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8 The use of ECAS in plant protection: a green and efficient
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11 antimicrobial approach that primes selected defense genes
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4 **ABSTRACT**
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7 The use of highly polluting chemicals for plant and crop protection is one of the components of
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9 the negative environmental impact of agricultural activities. In the present paper, an
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11 environmentally friendly alternative to pesticide application has been studied, based on the so-
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13 called *electrochemically activated solutions* (ECAS). Experiments have been carried out, by
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15 applying ECAS having different contents of active ingredients, on tobacco plants at a laboratory
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17 scale and on apple trees at fruit garden scale. The results, accumulated during a couple of years,
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19 have shown that properly selected dilute solutions of chlorides, once activated by an
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21 electrochemical treatment, exhibit a very effective protecting action of plants, irrespective of their
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23 nature. Extension of the research has shown that the observed effect is the result of two distinct
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25 factors: the expected anti-microbial action of the electrochemically synthesized oxidants, and an
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27 unexpected priming of immune plant defenses, which is clearly due to the treatment with ECAS.
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29 Interestingly, the repetition of ECAS application triggers an even stronger activation of defense
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31 genes. No oxidative damages, due to the use of the activated solutions, could be detected.
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43 **KEYWORDS**
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45 Electrochemically activated solutions; endogenous plant defense; antimicrobial activity; green
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47 chemicals; oxidative stress; hypochlorous acid.
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Introduction

Since infestation by pests and diseases can severely reduce the yield of agricultural and horticultural crops, one of the main concerns of modern agriculture is the fight against pathogens. In order to protect crops, before and after harvest, chemical products (generically and collectively referred to as ‘pesticides’) are typically used. In some cases, these chemicals act by confusing insects or making crops less palatable for pests; more commonly, the effectiveness is directly exerted by killing the undesired insect, fungi or bacteria. Obviously, such pesticides could have severe undesirable effects if they are not strictly regulated. Since the entry into force, in 2009, of European Directive 2009/128/EC (which explicitly refers to the sustainable use of pesticides), no plant protection product can be used within the Member States, unless it has first been scientifically established that:

- (1) it has no harmful effects on consumers, farmers and local residents and passers-by;
- (2) it does not cause unacceptable effects on the environment;
- (3) it is sufficiently effective against one or more pests.

Based on the above, the use of environmental- and human-friendly molecules has become increasingly important, especially in the last few years. Researching for active substances capable of killing pathogens without harming the environment, the so-called *electrochemically activated solutions* (ECAS) captured our attention. ECAS are slightly saline solutions that, after electrolysis, contain reactive oxidative species (ROS) characterized by a short life span and a potent antimicrobial activity (Hyang et al., 2008 and references therein; Thorn et al., 2012; Mukhopadhyay and Ramaswamy, 2012). The electrochemical treatment of a halogenides-containing solution typically leads to the formation of hypohalogenides (e.g., hypochlorite), which are well known not only because they are effective disinfectants (FAO and WHO, 2009; Gómez-López, 2012), but also for their capability in removing stains and odors, as well as for

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4 their bleaching properties (A.I.S.E. scientific dossier, 1997). Actually, hypochlorous acid (HOCl)
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6 is also an important component of vertebrate nonspecific immune system. Thanks to the action of
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8 the myeloperoxidase enzyme, it is synthesized from hydrogen peroxide (H₂O₂) and chloride
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10 anion (Cl⁻), during the neutrophils' respiratory burst (Klebanoff, 2005).
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14 On the other hand, plants have developed complex strategies of defense. Differently from
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16 animal immune system, plants do not have a mobile immune system but each cell is able to
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18 trigger a defense mechanism. These defenses can be divided into two classes: passive and active
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20 (Hammond-Kosak and Parker, 2003; Dangl and Jones, 2001; Niks and Marcel, 2009; Ahmad et
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22 al., 2010). Passive or constitutive defenses, are physical or chemical barriers, always present in
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24 plant cells and tissues. These basic defenses can be easily overcome by the pathogen, especially if
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26 it manages to enter plant tissues through wounds or by enzymatic action. Yet, the plant can still
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28 fight back the pathogen action, by active or inducible defenses, deployed as a consequence of
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30 pathogen recognition by the plant. In fact, recognition is the key factor for the success of the
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32 strategies employed and it has a genetic base (Ahmad et al., 2010). This recognition activates a
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34 more robust multilevel defense (Jones and Dangl, 2006; Bent and Mackey, 2007; Carr et al.,
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36 2010). The general pattern of this active resistance is based on three important levels (Jones and
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38 Dangl, 2006; Bernoux et al., 2011). Firstly, a rapid and powerful *hypersensitive response* (HR)
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40 weakens the pathogens near the site of infection through an oxidative burst (Wojtaszek, 1997;
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42 Gozzo, 2003).
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50 Besides, from the infection site, a series of molecular signals are sent to the whole plant (ROS,
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52 salicylic acid or jasmonic acid, azelaic acid (Conrath, 2011; Conrath et al., 2015)) triggering a
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54 broad and potent defense in neighboring tissues. This kind of defense is very powerful and
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56 involves the synthesis and accumulation of a heterogeneous class of proteins, the so-called PR
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58 (pathogen related) proteins, which have different chemical and physical properties (Tuzun and
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4 Somanchi, 2006). The latter are induced in plants resistant not only to microorganisms but also to
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6 other biotic (insects, nematodes, herbivores) or abiotic stresses (Van Loon et al., 2006). 17
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8 classes of PR proteins are known so far (Van Loon et al., 2006; Sels et al., 2008): most of them
9
10 show antimicrobial and antifungal activity (chitinases, glucanases, proteinases). These
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12 phenomena trigger a further defensive wave that extends to distal organs creating an immunity
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14 called *systemic acquired resistance* (SAR). During SAR, plant tissues are prompted to induce a
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16 strong defense response upon new infections (Ahmad et al., 2010) thanks to one or more signals
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18 (SA, JA and others) (Pieterse et al., 2009; Shah, 2009; Shah et al., 2014) that originated from the
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20 initial site of infection and spread to the whole plant (Heil and Bostock, 2002). These three
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22 coordinated steps often ensure that the plant stays healthy, with no disease symptoms.
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28 The fight against diseases in agriculture is usually based on the use of active substances toxic
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30 for the pathogen, and often also for the environment and humans. The use of molecules able to
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32 activate the endogenous plant defenses is still uncommon, even if it has been receiving a lot of
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34 attention in the last few years (Shah et al., 2014; Du et al., 2012; Bai et al., 2011; Chaturvedi et
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36 al., 2012).
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43 *A prompt from a direct field experience.*

