

UNIVERSITÀ POLITECNICA DELLE MARCHE Repository ISTITUZIONALE

Induced resistance to control postharvest decay of fruit and vegetables

This is the peer reviewd version of the followng article:

Original

Induced resistance to control postharvest decay of fruit and vegetables / Romanazzi, Gianfranco; Sanzani, S. M.; Bi, Y.; Tian, S.; Gutierrez Martinez, P.; Alkan, N.. - In: POSTHARVEST BIOLOGY AND TECHNOLOGY. - ISSN 0925-5214. - STAMPA. - 122:(2016), pp. 82-94. [10.1016/j.postharvbio.2016.08.003]

Availability:

This version is available at: 11566/249024 since: 2022-05-25T15:39:26Z

Publisher:

Published DOI:10.1016/j.postharvbio.2016.08.003

Terms of use:

The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. The use of copyrighted works requires the consent of the rights' holder (author or publisher). Works made available under a Creative Commons license or a Publisher's custom-made license can be used according to the terms and conditions contained therein. See editor's website for further information and terms and conditions. This item was downloaded from IRIS Università Politecnica delle Marche (https://iris.univpm.it). When citing, please refer to the published version.

note finali coverpage

(Article begins on next page)

1	Induced resistance to control postharvest decay of fruit and vegetables
2	
3	
4	
5	Gianfranco Romanazzi ^a , Simona Marianna Sanzani ^b , Yang Bi ^c , Shiping Tian ^d , Porfirio
6	Gutierrez-Martinez ^e , Noam Alkan ^f
7	
8	^a Department of Agricultural, Food and Environmental Sciences, Marche Polytechnic University, Via
9	Brecce Bianche, 60131 Ancona, Italy, e-mail g.romanazzi@univpm.it
10	^b Department of Soil, Plant, and Food Sciences, University of Bari Aldo Moro, Via Amendola 165/A
11	70126 Bari, Italy
12	^c College of Food Science and Engineering, Gansu Agricultural University, Lanzhou 730070, PR
13	China
14	^d Key Laboratory of Plant Resources, Institute of Botany, Chinese Academy of Sciences, Beijing
15	100093, China
16	^e Integral Laboratory on Food Science and Biotechnology Research. Technological Institute of Tepic.
17	63175. Nayarit, México
18	^f Department of Postharvest Science of Fresh Produce, Volcani Center, Agricultural Research
19	Organization, Bet Dagan, 50250, Israel
20	
21	
22	Highlights
23	
24	- Several treatments applied to fruit and vegetables can trigger host defenses
25	- Induced resistance can be systemic acquired resistance or induced systemic resistance
26	- Biocontrol agents, physical means, and natural compounds elicit host defense system
27	- Induced resistance is a strategy that contributes to control of postharvest decay
28	- Induced resistance can increase amounts of beneficial antioxidant compounds
29	
30	

31	Contents
32	
33	1. Introduction
34	2. Mechanisms involved in induced resistance
35	2.1. Systemic acquired resistance (SAR)
36	2.2. Induced systemic resistance (ISR)
37	2.3. Other mechanisms of induced resistance
38	3. Induced resistance by biocontrol agents
39	4. Induced resistance by physical means
40	4.1 UV-C irradiation
41	4.2. Heat treatment
42	4.3. Hypobaric and hyperbaric treatment
43	5. Induced resistance by natural and synthetic chemicals
44	5.1. Phytohormones and chemical elicitors
45	5.1.1. Salicylic acid and its analogs
46	5.1.2. Benzothiadiazole (BTH)
47	5.1.3. 2,6-Dichloronicotinic acid (INA)
48	5.1.4. Jasmonic acid (JA) and methyl jasmonate (MeJA)
49	5.1.5. Other chemical elicitors
50	5.2. Biological elicitors
51	5.2.1. Bacterial effectors
52	5.2.1.1. Harpin
53	5.2.1.2. Oligandrin
54	5.2.2. Fungal effectors
55	5.2.2.1. Chitosan
56	5.2.2.2. Burdock fructooligosaccharide
57	5.2.3. Other biological elicitors
58	5.3. Inorganic elicitors
59	5.3.1. Silicon (Si)
60	5.3.2. Nitric oxide (NO)
61	5.3.3. Sodium carbonate
62	6. Induced resistance by disinfecting agents
63	6.1. Ozone
64	6.2. Electrolyzed water

