 7 days after inoculation with *P. infestans* or *B. cinerea* based on a scale from I to IV [49], which represented the
percentage of leaf area covered by disease symptoms (I = 1–9%; II = 10–24%; III = 25–49%; IV = 50–100%).

Statistical analysis

All results are based on three independent experiments, each with at least three biological replicates. For each experiment, means of the obtained values were calculated along with standard deviations. The analysis of variance was conducted and the least significant differences (LSDs) between means were determined using Tukey’s test at the level of significance $\alpha = 0.05$.

Results

NO generation and transfer of NO bioactivity to potato defense system

Nitric oxide was generated early, within one hour after challenge with a hemibiotrophic or necrotrophic pathogen in both potato genotypes (Fig 1A and 1B). In the interface between cv. Bzura and *B. cinerea* a rapid and high NO burst was observed that reached a maximum (10-fold increase) after 1 hpi of the challenge, and it was followed by a prominent second burst.
of NO generation (14-fold increase) at 3 hpi in comparison to the mock-inoculated control leaves (Fig 1A). Avr P. infestans induced the program of boosted NO generation at a considerably lower level (5- to 7-fold increase) than in the necrotroph-dependent NO formation. In turn, an approximately 4-fold rise in NO production was found in the potato Bintje–B. cinerea interaction, in which it was significantly more abundant when compared to the virulent (vr) P. infestans–potato response (Fig 1B).

To examine the NO bioactivity transferred to NO-affected proteins we localized total S-nitrosothiols in potato leaves using a fluorescent probe, Alexa Fluor 405 Hg-link (Fig 2A–2C and 2G–2I). The most intensive fluorescence related to the cellular S-nitrosothiol (SNO) pool was observed in the avr P. infestans–potato interaction, as well as in the B. cinerea–genotype Bzura interaction at 48 hpi (Fig 2B and 2C). A weak ring of blue fluorescence attributable to SNOs was only observed in the vascular tissue of the petiole in non-infected leaves of cv. Bintje and Bzura (Fig 2A and 2G).

Independently, based on a precise chemiluminescence method, it was found that P. infestans treatment enhanced the total SNO production approximately 3-fold and more than 7-fold at 3 and 24 hpi, respectively, mainly in leaves of the resistant cv. Bzura. For comparison, a 4.5-fold rise in the SNO pool was observed at 24 hpi in leaves of the same genotype infected by B. cinerea (Fig 2I). By contrast, no drastic overproduction was recorded in the susceptible genotype Bintje in response to both hemibiotrophic and necrotrophic pathogens. Immunolocalization of S-nitrosoglutathione (GSNO) with a specific antibody also revealed the most abundant orange fluorescence in the vascular and phloem tissues of potato in response to avr P. infestans, while it was much lower in the Bzura–B. cinerea interaction (Fig 2D–2F).

Analyses of peroxynitrite formation showed that both potato genotypes challenged with B. cinerea produced more peroxynitrite than in the case of P. infestans (Fig 3A and 3B). It was correlated with an immunoassay of nitrated proteins using an antibody against 3-nitrotyrosine (Fig 3C–3F). Intensive orange fluorescence related to highly nitrated proteins was observed in the whole cross-section of petiole and potato leaves infected mainly by B. cinerea (Fig 3D and 3F) when compared to P. infestans (Fig 3C and 3E). Similarly, a relative quantification of the microscopic images revealed the highest intensity of orange illumination in potato challenged with B. cinerea, in contrast to the lowest intensity of fluorescence recorded in the avr P. infestans–potato system (Fig 3G).

ROS generation and enzymes involved in their production

Inoculation of resistant potato with avr P. infestans caused an early and significant rise in superoxide generation starting from 0.5 hpi with a peak at 6 hpi (5-fold increase) (Fig 4A). In susceptible potato leaves challenged with vr P. infestans we found a constant 2-fold higher level of the superoxide at the successive time points after pathogen treatment (Fig 4B). In turn, superoxide production was blocked during the first 24 hpi in leaves of both potato genotypes infected with B. cinerea (Fig 4A and 4B).

To explore whether superoxide managed by NADPH oxidase might be regulated by the NO level, activity of this enzyme was determined in infected potato. Challenge with avr P. infestans caused an early increase in NADPH oxidase activity starting from 0.5 hpi in resistant potato (Fig 4C). In contrast, B. cinerea suppressed this enzyme activity during the first day after the necrotroph attack both in the Bzura and Bintje genotypes (Fig 4C and 4D). In turn, treatment of susceptible potato leaves with the virulent P. infestans showed a weak increase in NADPH oxidase activity only at 6 hpi (Fig 4D).