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45 In order to test a non-conventional plant-protecting treatment, in 2005 an extensive field
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47 experiment was carried out in a 2 ha apple orchard severely infected by *Nectria galligena*. The
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49 situation was so serious that agronomists contacted to resolve the problem could but suggest
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51 uprooting of orchard trees and wood incineration. At that time, our experiences with ECAS were
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53 just at the beginning, and we thought to spray the trees with the disinfectant solution, as
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55 suggested in initial papers on the subject (*e.g.*, Al-Haq et al., 2005; Buck et al., 2003, and
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57 literature therein). Quite surprisingly, the treatment caused a drastic interruption of the
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4 pathogenic cycle and a robust healing of the lesions. At the same time, plant growth and crop
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6 production were considerably improved. As a possible comparison (control), untreated
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8 neighboring orchards of the same nature and with the same infections remained affected by the
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10 diseases and have been explanted one year later.
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14 To understand these phenomena, we started an investigation based on molecular and cellular
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16 analyses, in order to understand whether the ECAS treatments were only responsible for a
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18 superficial sanitation of the infected plants or they were also able to induce specific plant defense
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20 strategies. The study took into consideration both *Nicotiana tabacum* (tobacco) and *Malus*
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22 *domestica* (apple) plant species. The former is often considered as a model in plant biology, while
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24 the latter is interesting especially for its commercial value.
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28 Here we report results suggesting that ECAS treatments are able to trigger endogenous plant
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30 defenses, in particular genes belonging to the PR family and other cellular defenses. Accordingly,
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32 the ECAS seem to represent a new-generation of active substances that combines a powerful
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34 antibacterial and antimicrobial effect, a marked effect on the plant immune system and the
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36 absence of environmental impact.
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44 **Materials and Methods**

45 *Preparation of electrochemically activated solutions*

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47 As briefly discussed in a recent review by Reynolds and coworkers (Thorn et al., 2012),
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49 electrochemically activated solutions can be synthesized with different approaches. Investigations
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51 began in Russia in the early 70s, thanks to the work of Academician V. Bakhir (Bakhir, 1985;
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53 Prilutsky and Bakhir, 1997), who devised a flow-through electrochemical module, based on
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55 concentric electrodes separated by a ceramic membrane (Bakhir et al., 1995). A separator
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4 between the electrode compartments is required when an *anolyte* (an oxidizing, generally acidic
5 solution) and a *catholyte* (a reducing, alkaline solution) are sought. In addition, a neutral anolyte
6 can be produced by suitably adjusting the (hydraulic) mixing of the above liquids. In the presence
7 of chlorides, gaseous chlorine (Cl₂) is synthesized at the anode of the electrochemical reactor.
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9 Once formed, chlorine dissolves in water producing a *free* (or *active*) *chlorine* solution. The
10 solution pH decides the forms in which chlorine is present, hypochlorous acid (HOCl),
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12 hypochlorite (ClO⁻) or gaseous chlorine (Cl₂), and eventually represents the key factor to explain
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14 the bactericidal effectiveness of the anolyte (Len et al., 2000; Abadias et al., 2008; Xiong et al.,
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16 2010).
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26 In the present investigation, we took into account two specific pH values (6.5 and 9.0), in order
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28 to assess the roles of hypochlorous acid and hypochlorite ion (these two species are related by a
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30 chemical equilibrium: HOCl + H₂O ⇌ ClO⁻ + H₃O⁺, pK_a ≈ 7.53). At the lower pH, most of the
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32 *active chlorine* is present as HOCl (90%), the hypochlorite form contributing for the remaining
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34 10%; at pH 9, on the contrary, the opposite is observed (HOCl 5%, ClO⁻ 95%).
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38 Solutions were obtained by starting from dilute brines (5 g L⁻¹ of either NaCl or KCl, in tap
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40 water), which were electrolyzed for a few minutes in electrochemical reactors provided with or
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42 without a separator, in order to obtain the desired pH values. After the synthesis, the solution pH
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44 was measured with a pH-meter and adjusted to the required value by adding small amounts of
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46 hydrochloric acid or sodium hydroxide. The *active chlorine* content (always comprised between
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48 700 and 1000 mg L⁻¹) was quantified by means of the standard *N,N*-diethyl-1,4-
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50 phenylenediamine colorimetric method, on specimens diluted with distilled water (ISO 7393-2,
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52 2002).
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Plant materials

Plants of *Nicotiana tabacum* cv. Petite Havana SR-1 and *Malus domestica* cv. Fuji were grown in a controlled growth chamber with a 16/8 h photoperiod ($150 \mu\text{E m}^{-1} \text{s}^{-1}$) at 24 °C and 60% of humidity. The mutant tobacco plants *NahG* (Friedrich et al., 1995) (kindly provided by Dr. Luis Mur, University of Aberystwyth, Wales, UK), unable to accumulate salicylic acid, and the transgenic tobacco plants carrying the GUS reporter gene under the control of the pathogen related gene PR1 (*PR1a-GUS* (Grüner and Pfitzner, 1994), kindly provided by Dr. Ursula Pfitzner, University of Hohenheim, Stuttgart, Germany) were grown in the same conditions. Field tests were done in a local farm on a parcel of *Malus domestica* cv. Dallago (20 y old).

Treatments and samplings

Electrochemically activated solutions containing 250 mg L⁻¹ of *active chlorine* (AC) and adjusted at the desired pH (6.5 or 9.0) were sprayed on plants of tobacco (2-months old) and apple (1-year old), until their complete wetting. In parallel, plants of tobacco and apple were sprayed with a control non-electrolyzed solution, having a similar pH and chloride content. 2 h, 6 h, 18 h and 24 h after the first treatment with ECAS and control solutions, the fourth and fifth leaves were harvested; upon a second treatment, leaf samples were collected after 2 h, 6 h, 12 h, 24 h, 48 h or 96h.

In field tests, due to the hardness of trees, the concentration of ECAS and control solutions was increased to 500 mg L⁻¹ and leaf pool were collected at 24 h, 48 h and 96 h.

After sampling, the leaves were immediately frozen in liquid nitrogen, grounded in liquid nitrogen and stored at -80 °C.

RNA extraction and real-time PCR

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4 Total RNA was extracted from 100 mg of powdered leaves by “Spectrum™ Plant Total RNA
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6 Kit” (Sigma). To avoid genomic DNA contaminations, a DNase treatment was carried out as
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8 suggested by the manufacturer. Total RNA integrity was then controlled on 1% (w/v) agarose
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10 gel, while the genomic contamination was assessed by conventional PCR. The relative
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12 quantification of mRNAs was performed by means of a two steps real-time analysis. In the first
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14 step, single-strand cDNA was prepared using iSCRIPT™ cDNA synthesis Kit (Bio-Rad). In the
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16 second step, the real-time PCR was performed using a SYBR green I mix (SsoFast EvaGreen
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18 Supermix-Bio-Rad) on a Chromo4 real-time PCR system (Bio-Rad). The efficiency of the cDNA
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20 synthesis was evaluated by real-time PCR amplification using the control Actin gene. Only the
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22 cDNAs that showed an Actin Ct value of 22 ± 2 were considered for the subsequent gene
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24 expression analyses. Chosen genes are listed in Table 1.
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34 --- *Table 1* ---
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38 The relative expression level of all genes investigate were obtained using the geNorm
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40 algorithm (geNorm, 2008). This algorithm requires the use of two stable reference genes (in our
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42 case Actin and Elongation Factor 1 α). At each time, the treated and control samples were
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44 compared with geNorm. qPCR reactions have been performed in triplicate and intra-assay
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46 variance was considered acceptable when the standard deviation was lower than 0.5. Given the
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48 high biological variability, another specific algorithm has been used to compare the replicates
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50 (Vandesompele et al., 2002; Willems et al., 2008).
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53 The results obtained have been expressed as “fold changes”, *i.e.* as the over-expression of the
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55 genes induced by the electrolyzed solution, compared to the control solution.
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4 *Histochemical analyses*