65	6.3. Ethanol
66	7. Induced resistance by microbial and plant volatile organic compounds (VOCs)
67	7.1. Induced resistance by microbial volatile organic compounds (MVOCs)
68	7.2. Induced resistance by plant volatile organic compounds (PVOCs)
69	8. Induced resistance perspectives
70	
71	
72	

73 Abstract

74

More than one third of harvested fruit and vegetables are lost and do not reach the customers mainly 75 due to postharvest decay. During the last decade, several postharvest fungicides have been excluded 76 from the market, or their allowed residues have been significantly decreased. Therefore, there is 77 growing interest in eco-friendly and safe alternatives to synthetic fungicides. Induced resistance has 78 79 gained increasing attention as a sustainable strategy to manage postharvest decay of fruit and vegetables. Their natural resistance can be increased by various means, such as biocontrol agents or 80 their secreted elicitors. Alternatively, physical means, such as UV-C, ozone, and heat treatment, can 81 prime plant resistance through abiotic stress. Moreover, various defense-related phytohormones, 82 biological elicitors, non-organic elicitors, and volatile organic compounds have been shown to induce 83 plant resistance. During the last decades, new technologies have enabled the evaluation of gene 84 expression, such as quantitative real time PCR and the most recent next-generation sequencing, and 85 thus the quantification of physiological changes, which have revealed new knowledge about 86 87 preharvest and postharvest induced resistance in response to various treatments. These techniques allow optimization of postharvest application of the control means, although these data cannot 88 89 disregard the evaluation of in vivo effectiveness. The elicitation of host defenses prevents the appearance of resistant isolates of pathogens. Induced resistance can lead to increased levels of 90 phenolic compounds in the plant tissues, which often have antioxidant properties that are highly 91 beneficial to humans. Moreover, induced resistance preserves the natural microflora, which is rich in 92 potential biocontrol agents, and which provides a combined approach in the control of postharvest 93 decay that is sustainable and safe for both growers and consumers. This approach meets the 94 requirements of integrated disease management on sustainable use of pesticides that in the EU is 95 implemented through Directive 128/2009. This review summarizes recent achievements and 96 knowledge of the elicitation of host defenses to control postharvest decay of fruit and vegetables, and 97 provides an outlook on the new challenges in this fascinating subject. 98

99

Keywords: biostimulants, elicitors, induced systemic resistance, resistance inducers, systemic
 acquired resistance

- 102
- 103

104 **1. Introduction**

105

Recent investigations have shown that more than one third of harvested fruit and vegetables are lost 106 (FAO, 2011; USDA, 2014; OECD, 2014). Most losses occur due to pathogen infections in the field 107 or after harvest, which lead to postharvest decay, when fruit ripen and vegetables senesce. Moreover, 108 during the last decade, several postharvest fungicides that often had wide spectra of targets have been 109 withdrawn from the market, due to: (i) selection of resistant fungal isolates; (ii) toxicity to humans 110 and the environment; (iii) increasing consumer concern toward risk of pesticide residues on products, 111 with the consequent strict requirements from several major supply chains for the quantity and number 112 of active ingredient(s) on foodstuffs, as percentages of maximum residue limit; and (iv) increasing 113 costs of registration and re-registration (Romanazzi et al., 2016a). Therefore, there is growing interest 114 in finding cheap, safe, and eco-friendly alternatives to synthetic fungicides for the control of 115 116 postharvest decay of fresh produce. Induction of plant resistance by biological, chemical, or physical means is considered a sustainable strategy to manage postharvest decay of fruit and vegetables. This 117 118 approach has gained increasing interest during recent years, in which we can see a high trend in papers dealing with induced resistance, from few ones recorded 30 years ago to more than 800 recorded 119 120 yearly in 2013-2015 (Fig. 1), and due to new tools, further knowledge has been obtained on host responses to various methods of control (Hershkovitz et al., 2013; Gapper et al., 2014). 121