Next we investigated whether the total H$_2$O$_2$ level was also differentially modified by potato exposure to hemibiotroph and necrotroph pathogens. Unexpectedly, a massive rise (3-fold at 3}
Fig 2. Effect of hemibiotroph (*P. infestans*) or necrotroph (*B. cinerea*) inoculation on S-nitrosothiols in potato. Total contents of S-nitrosothiols (SNOs) in leaves of potato (A-C) cv. Bzura and (G-I) cv. Bintje inoculated with *P. infestans* or *B. cinerea* at 48 hpi. Detection of SNOs in potato leaves was performed by immunofluorescence histochemistry using Alexa Fluor 405 Hg–Link reagent phenyl mercury; blue fluorescence is attributable to SNOs. Images of leaf cross section show general phenomena representative of three individual experiments. Representative images illustrating the immunodetection of endogenous GSNO in (D) control leaves of potato cv. Bzura and in leaves inoculated with (E) *P. infestans* or (F) *B. cinerea*. (J) Total SNO content in leaves of potato determined by chemiluminescence assay using Sievers Nitric Oxide Analyzer NOA 280i; (*) significantly different from mock-inoculated potato leaves, *P*<0.05. Values represent the mean ± SD of at least three independent experiments. (K) Potato leaf cross section viewed under white light: 1 –palisade mesophyll, PM; 2 –spongy mesophyll, SM; 3 –parenchyma, P; 4 –xylem, Xm; 5 –epidermal cells, Ep. Bars indicate 200 μm (A-C, G-I, K) and 100 μm (D-F).

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(hpi) of H$_2$O$_2$ was found in cv. Bzura after challenge with B. cinerea, and a comparable level was also found in cv. Bintje at the same time after inoculation (Fig 5A and 5B). Likewise, the oomycete P. infestans induced early H$_2$O$_2$ synthesis in potato leaves, much more pronounced in the resistant than the susceptible response. Furthermore, the constitutive level of H$_2$O$_2$ was 2-fold lower in cv. Bintje than in cv. Bzura (Fig 5A and 5B).

When examining the effect of pathogen infection on germin-like activity producing H$_2$O$_2$ it was found that GLP activity also increased temporarily within 0.5 to 3 hpi after the avr hemibiotroph or necrotroph treatment (Fig 5C). Generally, early induction of GLP activity (at 3 hpi) was higher in the B. cinerea–potato interaction than in the avr P. infestans–potato one, and it was more pronounced than in the susceptible responses (Fig 5D).

These data imply that the pathogen-mediated early ROS production is largely dependent on its lifestyle. Shortly after inoculation, the necrotrophic pathogen reduced superoxide production in potato, in contrast to the hemibiotroph. In the resistant potato–P. infestans interaction, enhanced ROS production was observed via time-dependent activation of NADPH oxidase.

**Fig 3. Peroxynitrite formation and tyrosine nitration in potato leaves challenged with P. infestans or B. cinerea.** Peroxynitrite content in potato challenged with P. infestans or B. cinerea, respectively, in (A) cv. Bzura and (B) Bintje. (*) significantly different from mock-inoculated (control) potato leaves, P<0.05. Values represent the mean ± SD of at least three independent experiments. (C-F) Representative images illustrating microscopic localization of protein 3-nitrotyrosine in leaf cross-sections of both potato genotypes challenge inoculated with the pathogens at 24 hpi. Bars indicate 200 μm (C-F). (G) Signal intensities were quantified using Multi Guide V2.2 software (Fujifilm) and expressed in arbitrary units.
Though the necrotroph temporarily blunted NADPH oxidase and superoxide generation, potato leaves restored $\text{H}_2\text{O}_2$ production owing to germin-like activity.

**Components involved in NO and ROS defense-associated metabolism**

To understand how both pathogens change the host redox status and the associated signaling mechanism we next analyzed SA production and SA-dependent defense gene expression. The obtained data indicate that free SA synthesis was mainly triggered by *P. infestans* (Fig 6A and
6B). Inoculation with avr *P. infestans* induced a systemic rise in the free SA level from 0.5 to 48 hpi (Fig 6A), while the virulent *P. infestans* significantly potentiated SA synthesis after 6 hpi (Fig 6B). In contrast, *B. cinerea* constantly antagonized SA production in both potato genotypes (Fig 6A and 6B).