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6 *Superoxide and H₂O₂ staining.* The O₂^{•-} and H₂O₂ production was evaluated by histochemical
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8 staining with nitroblue tetrazolium (NBT) and 3,3-diaminobenzidine (DAB), respectively. After
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10 ECAS treatment, full leaves from treated and control plants were cut at different time points and
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12 the *in situ* presence of superoxide and hydrogen peroxide was evaluated as described previously
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14 (Jabs et al., 1996; Dutilleul et al., 2003; Thordal-Christensen et al., 1997). Positive control plants
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16 were treated with Paraquat 20 μM.
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23 *GUS histochemical assay.* To evaluate if the treatments with ECAS were able to induce the SAR
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25 molecular marker PR1a, transgenic plant PR1a-GUS were treated either with ECAS or with the
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27 control solution. The GUS histochemical assay was carried out as previously described (Vitha et
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29 al., 1995). Briefly, tobacco leaves harvested 6 and 24 h after the treatment were incubated for 30
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31 minutes in 5 mL of fixative buffer, washed several times for 30-40 minutes in washing buffer and
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33 vacuum-infiltrated in the X-gluc substrate for 30 minutes. Staining was performed in darkness at
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35 37 °C for 16-20 h. To remove chlorophyll, leaves were washed with 70% ethanol. Photographs
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37 were taken with a Nikon Eclipse E200 optical microscopy.
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47 **Results and Discussion**

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49 *Results of gene expression on Tobacco*

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51 Electrochemically activated solutions are potent sterilizing solutions, able to kill several kinds of
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53 phyto-pathogenic microorganisms. Their application on vegetative parts of diseased plants may
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55 heal the plant organs but it is not clear whether this effect is only due to the biocide activity or
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57 there is also a positive effect on the plant. To evaluate the plant responses to ECAS treatment, we
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4 chose healthy tobacco plants and analyzed the expression profiles of several genes involved in
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6 plant defenses against pathogens (members of the PR family, enzymes and regulatory proteins)
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8 comparing the fold change observed in ECAS-treated plants with control-treated plants. The
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10 relative quantification of the mRNA levels for each gene was performed using real-time PCR by
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12 considering two other genes as reference (Actin and EF1 α). To take into consideration the PCR
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14 efficiency, the geNorm algorithm was employed (geNorm, 2008; Vandesompele et al., 2002).
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19 Tobacco plants were treated with ECAS (250 mg L⁻¹) or control solution, and leaf samples
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21 collected at different times (2-24 h) from different plants, in order to exclude wound-induced
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23 responses. Then, total RNA was extracted, retro-transcribed and amplified in real-time PCR. PR
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25 and other defense-related genes were examined and their expression level normalized with
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27 reference genes. In ECAS-treated plants, the activation of genes was expressed as fold change in
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29 comparison with control-treated plants. The results shown in Fig. 1 reveal that a single ECAS
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31 treatment can increase the expression level of several PR genes (especially PR1a, PR2 and PR17,
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33 10 fold increase) starting from 6 h after the treatment. No PR up-regulation was detected 2 h after
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35 the ECAS treatment, while at later stages only a few genes were still expressed (PR1a, PR2,
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37 PR3a and PR17). A significant and fast up-regulation was also observed for the HR-related gene
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39 HSR203J during the first 2-6 h, while PAL and WRKY3 genes showed a 10-fold increase 6 h
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41 after treatment and then returned to basal levels. Other genes (NPR1, non-expressor of PR1, is a
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43 key regulator of systemic acquired resistance; ICS, isochorismate synthase; PR2a, basic Class I
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45 glucanase; and PR3, basic class I endochitinase) were also analyzed but did not show any
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53 induction upon treatment.

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55 Control-treated plants did not show significant differences in terms of gene expression profiles
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57 when compared to untreated plants (data not shown), confirming that the activation observed
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4 after treatment with ECAS was not merely due to the dilute brine solution itself, but rather to the
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6 (more or less labile) chemical species generated by electrochemical activation.
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9 The same tobacco plants were treated again with ECAS 14 d after the first treatment. This
10 allowed determining whether plant molecular defense responses vary with time and if there was a
11 priming phenomenon associated with ECAS treatments. Leaf samples were collected at different
12 time points from independent plants (to avoid wound-induced effects) and tested in quantitative
13 real-time. Fig. 2 reveals that a second ECAS treatment enhances PR gene expression, especially
14 24 h after the treatment, with very high levels of induction for PR1a and PR2 (up to 1000x fold
15 change). Other PR genes showed expression profiles similar to those observed after the first
16 treatment. In terms of the other genes analyzed, PAL gene was again rapidly induced at high
17 levels (40x) but later its expression level decreased. Interestingly, most of the genes analyzed
18 appear to be still overexpressed 48 h after the treatment; this outcome was not observed after the
19 first treatment. No sign of phyto-toxicity was ever observed.
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36 When the second treatment was performed on tobacco plants 35 d after the first treatment, the
37 gene expression analysis revealed a significant activation of several PR genes (PR1a 100x fold
38 change) that lasted up to 96 h. On the other hand, there were no differences in the expression
39 levels of other defense genes (PAL, WRKY3, etc.; data not shown). This would suggest that
40 ECAS treatment can induce a priming phenomenon that appears quite strong during the first
41 weeks and then it decreases in intensity while remaining still detectable after one month.
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53 *Dependence of gene expression on ECAS concentration and pH*

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55 To establish if the observed gene induction is influenced by the concentration of active chlorine
56 present in the electrochemically activated solution, we sprayed tobacco plants also with solutions
57 containing 125 and 500 mg L⁻¹ of AC. Leaves were collected from independent plants after 12
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4 and 48 h. After a first treatment, quantitative real-time amplification and fold change analysis
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6 revealed that the induction in the expression of the defense-related genes (both PRs and other
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8 genes) was comparable to those of the previous treatments (with 250 mg L⁻¹ of AC, data not
9
10 shown). On the other hand, a second treatment 14 d after the first revealed that the priming
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12 phenomenon is much more intense with ECAS at 250 mg L⁻¹. At 125 mg L⁻¹, the defense-genes
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14 overexpression was quite weak, even weaker at 500 mg L⁻¹ as compared to the overexpression
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16 observed with ECAS at 250 mg L⁻¹ (data not shown).
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21 These molecular data thus suggest that an AC content of 250 mg L⁻¹ is an optimal concentration
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23 for tobacco plants since it allows achieving the highest expression level for the longest period. It
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25 is worth noting that no symptoms of phyto-toxicity were observed on plants treated even at
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27 higher ECAS concentrations (up to 800 mg L⁻¹, data not shown).
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31 We also assessed if the observed gene overexpression was linked to the pH value of the
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33 electrochemically activated solution. As previously mentioned, the pH value of the solution
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35 changes considerably the nature of the molecules present in the electrolyzed solution. At neutral
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37 pH, hypochlorous acid is prevalent, while at alkaline pH the hypochlorite form is predominant.
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39 We treated tobacco plants with ECAS containing 250 mg L⁻¹ of AC at either pH 6.5 or pH 9.0,
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41 and carried out mRNA analyses via real-time PCR on leaves collected at various time points. As
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43 shown in Fig. 3, the electrochemically activated solution with alkaline pH triggers an
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45 overexpression that is limited to some of the PR genes (PR1a and PR2), while other genes (PRs,
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47 PAL, HSR, RBOH and WRKY3) are not up-regulated. When a second treatment is performed 14
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49 d after the first, a partial priming and enhancement of gene expression has been evidenced by
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51 real-time PCR analysis 24-48 h after the treatment, but they are not comparable to the
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53 enhancement obtained with electrochemically activated solutions at pH 6.5. PR1a and PR2 show
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55 a 100x and a 10x fold increase, respectively, with ECAS at pH 9 (24 h) while they showed a
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4 1000x and a 100x fold increase 24 h after the second treatment in the case of ECAS at pH 6.5.