The beneficial effects of induced resistance in the postharvest environment were originally 122 demonstrated about two decades ago. For example, the use of heat treatment to decrease chilling 123 injury and disease incidence in fruit through the induction of host resistance has been extensively 124 studied (Lurie and Pedreschi, 2014). Ultraviolet-C (UV-C) irradiation and exposure to sunlight have 125 been shown to induce resistance to pathogens and chilling tolerance in many harvested commodities 126 (Wilson et al., 1994; Ruan et al., 2015; Sivankalyani et al., 2016). More recently, different inducers, 127 such as cell-wall components, plant extracts, compounds of biological origin, and synthetic 128 chemicals, have been shown to trigger plant resistance to pathogen attack locally and systemically 129 (Walters and Fountaine, 2009). Moreover, biological control agents can induce plant resistance to 130 pathogens (Vallad and Goodman, 2004; Da Rocha and Hammerschmidt, 2005; Lyon, 2007). 131 However, to correctly induce resistance in different plants, it is necessary to know and understand the 132 host-microbe interactions, and the effects on postharvest physiology and handling of the different 133 fruit and vegetables (Da Rocha and Hammerschmidt, 2005). 134

Here, we review the different biological, physical, and chemical inducers that have been shown to control postharvest diseases of fruit and vegetables, and highlight their proposed mechanisms of action. 138

139 2. Mechanisms involved in induced resistance

140

Various biotic inducers (e.g., fungi, bacteria, viruses, phytoplasma, insects) and abiotic stresses (e.g., 141 chemical and physical inducers) can trigger resistance in plants, which is known as 'induced 142 resistance' (Pieterse et al., 2012; Walters et al., 2013; Pieterse et al., 2014). These can produce rapid 143 expression of defense responses (Conrath et al., 2002; Fu and Dong, 2013). Examples of treatments 144 able to induce resistance in host tissues and of representative mechanisms involved are reported in 145 Fig. 2. We can imagine induced resistance as produced by an array of treatments that elicit a cloud of 146 defense responses. There are two types of induced resistance in plants: systemic acquired resistance 147 (SAR) and induced systemic resistance (ISR). Both of these mechanisms can induce defenses that 148 confer long-lasting protection against a broad spectrum of microorganisms, and are mediated by 149 phytohormones, such as salicylic acid (SA), jasmonic acid (JA), and ethylene (ET). SAR requires the 150 signal molecule SA and is associated with accumulation of pathogenesis-related (PR) proteins, which 151 152 are believed to contribute to resistance (Durrant and Dong, 2004). Instead, the ISR pathway functions independently of SA, while it is dependent on JA and ET (van Wees et al., 1999). 153

This induced resistance does not directly activate plant defense responses, but activates the plant to a state of 'alertness', so that a future pathogen attack will be strongly and efficiently responded to. This phenomenon is also known as the 'priming effect' (Conrath et al., 2006; Jung et al., 2009), and one of the most known priming effects is root colonization by plant-growth-promoting rhizobacteria (PGPR), which induce plant development and ISR-mediated resistance (Vallad and Goodman, 2004; Verhage et al., 2010). While PGPR induces ISR, other inducers can activate SAR or both of these systems.