Evidence is accumulating that ET together with jasmonic acid plays an important role in the signaling network, creating resistance against necrotrophs; therefore ET production was also detected. *Botrytis cinerea* triggered enhanced ET synthesis peaking at 3 hpi, with similar intensity and kinetics in both potato genotypes (Fig 6C and 6D). The response of susceptible potato

**Fig 5.** Effect of hemibiotroph *P. infestans* or necrotroph *B. cinerea* inoculation on hydrogen peroxide production and germin-like oxidase in potato. (A, B) Hydrogen peroxide accumulation and (C, D) germin-like oxidase activity in potato leaves of cv. Bzura and cv. Bintje. (*) significantly different from mock-inoculated (control) potato leaves, *P*<0.05. Values represent mean ± SD of at least three independent experiments.

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to *P. infestans* did not reveal ET production (Fig 6D), while the resistant response showed an incidental rise in ET activation only at 3 hpi (Fig 6C).

In order to clarify how redox-based changes induced by pathogens targeted the *PR* genes, we analyzed the amount of mRNA coding for PR-1, PR-2 and PR-3. In the *avr P. infestans*–potato interaction the highest amount of mRNA (over 15-fold increase) coding for PR-1 was observed starting from 1–6 hpi, together with increased (over 25-fold) expression of the *PR-2* gene coding for glucanase at 3 hpi (Fig 7A and 7C). Leaves of the susceptible potato cultivar showed strong induction of genes encoding *PR-1* and *PR-3* relatively late (at 24 hpi) after...
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A

**Bzura**
- control
- *avr P. infestans*
- *B. cinerea*

**PR1**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24

B

**Bintje**
- control
- *avr P. infestans*
- *B. cinerea*

**PR1**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24

C

**Bzura**
- control
- *avr P. infestans*
- *B. cinerea*

**PR2**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24

D

**Bintje**
- control
- *avr P. infestans*
- *B. cinerea*

**PR2**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24

E

**Bzura**
- control
- *avr P. infestans*
- *B. cinerea*

**PR3**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24

F

**Bintje**
- control
- *avr P. infestans*
- *B. cinerea*

**PR3**

relative gene expression [fold]

0.5 1 3 6 24

relative gene expression [fold]

0.5 1 3 6 24
virulent \textit{P. infestans} infection (Fig. 7B and 7F). In turn, \textit{B. cinerea} up-regulated PR-1 and PR-2 mainly in potato cv. Bzura (1–3 hpi), but in comparison to the hemibiotrophic pathogen the level of \textit{PR} gene expression was lower and it occurred later after necrotroph inoculation.

Contrasting functions of induced cell death in potato defense strategy against hemibiotrophic and necrotrophic pathogens

High cellular NO and H$_2$O$_2$ levels in resistant potato attacked by \textit{P. infestans} or \textit{B. cinerea} positively correlated with symptoms of active cell death, which was identified by the TUNEL-positive assay illustrating the programmed DNA fragmentation (Fig. 8A and 8C). After 24 h of the challenge, approximately 95% of potato cells induced by \textit{avr P. infestans} contained green-colored nuclei (Fig. 8A), while cells elicited by \textit{B. cinerea} exhibited only 65% of nuclei with positive TUNEL staining (Fig. 8C). In susceptible potato challenged with \textit{B. cinerea} very few cells displayed HR-like cell death (Fig. 8D), while virulent \textit{P. infestans} elicited cell death identified as TUNEL-negative (Fig. 8B).

Collectively, these findings imply that \textit{B. cinerea} triggered HR-like death, which did not arrest the necrotrophic pathogen at the site of its entry, and then these dying cells expanded to

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Fig 7. Effect of hemibiotroph \textit{P. infestans} or necrotroph \textit{B. cinerea} inoculation on \textit{PR} gene expression. (A, B) \textit{PR}-1, (C, D) \textit{PR}-2 and (E, F) \textit{PR}-3 gene expression in potato leaves of cv. Bzura and cv. Bintje. (*) significantly different from mock-inoculated (control) potato leaves, P<0.05. Values represent mean ± SD of at least three independent experiments.