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7 The other defense-related genes were only slightly over-expressed, while with ECAS at pH 6.5
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9 they were significantly over-expressed in both the 24 and the 48 h samples.

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11 Accordingly, these analyses indicate that hypochlorous acid, rather than hypochlorite, is essential
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13 to achieve a strong and long lasting activation of plant defenses.
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16 Since the observed results are related to defensive, salicylic acid-dependent protein activation,
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18 it might be that ECAS is able to trigger an increase in the production of salicylic acid. The
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20 quantification of the salicylic acid hormone (an important signaling molecule, activator of the
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22 endogenous defenses) was carried out on tobacco leaf tissues collected at different time points,
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24 after a first or a second treatment (14 d). The quantification of salicylic acid in leaves was
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26 performed by GC-MS analysis (Deng et al., 2003); results, expressed as the ratio of
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28 concentrations found in the treated plants against those used as controls, are shown in Fig. 4. The
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30 chemical data confirmed those obtained by molecular analysis, proving that the signal molecule
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32 of salicylic acid is effectively synthesized after a single ECAS treatment (10 times more abundant
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34 than in control plants). The hormone levels increased drastically even after a second treatment,
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36 with an induction profile quite similar to that described for the gene activation. As a result, these
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38 data seem to suggest that salicylic acid is at least partially implicated in the immune response of
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40 ECAS-treated plants.
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48 In order to confirm the role of salicylic acid in the ECAS-triggered up-regulation of genes, tests
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50 were carried out also on transgenic tobacco plants that are unable to accumulate this hormone
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52 (*NahG*). As can be seen from Fig. 5, tests carried out using the *NahG* plants show a gene
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54 activation profile much less pronounced compared to wild-type plants. In particular, a weak
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56 activation of PR1a, a typical salicylic-dependent gene, is evident after a single ECAS treatment
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58 and even weaker after the second treatment (maximum fold change 100x as compared to 1000x
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4 for wild-type plants). Nevertheless, many defense genes appeared activated despite the absence
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6 of SA, thus suggesting that the activation of defenses induced by treatment with ECAS takes
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8 place along with the synthesis of salicylic acid, but the latter does not seem to be the main cause
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10 of activation of genes. SA participates in the gene activation and in priming the plants, but it is
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12 not the sole responsible, therefore other pathways might be involved.
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18 *Histochemical investigations*

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20 Since the ECAS application can possibly induce an oxidative stress, we evaluated the
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22 accumulation of reactive species such as H₂O₂ and O₂^{•-} in tobacco leaves sprayed with ECAS
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24 through histochemical staining. The synthesis of hydrogen peroxide was analyzed using 3,3'-
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26 diaminobenzidine (DAB) (Dutilleul et al., 2003), whereas the nitroblue tetrazolium (NBT)
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28 staining method was used to highlight the presence of the superoxide anion (Dutilleul et al.,
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30 2003). Tobacco leaves were collected and stained 2 and 6 h after ECAS application; Paraquat
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32 was used as a positive control. The degree of staining of ECAS-treated leaves was not significant
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34 (Fig. 6), as compared to the positive control, thus suggesting that ECAS treatments do not
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36 generate any oxidative stress in tobacco plants.
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43 The gene expression data were also compared to histochemical data obtained from transgenic
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45 tobacco plants bearing a PR1a-GUS transgene able to express the GUS reporter protein as a result
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47 of the PR1a promoter activation (Grüner and Pfitzner, 1994). Tobacco PR1a-GUS plants were
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49 treated with ECAS analogously to WT plants treated before and leaf samples were collected at
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51 different time points. Then, the tissues were immediately fixed and stained overnight in X-Gluc
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53 solution. After chlorophyll removal, ECAS-treated samples were clearly stained after either the
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55 first or the second treatment (Fig. 7) while control-treated plants did not show any GUS-specific
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4 staining. This evidence confirms what previously observed via real time PCR: the PR1a gene is
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6 overexpressed upon ECAS treatment and enhanced upon a delayed second treatment. As a
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8 positive control, we treated PR1-GUS plants with the SA analog BTH (benzo-(1,2,3)-thiadiazole-
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10 7-carbothioic acid S-methyl ester), a potent activator of PR1a (Görlach et al., 1996). 24 h after
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12 the treatment, an intense staining was observed due to the strong PR1a activation (also confirmed
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14 at the molecular level: PR1a 1000x fold change by RT-PCR).
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21 *Results of gene expression on Apple*

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23 Based on the molecular data obtained on tobacco, we extended the investigation and gene
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25 activation analysis to a species of great agronomic interest such as apple. ECAS applications
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27 were carried out on 1 year-old plants grown in a growth chamber, as well as on mature trees
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29 (about twenty year-old), in orchard. The young plants were treated with an electrochemically
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31 activated solution at pH 6.5 containing 250 mg L⁻¹ of *active chlorine* (Fig. 8, A and B). Since the
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33 old plants in the orchard showed thicker leaves and a greater vigor, as compared to those in the
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35 growth chamber, we increased the *AC* concentration to 500 mg L⁻¹ (Fig. 8, C and D).
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41 Similarly to what observed in tobacco, a first treatment with ECAS is able to trigger a
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43 defensive response in apple trees. All tested PR genes as well as the PAL gene were up-regulated
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45 (10-30x fold change), especially in the case of plants in orchard, whose responses were stronger
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47 than those of plants in the growth chamber. Differently from what observed in tobacco plants, a
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49 second ECAS treatment, 14 d after the first, is capable of triggering a defensive response in apple
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51 plants, but with weak “enhancement effect”. In apple trees, the gene expression remains
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53 approximately constant, with no significant variations with the gene expression level observed
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55 after the first treatment.
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4 Organoleptic tests (color, and sugar content of apple fruits) as well as chlorophyll and total
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6 nitrogen contents (in leaves) did not indicate changes, neither in the properties and quality of
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8 fruits, nor in the vigor of treated plants (data not shown). As regards the antimicrobial effects
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10 associated with treatments, SEM micrographs of orchard apple leaves taken before (Fig. 9, A and
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12 B) and after (Fig. 9, C and D) the ECAS application clearly show how the treatment substantially
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14 leads to the disappearance of any microorganism from the surfaces. Pictures taken on leaves
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16 treated with the control solution (KCl 5 g/L, pH 6.5) did not show any noticeable change as
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18 compared to untreated leaves (data not shown). This is a somewhat expected outcome: the leaves
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20 are normally subjected to bad weather, in nature, and the simple washing with water or with a
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22 diluted salt solution is not sufficient to wash away the microorganisms. On the contrary, the
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24 washing with ECAS is effective, plausibly because the active chlorine is able to destroy the
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26 biofilm (Ozaki et al., 2012) and the microorganisms adhesion as well. Overall, this represents a
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28 further proof of the potential of ECAS as a green alternative to other pesticides.
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38 **Conclusions**