161

162 2.1. Systemic acquired resistance (SAR)

The mechanisms of SAR are based on SA-mediated defense. The transcription factor Nonexpressor 163 of pathogenesis-related genes 1 (NPR1) is considered to be the master regulator of SA and SAR. 164 Here, biotic, abiotic, chemical, and physical inducers can trigger defense responses locally, and can 165 also induce the production of suggested mobile immune signals, including SA, methyl salicylic acid 166 (MeSA), azelaic acid (AzA), glycerol 3-phosphate, and abietane-diterpenoid-dehydroabietinal (Park 167 et al., 2007; Chaturvedi et al., 2012). One or more of these signals can lead to systemic defense 168 'memory' that can last for weeks to months, to protect the plant from future infection (Jung et al., 169 2009). 170

Cellular redox and reactive oxygen species (ROS) are modified during SAR. Both primary 171 and secondary oxidative bursts are required for the onset of SAR (Alvarez et al., 1998). Furthermore, 172 cellular redox has been shown to be modified during SAR, as initially oxidized, and then reduced. 173 NPR1, a master regulator of SA and SAR, was shown to be reduced by thioredoxin. The NPR1 174 oligomer is disrupted and its monomers enter the nucleus, and activate SA-mediated defense (Tada 175 et al., 2008). The transcription factor NPR1 and the activated SA-mediated defense response result in 176 SAR and the activation of approximately 10% of the plant transcriptome (Fu and Dong, 2013). This 177 defense response includes direct targets of the WRKY domain transcription factor family, and 178 179 synthesis and secretion of various PR proteins, with activation of the mitogen-activated protein kinase (MAPK) cascade, the DNA repair machinery, histone modifications, and a whole arsenal of defense-180 181 related transcripts (Fu and Dong, 2013).

182

183 2.2. Induced systemic resistance (ISR)

Induced systemic resistance is known to reprogram plant-induced mechanisms based on JA and ET 184 185 defenses that alleviate physiological, abiotic, and biotic stresses. Elicitors and effectors known as 'microbe-associated molecular patterns' can be identified by plant receptors (Bent and Mackey, 186 187 2007). This recognition has a key role in activation of innate immunity. The ISR responses to biotic or abiotic stresses are diverse and can elicit plant defense responses. These responses include: MAPK 188 signaling, generation of ROS, the octadecanoic pathway (which synthesizes oxidized fatty acid 189 signals known as oxylipins), the phenylpropanoid pathway (which is involved in terpenoid and 190 phytoalexin biosynthesis), increased levels of phenolic compounds, lignification at the site of 191 pathogen infection, and cell-wall metabolism (Shoresh et al., 2010; Lloyd et al., 2011). ISR activates 192 hydrogen peroxide producers on the one hand, such as oxalate oxidase and glucose oxidase, and 193 antioxidants on the other hand, such as peroxidase (POD) and superoxide dismutase (SOD) (Shoresh 194 et al., 2010). The balance between these two determines the ROS levels, and high ROS levels can 195 196 lead to lipid peroxidation (Mittler, 2002). Additionally, several key transcripts, such as lipoxygenase (LOX1), phenylalanine ammonia lyase (PAL), and heat-shock proteins (HSPs), have been shown to 197 198 be induced during ISR (Bi et al., 2007; Shoresh et al., 2010).

199

200 2.3. Other mechanisms of induced resistance

While the main induced resistance mechanisms are SAR and ISR, some processes of induced resistance combine these two resistance mechanisms in various ways. For example, β -aminobutyric acid (BABA)-induced resistance involves both SA-dependent and abscisic acid (ABA)-dependent defense mechanisms (Buonaurio et al., 2009; Pieterse et al., 2009). The relative importance of these phytohormone-dependent defenses varies according to the nature of the challenge pathogen. Indeed,
BABA-induced resistance against *Botrytis cinerea* resembles SAR and requires SA accumulation
(Zimmerli et al., 2000), while the ABA-dependent pathway, which is associated to callose deposition,
is necessary against *Hyaloperonospora parasitica* and *Plectosphaerella cucumerina* (Zimmerli et al.,
2000; Ton and Mauch-Mani, 2004).