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Fig 8. TUNEL test illustrating programmed DNA fragmentation in potato leaves inoculated with the hemibiotroph \textit{P. infestans} or with the necrotroph \textit{B. cinerea}. (A) TUNEL fluorescein-positive reaction (green nuclei in fluorescence microscopy) detected in the \textit{avr P. infestans}–Bzura interaction at 24 hpi; (B) TUNEL fluorescein–negative reaction in \textit{P. infestans} susceptible potato cv. Bintje inoculated with the hemibiotroph; TUNEL fluorescein-positive reaction in (C) the Bzura–\textit{B. cinerea} and (D) the Bintje–\textit{B. cinerea} interactions at 24 hpi. Bars indicate 15 μm (A–D).

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form a spreading necrosis. The index of disease assayed 7 days after inoculation showed that \textit{B. cinerea} induced development of a strong gray mold covering over 80% and 67% leaf area with disease symptoms in susceptible and resistant potato genotypes, respectively (Fig 9). In the case of \textit{avr P. infestans}-inoculated leaves we detected late blight spots covering less than 3% of total leaf area, whereas virulent \textit{P. infestans} triggered severe 95% foliar damage observed in each leaf of susceptible potato (Fig 9).

### Discussion

Since the impact of NO on plant metabolism is related to its concentration being precisely regulated by the plant, depending on its abundance NO may provoke opposite effects under physiological and stress conditions [23, 50]. In this study we reported how potato leaves from the same genotypes challenged with pathogens of different lifestyles, e.g. \textit{P. infestans} and \textit{B. cinerea}, at the sites of attempted infection trigger varying magnitudes of NO burst and subsequent NO/ROS-associated defense signaling linked to an effective hypersensitive response or impaired resistance to the attacker. Our findings show that in contrast to the hemibiotrophic \textit{P. infestans}, the necrotrophic \textit{B. cinerea} strongly potentiates the NO burst and in synergy with H$_2$O$_2$ overproduction induces programmed cell death and finally exploits the potato defense system to its own benefit.

It has been well documented by other authors that the necrotrophic \textit{B. cinerea} produces non-host specific phytotoxic metabolites, including oxalic acid and botrydial, which by
manipulating the host redox milieu favor fungal penetration and colonization of plant tissue [51–53]. In turn, in an avr P. infestans–potato interaction, boosted NO generation in an induced program was observed to be at a much lower level than the necrotroph-dependent NO formation. Weaker NO-mediated activation was altogether sufficient to culminate in the prompt execution of the HR, found as the TUNEL-positive cell death, which arrested the hemibiotroph at the sites of its attempted entry. At the same time, very low production of NO during vr P. infestans–potato interactions might be crucial in avoiding active cell death (identified as TUNEL-negative) and could even be an important requirement for pathogen virulence [54].

S-nitrosylation as a redox-based post-translational modification represents the central transfer of NO bioactivity in plants [23, 55]. Considering the augmented NO production detected in infected potato leaves, we next examined whether NO might be converted into SNO storage. The results provided by the chemiluminescent method showed that enhanced NO synthesis found in infected leaves of potato cv. Bzura was accompanied by a significant rise in SNO pools, which was much more pronounced when challenged with P. infestans rather than B. cinerea. Relatively high levels of GSNO and SNOs detected immunocytochemically indicated the mobile function of these compounds as a reservoir of NO bioactivity concentrated in the main vein of potato leaves. The tissue-specific location of GSNO was also observed in sunflower hypocotyls exposed to high temperature and mechanical wounding [56, 57]. In our recent paper [44] we reported that in the avr P. infestans–potato system the NO-coding message was stored and converted into an enhanced total SNO pool and particularly S-nitrosylation of targeted proteins. Among previously identified proteins suggested for S-nitrosylation a reversible and redox-dependent modification of the germin-like protein converting oxalic acid to H₂O₂ was found [44, 58]. The proteomic analysis of S-nitrosylated proteins in Arabidopsis undergoing a hypersensitive response to Pseudomonas syringae pv. tomato (Pst avrB) also identified a germin-like protein related to its critical role at the onset of the cell death program [59].