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40 The presented results suggest that electrochemically activated solutions, thanks to their proven
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42 antimicrobial- and antifungal-activity along with the efficacy in activating the molecular defenses
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44 of plants (especially after repeated treatments) and the low phyto-toxicity, are a promising
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46 alternative to conventional pesticides. Owing to its various possibilities of attack, the active
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48 ingredient is not expected to induce resistance in the target pathogens; actually, the lack of
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50 specificity represents also the weak point of the application, which requires to be suitably timed,
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52 in accordance with plant growth and, hopefully, within an integrated pest management program.
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55 This green alternative to the pesticides has the advantage of being safe for the users and do not
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4 leave residuals on treated fruits. Lastly, but no less important, ECAS do not pose environmental
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6 risks and can be produced when required, at low costs.
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34 **Author Contributions**

35
36 The manuscript was written through contributions of all authors. All authors have given approval
37
38 to the final version of the manuscript. § These authors contributed equally.
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41

42 **Conflict of interest**

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45 The authors declare that they have no conflict of interest.
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Table 1. Panel of analyzed genes, for both tobacco and apple plants, with respective GenBank accession numbers.

Gene	Function	Tobacco	Apple
PR1a	Antifungal	X06361	DQ318212
PR2	β -1,3-glucanase	M60460	AY548364
PR3a	Acidic Chitinase	X51426	AF494395
PR4a	DNA-RNase ?	X58547	
PR5	Thaumatin-like, antifungal	S44889	DQ318213
PR8	Endochitinase		DQ318214
PR17	Unknown	AB024600	
PAL	Phenylalanine ammonia-lyase	D17467	AF494403
WRKY3	Transcriptional factor	AF193770	
RBOH	Hypersensitive response-related protein	AJ309006	
HSR203J	Hypersensitive response-related protein	X77136	
Actin	Reference	U60493	GQ339778
EFNT1 α	Reference	AF120093	DQ341381

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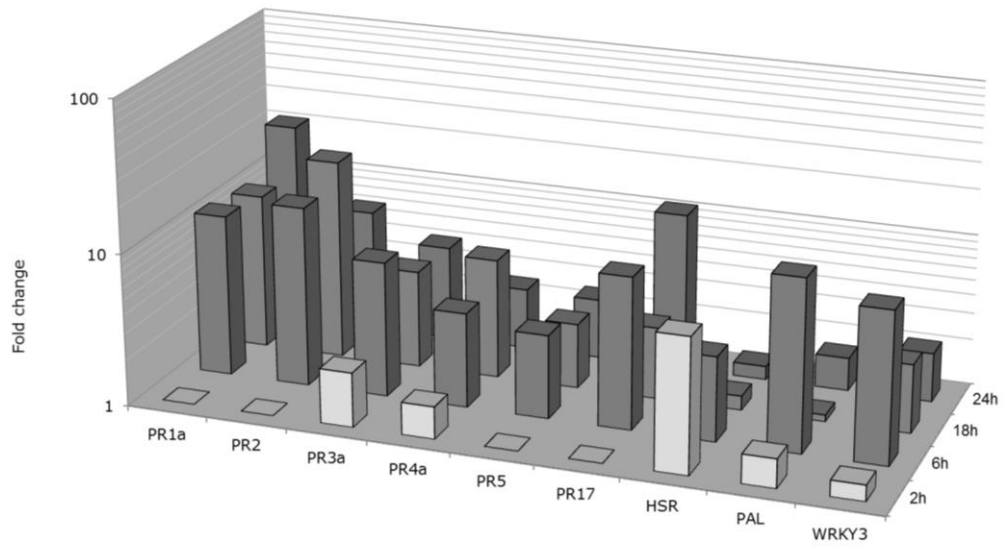


Fig. 1 Gene expression on tobacco plants, treated once with ECAS containing 250 mg L⁻¹ of AC at pH 6.5. Fold-change values presented a standard deviation always lower than 20%.