210

3. Induced resistance by biocontrol agents

212

213 Numerous reports have indicated that biocontrol agents, such as antagonistic yeast, can increase fruit resistance against postharvest diseases (Ippolito et al., 2000; Tian et al., 2006; Janisiewicz et al., 2008; 214 Droby et al., 2016; Spadaro and Droby, 2016) (Tab. 1). As defense responses in plants are complex 215 and involve both biochemical and structural barriers, the mechanisms of biocontrol agents are usually 216 multiple. One is the secretion of extracellular lytic enzymes, as for Pichia membranifaciens and 217 Cryptococcus albidus, where these can attach and degrade the hyphae of Monilinia fructicola, 218 219 Penicillium expansum and Rhizopus stolonifer, both in vitro and in vivo (Chan and Tian, 2005). The second is the accumulation of host PR proteins (Jijakli and Lepoivre, 1998). These are strongly 220 221 induced in response to wounding or infection by pathogens, and they accumulate abundantly at the site of infection, to contribute to SAR (Ryals et al., 1996). Treatment with Cryptococcus laurentii 222 noticeably stimulated expression of the β -1,3-glucanase (*Glu-1*) gene in jujube fruit (Tian et al., 223 2007), which suggested that *Glu-1* has a role in defense responses to fungal pathogens. C. laurentii 224 and *P. membranifaciens* have been shown to reduce disease incidence in pears (Tian et al., 2006), 225 peaches (Xu et al., 2008a), and table grapes (Meng and Tian, 2009), via enhancement of defense-226 related enzyme activities, such as chitinase (CHT), β-1,3-glucanase (GLU) and PAL. Aureobasidium 227 pullulans induced the production of CHT, GLU and POD in apple tissues starting 24 h after treatment, 228 which reached maximum levels 48 h and 96 h after treatment (Ippolito et al., 2000). Then, the 229 230 induction of host antioxidant enzymes and specific proteins has a relevant role. P. membranifaciens can induce host hydrogen peroxide metabolism, to enhance the resistance of sweet cherry against blue 231 mold by P. expansum (Chan and Tian, 2006). Additionally, four antagonistic yeasts (P. 232 membranifaciens, C. laurentii, Candida guilliermondii, Rhodotorula glutinis) can stimulate catalase 233 (CAT) and POD activity, and reduce the levels of protein carbonylation in response to ROS caused 234 by M. fructicola in peach fruit (Xu et al., 2008b). These results suggest that yeast treatments can 235 alleviate protein carbonylation and pathogen-induced oxidative damage, which implies that the 236 antioxidant defense response is involved in the mechanisms of microbial biocontrol agents against 237 238 fungal pathogens. Other yeasts, such as P. membranifaciens, C. guilliermondii, and R. glutinis, have

also been shown to stimulate the expression of both PR proteins and redox enzymes (e.g., CHT, GLU, CAT, POD) in peach fruit during all storage periods (Xu et al., 2008b). *Candida saitoana* induced postharvest systemic resistance in apple fruit against *B. cinerea*, with increased activities of CHT and GLU reported (El Ghaouth et al., 2003b). *Candida intermedia* induced strawberry fruit defense mechanisms, which included β -1,3-exoglucanase, and was effective in postharvest control of *B. cinerea* (Huang et al., 2011a). Higher PAL, CHT and GLU activities were observed when *Pichia guilliermondii* and *C. laurenti* were applied to harvested tomatoes (Zhao et al., 2008).

246

247 **4. Induced resistance by physical means**

248

Several physical means, such as ultraviolet-C (UV-C) light, heat, hypobaric and hyperbaric treatments
are known to be effective in controlling postharvest decay of fruit and vegetables (Usall et al., 2016).
The advantage of most of these relies on direct effects on the pathogen without leaving residues on
the fruit (Sanzani et al., 2009a). Moreover, these can induce several changes in host tissues, including
increased resistance to abiotic and biotic stress (Tab. 2).