Tyrosine nitration as the post-translational modification mediated by peroxynitrite is often considered a marker for certain pathological conditions; however, the nitration process might be involved in the signaling cascade under physiological conditions as well [45]. Probably the excess of ONOO⁻ generated under a pathophysiological state will determine the occurrence of negative modulations translated into an altered protein conformation or catalytic activity of targeted enzymes [60]. In this study, mainly in the 'Bintje' genotype treated with the necrotrophic pathogen, we demonstrated boosted tyrosine nitration in proteins to be closely correlated with a late time-dependent rise in the peroxynitrite level and nitro-tyrosine immunoassay. In turn, in an avr P. infestans-potato system NO was fine-tuned with O₂⁻⁻ formed peroxynitrite, probably resetting the signal following pathogen stimulation, resulting in weak ONOO⁻-dependent fluorescence and potato leaf nitration of target proteins. In contrast, a drastic increase in the pool of nitrated proteins, observed in the necrotroph colonized potato tissue, could be considered as a footprint of nitrosative stress [43, 45].

It has been generally accepted that rapid ROS generation is one of the immunity hallmarks in the early host response during an incompatible reaction between the plant and the biotrophic pathogen [4, 61]. In this study inoculation of potato leaves with avr P. infestans caused early and transient accumulation of O₂⁻⁻, reflected in the activation of NADPH oxidase and induction of HR resistance to this biotrophic pathogen. Probably ROS fine-tuned with NO can help the plant to stimulate a particular pathway leading to resistance to the pathogen [62]. Likewise, the results presented earlier [63] indicated that RBOHB silencing in Nicotiana benthamiana, in contrast to mutant of NOA1-mediated NO burst, reduced the effective defense against P. infestans. In Arabidopsis with high SNO concentrations NO governed a negative feedback loop compromising HR, mediated by S-nitrosylation of AtRbohD, at Cys 890, blunting its
ability to synthesize ROS [64]. Recently it has been reported that although AtRbohD mediates apoplastic ROS production in attacked cells, AtRbohF is crucial for reprogramming plant metabolism and establishing efficient resistance responses in neighboring cells during a susceptible interaction [65].

The present experiments revealed that in contrast to the hemibiotrophic *P. infestans*, the necrotrophic *B. cinerea* provoked suppression of the superoxide burst and concomitant defense responses in potato leaves. This means that huge amounts of NO, induced mainly by *B. cinerea* toxins, might temporarily antagonize the NADPH oxidase dependent O$_2$$^•$− production at the preliminary stage of inoculation.

Similarly, recorded data clearly documented that *Sclerotinia sclerotiorum*, a necrotroph related to *B. cinerea*, via elicitation of oxalic acid (OA) dampened the host oxidative burst and callose deposition by manipulating the plant redox environment [66]. However, at the later stages when infection was established, this necrotroph induced ROS generation leading to damage of the host tissue. Additionally, tomato leaves inoculated with the non-pathogenic OA-deficient mutant produced oxidative burst and displayed callose deposition [66].

As mentioned before, the host plant provoked by a necrotroph can produce H$_2$O$_2$ by germin-like proteins and remove apoplastic oxalate [13]. It was found in our study that mainly inoculation with *B. cinerea* caused a prompt increase in GLP activity, much greater than *avr P. infestans*.

The level of H$_2$O$_2$ accumulated in bean leaf discs inoculated with different *B. cinerea* isolates correlated with their aggressiveness [67]. Our previous experimental approach showed that leaves of ivy pelargonium (*Pelargonium peltatum*) in response to *B. cinerea* showed a strong NO burst and H$_2$O$_2$ accumulation through NO-dependent reversible inhibition of catalase and ascorbate peroxidase [29]. Nitric oxide together with H$_2$O$_2$ was produced during the initial stages of tissue invasion and also in dying cell layers in *Arabidopsis* mutant genotypes challenged with three pathogenic *Botrytis* species [28].

A growing body of evidence suggests that it is H$_2$O$_2$ rather than superoxide that synergizes with NO to execute cell death [68]. Accumulation of H$_2$O$_2$ is known to represent a defense response which is highly effective against biotrophic pathogens [69] but not against necrotrophs [70]. It is well documented that *B. cinerea* possesses multiple enzymes that are involved in massive oxidative burst driven by the NADPH oxidase complexes [71] or Cu-Zn superoxide dismutase [11] upon infection in the host plant. Although it is difficult to discriminate between pathogen and plant contribution to ROS generation, this necrotroph potentiates the host oxidative burst and requires ROS-dependent responses to achieve full pathogenicity [72]. The absent of *B. cinerea* *bcsod1* triggered oxidative imbalance (high superoxide and low H$_2$O$_2$ level) in *Arabidopsis* and tomato plants, reduced *B. cinerea* virulence and promoted efficient defense responses [11].