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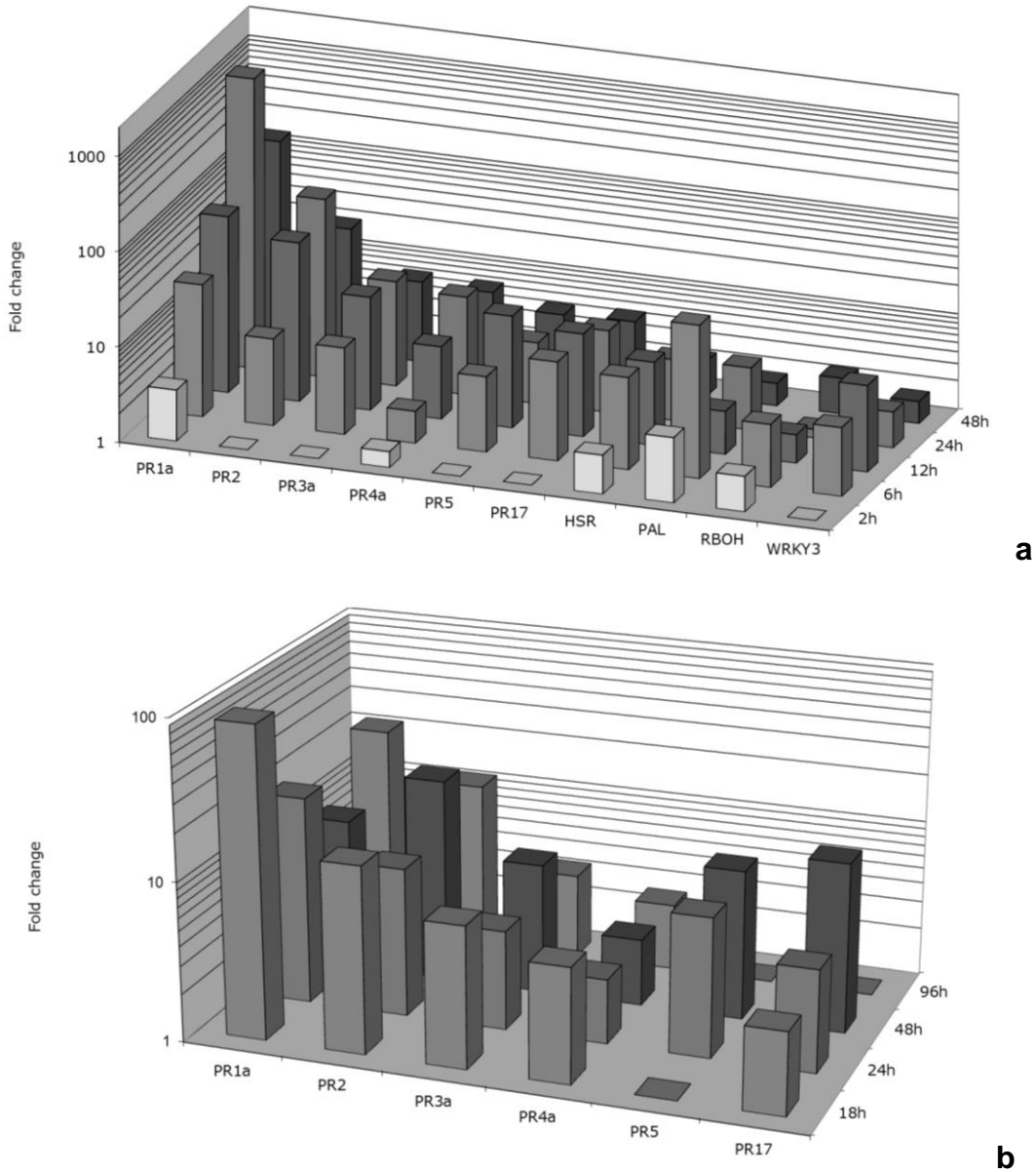
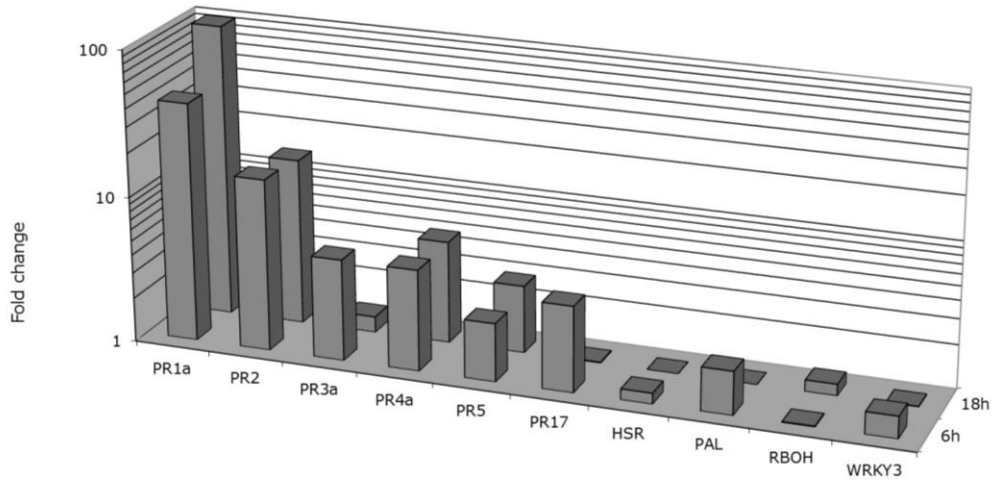
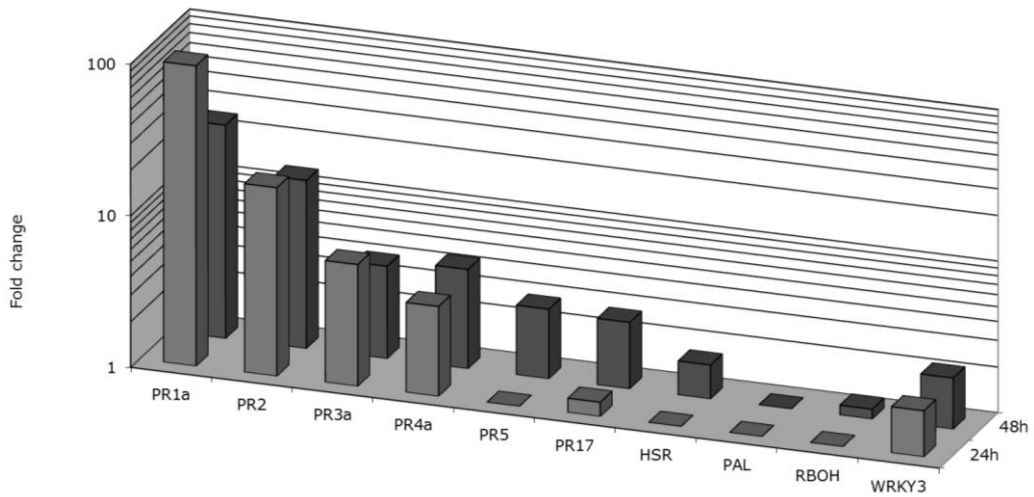


Fig. 2 Gene expression on tobacco plants, treated twice with ECAS containing 250 mg L⁻¹ of AC at pH 6.5. **(a)** The 2nd treatment was performed 14 d after the first one; **(b)** the 2nd treatment was performed 35 d after the first one. Fold-change values presented a standard deviation always lower than 20%.

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Fig. 3 Gene expression on tobacco plants, treated with ECAS containing 250 mg L⁻¹ of AC at pH 9.0. **(a)** 1st treatment; **(b)** the 2nd treatment was performed 14 d after the first one. Fold-change values presented a standard deviation always lower than 20%.

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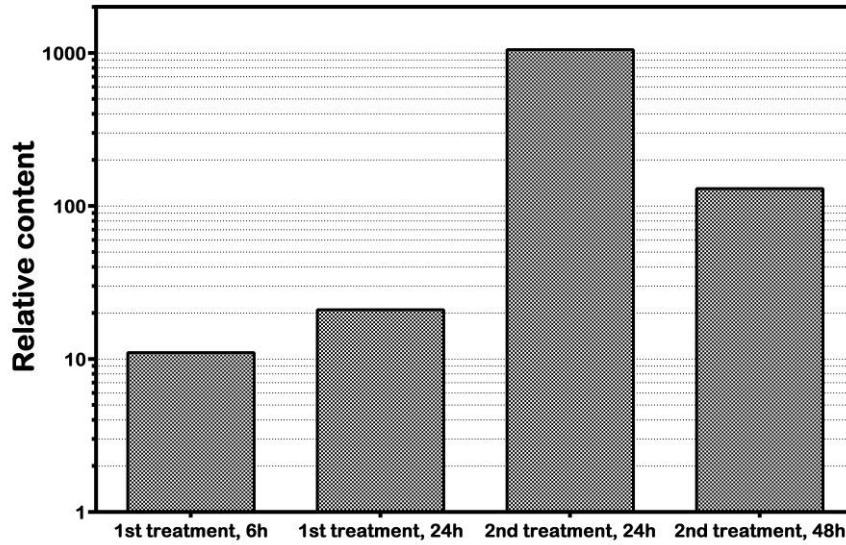


Fig. 4 Salicylic acid content in tobacco leaves sprayed with ECAS containing 250 mg L⁻¹ of AC, at pH 6.5. Data refer to the first and second treatments (after 14 d).

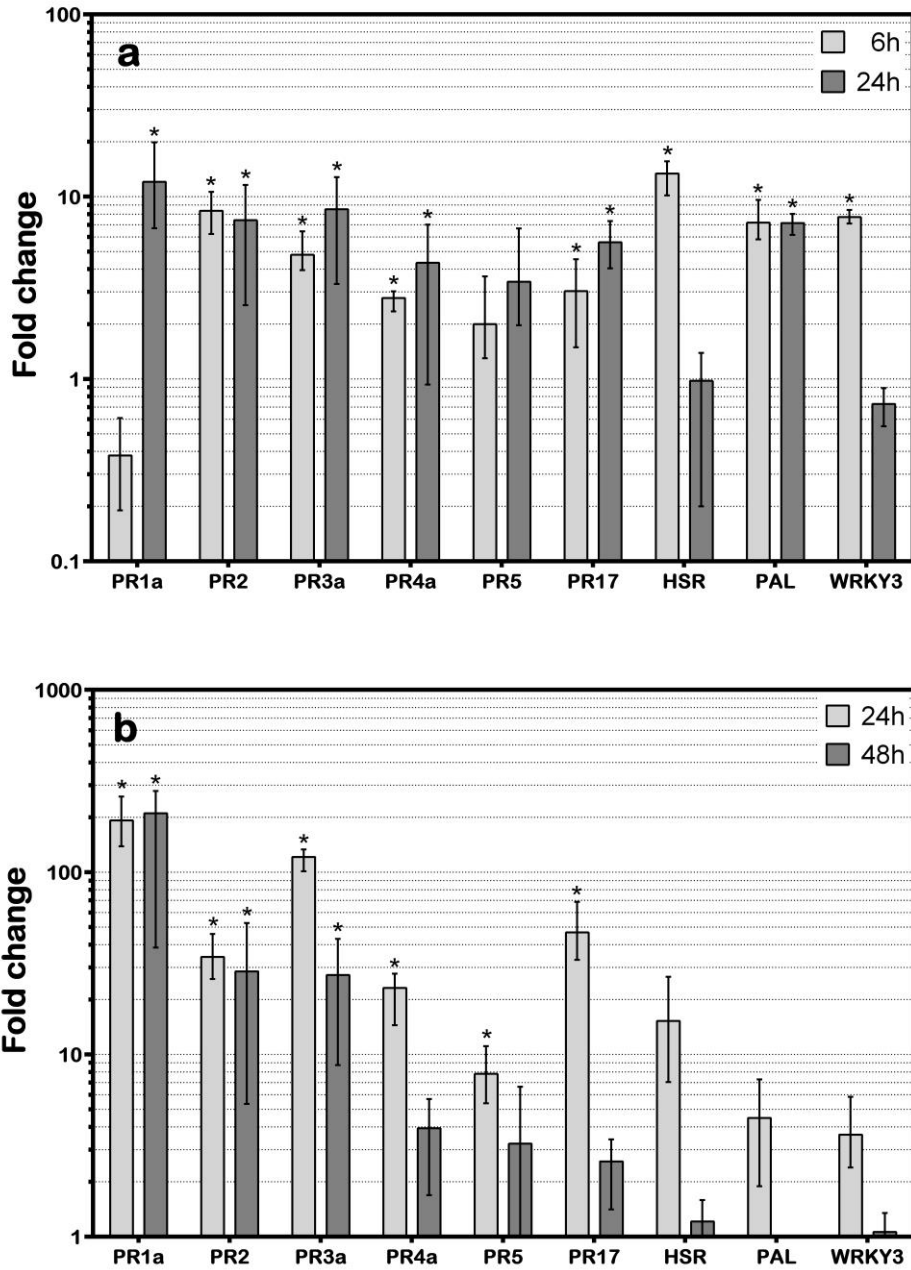







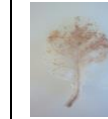



Fig. 5 Gene expression on transgenic (*NahG*) tobacco plants, treated with ECAS containing 250 mg L⁻¹ of AC at pH 6.5: (a) 1st treatment; (b) 2nd treatment (after 14 d). Stars indicate data for which a statistical significance has been ascertained according to Student's t-test ($p < 0.05$).

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a 1 st treatment				2 nd treatment (14 d after 1 st)				Positive control
Control	ECAS	Control	ECAS	Control	ECAS	Control	ECAS	Paraquat
								
2h post treat.		6h post treat.		2h post treat.		6h post treat.		




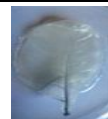




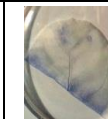
b 1 st treatment				2 nd treatment (14 d after 1 st)				Positive control
Control	ECAS	Control	ECAS	Control	ECAS	Control	ECAS	Paraquat
								
2h post treat.		6h post treat.		2h post treat.		6h post treat.		

Fig. 6 DAB and NBT histochemical essays performed on tobacco leaves sprayed with ECAS containing 250 mg L⁻¹ of AC, at pH 6.5: **(a)** a brown staining would reveal the presence of H₂O₂; **(b)** a blue staining would reveal the presence of O₂^{•-}.

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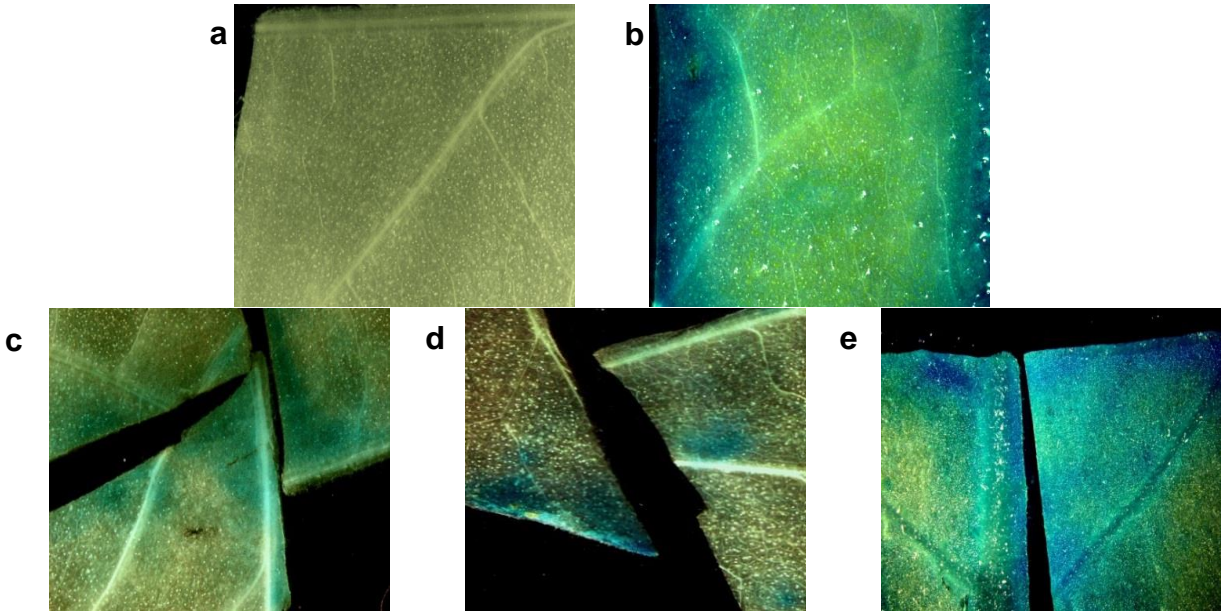


Fig. 7 GUS histochemical staining on PR1a-GUS transgenic tobacco plant after different treatments: **(a)** control solution, 24 h after 1st treatment; **(b)** positive control BTH (130 mg L⁻¹) 24 h after 1st treatment; **(c)** ECAS (250 mg L⁻¹) 24 h after 1st treatment; **(d)** ECAS (250 mg L⁻¹) 24 h after 2nd treatment, 14 d after the 1st; **(e)** ECAS 48 h after 2nd treatment, 14 d after the 1st.