Next, timing and precise control of NO and H$_2$O$_2$ concentrations are essential in order to orchestrate the programmed cell death with the host resistance to the invader. Experimental data published to date have not been clear whether the cooperation of NO/H$_2$O$_2$ plays a role in the elicitation of cell death or cell death propagation [24, 73].

Analyses of pathogen-dependent programmed cell death in the Bzura potato genotype showed that *B. cinerea* triggered HR-like death, which did not arrest the necrotizing pathogen at the site of its entry, and then these dying cells expanded to form a necrotic region. It is possible that this form of cell death may be the effect of host metabolic dysfunction, in which high NO concentrations play a major role.

Predominantly, the hypersensitive response to a biotroph involves both the early NO burst and SA-dependent signaling, which promotes ROS production [25]. In our experiment despite the high constitutive SA level in potato leaves, a further rise in the free SA level was observed mainly in the potato resistant to *avr P. infestans*. In turn, a high NO burst observed after challenging potato leaves with *B. cinerea* negatively regulated SA production but not PR expression.
The existence of additional signals that contribute to the induction of PR expression cannot be excluded [74]. As it was revealed, the proton electrochemical gradient across the plasma membrane and sugar metabolism could play a regulatory role in the SA-independent induction of PR gene expression [75, 76].

It also should be taken into account that the induction of a signaling route can activate its own negative feedback, and sometimes this complex network is difficult to interpret. Recent findings in an Arabidopsis mutant line (nox overexpression mutant) challenged with P. syringae (PstD3000) also suggested that enhanced NO accumulation compromised multiple modes of disease resistance by blunting both SA synthesis and SA signaling [77]. Also, abscisic acid decreased resistance to B. cinerea in tomato via reduction of NO, H₂O₂ and ET production [78].

We found that both avr P. infestans and B. cinerea caused an early ethylene burst in potato leaves, but the magnitude and timing of ET emission substantially differed between these responses. First of all, the B. cinerea-inducible program of boosted ET synthesis was stimulated early and lasted for the successive hours after inoculation; however, it was not linked with rapid SA-dependent gene up-regulation.

Essentially, PR-1 gene expression negatively correlates with resistance to B. cinerea, except for the Arabidopsis BOTRYTIS-INDUCED KINASE1 (BIK1) gene mutant highly susceptible to necrotrophic fungal pathogens [79].

Our final data indicate that precise control of early NO production in cooperation with ROS at the site of the pathogen contact with potato leaf tissue activates NO- and ROS-dependent sequences of events for the host’s effective defense against the invader. Mainly in the avr P. infestans–potato system NO well balanced with the H₂O₂ level was tuned with a battery of SA-dependent defense genes, facilitating the establishment of pathogen resistance. The genetic inability of the susceptible potato to accumulate suitable amounts of NO/H₂O₂ results in delayed PR-1 and PR-3 gene expression and tissue colonization by virulent P. infestans. Meanwhile, huge NO overproduction in B. cinerea-challenged potato accelerated HR-like cell death, although it abolished defense-related gene expression and finally plant resistance. These data could indicate that NO is a key player in both PAMP-triggered immunity (PTI) and effector-triggered immunity (ETI) strategies.

The present findings are in line with the concept on the organismal cost of having the machinery for HR as a potential ecological cost of biotroph resistance, causing increased sensitivity to necrotrophic pathogens [1, 80]. It may thus be stated that if plants possess specific signaling equipment to stimulate HR control via NO/H₂O₂, they might experience fitness benefits and fitness costs, associated with this defense strategy being overcome by the necrotroph. The necrotrophic B. cinerea–by manipulating both NO and superoxide synthesis–affects the host redox environment and exploits the potato defense system to its own advantage. Recently the chaperone-like CDC48 was identified as the NO target, and it was suggested that both NO and ROS are probably important components of the mechanism coordinating the ubiquitination system in the cell death pathway [58, 62]. For confirmation, S-nitrosylation of polyubiquitinated proteins was documented by us in potato cells facing a P. infestans attack [44].

Finally, deep insight into the involvement of NO and ROS closely related to other components and effectors interfering with the potato defense mechanism and inducing side effects might help us apply new environmentally safe methods to improve potato protection against pathogens of different lifestyles.

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