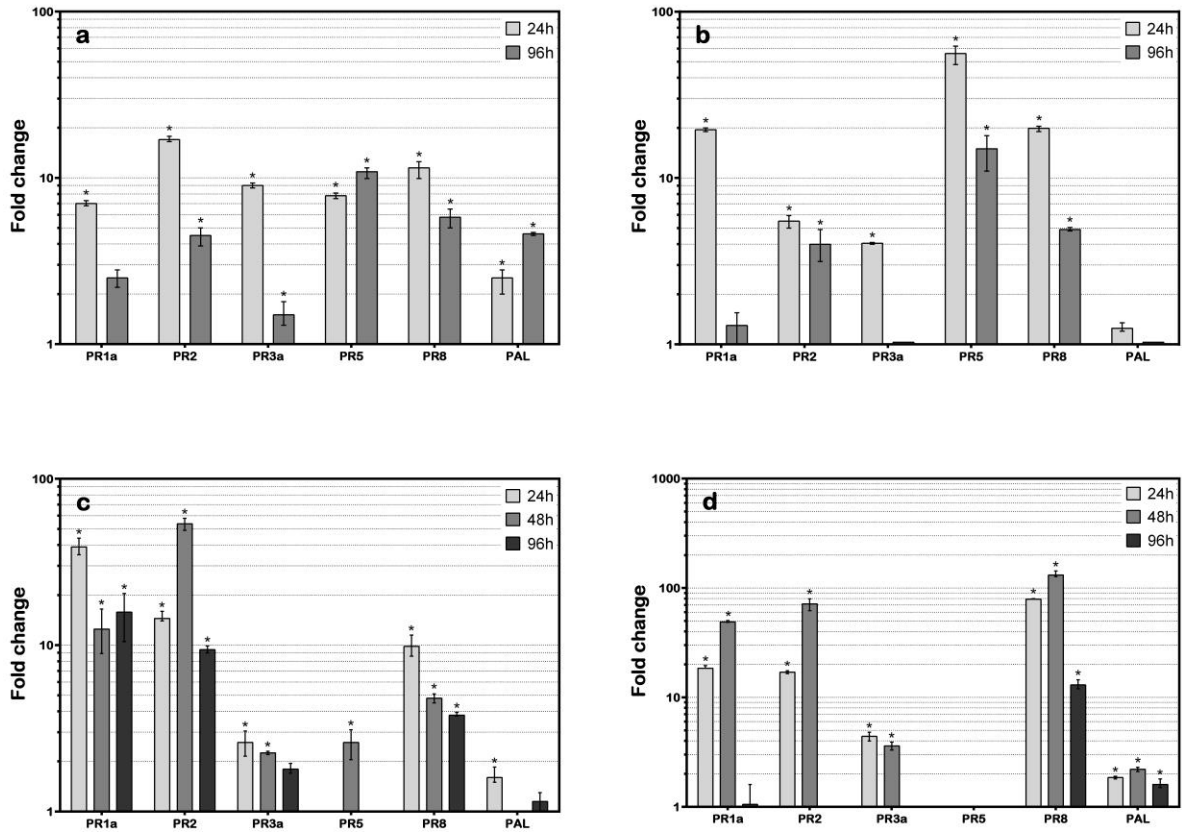


Fig. 8 Gene expression in apple plants, treated with ECAS at pH 6.5, containing either 250 mg L⁻¹ of AC (**a** and **b**, young plants) or 500 mg L⁻¹ of AC (**c** and **d**, plants in orchard); **a** and **c** refer to 1st treatment, while **b** and **d** refer to 2nd application (after 14 d). Stars indicate data for which a statistical significance has been ascertained according to Mann and Whitney's test ($p < 0.05$).

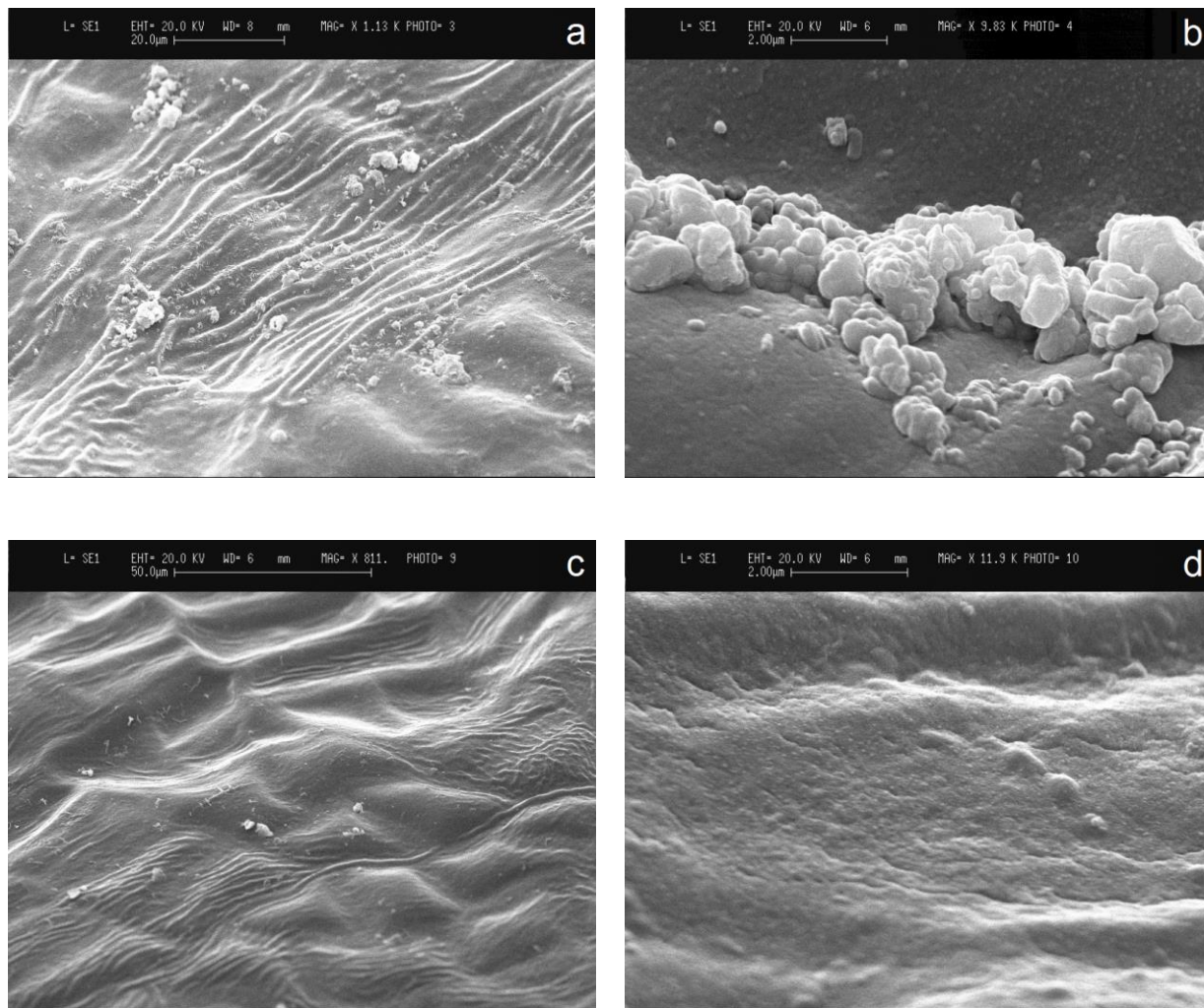


Fig. 9 SEM micrographs of apple leaves, before (**a** and **b**) and after (**c** and **d**) the treatment with ECAS at pH 6.5, containing 250 mg L⁻¹ of AC.