254

255 4.1. UV-C irradiation

The eliciting effects of UV-C irradiation have been studied for different fruit and vegetables (Charles 256 and Arul, 2007). The first studies on the effects of UV-C irradiation on host tissues were carried out 257 with carrots, the phytoalexin content of which was increased by the treatment (Mercier et al., 1993). 258 Strawberries exposed to UV-C at 0.50 kJ m⁻² and 1.00 kJ m⁻² increased their PAL activity 12 h after 259 treatment (Nigro et al., 2000). Peach fruit treated with UV-C showed increases in PAL, CHT and 260 GLU activities (El Ghaouth et al., 2003a). Tomatoes exposed to UV-C and later inoculated with R. 261 stolonifer showed 40% reduction in polygalacturonase (PG) activity, as compared to the control 72 h 262 after the challenge (Stevens et al., 2004). The application of UV-C to harvested table grape berries 263 increased the content of trans-resveratrol and catechin in the skin (Cantos et al., 2000; Romanazzi et 264 al., 2006). A higher production of both compounds occurred when the berries were treated with 265 266 chitosan 48 h before harvest and later exposed to UV-C irradiation (Romanazzi et al., 2006).

267

268 4.2. Heat treatment

Temperature (both high and low) is one of the oldest means to control postharvest diseases of fruit and vegetables. In particular, the use of low temperature is the most diffuse means of control, and nowadays imperfect management of this cold chain can cause heavy losses of fresh produce (Romanazzi et al., 2016a). The physiological changes in host tissues induced by heat treatment were

well described by Lurie and Pedreschi (2014). In peaches exposed to hot water treatment, cell-wall 273 genes involved in ripening, such as β-galactosidase, pectin lyase, PG, and pectin methyl esterase, 274 showed general decreased expression, while ROS scavenging genes and PAL, CHT, and HSP70 275 showed increased expression (Spadoni et al., 2014). A microarray analysis run on apples treated at 276 277 45 °C for 45 min and inoculated with P. expansum showed up-regulation of HSP, HS cognate protein, and HS transcription factor genes, which were linked to resistance of the fruit to the heat stress 278 (Spadoni et al., 2015). Thus, this short hot water treatment has been shown to induce resistance to 279 chilling and pathogens in various fruit (Fallik, 2004; Lurie and Pedreschi, 2014). 280

281

282 *4.3. Hypobaric and hyperbaric treatment*

Hypobaric and hyperbaric treatments rely on the use of pressure different from atmospheric pressure 283 over a short period of time. These treatments do not have direct effects on the pathogen, although 284 285 they affect the host tissues (Romanazzi et al., 2008). For hyperbaric treatment the occurrence of physiological changes is presumed, but not yet clarified, while the variations in host tissues induced 286 287 by hypobaric applications have been well known since the middle of the last century, when they were applied to fresh fruit to delay ripening (Burg and Burg, 1966). The reduced pressure affects ET 288 289 metabolism, which reduces respiration, delays ripening, and makes the fruit less prone to decay (Lougheed et al., 1978). Short hypobaric treatment has been shown to be an effective means of control 290 of postharvest decay of strawberries, sweet cherries, and table grapes (Romanazzi et al., 2001), and 291 increased activities of PAL, CHT, and POD were observed in strawberry exposed to 0.5 atm for 4 h 292 (Hashmi et al., 2013). In contrast, the mechanisms of action of hyperbaric treatments are still not 293 clear. However, this treatment changed lycopene accumulation in tomatoes, as it reduced during 294 storage and increased during ripening (Liplap et al., 2013). 295

296

297 **5. Induced resistance by natural and synthetic chemicals**

298 5.1. Phytohormones and chemical elicitors

Phytohormones are well-known in the control of defense responses to pathogens and in the 299 300 modulation of plant induced resistance (Alkan and Fortes, 2015). A central role in the regulation of plant immune responses has been ascribed to the defense hormones SA, JA, ABA, and ET in the 301 regulation of plant-pathogen interactions (Fujita et al., 2006; Spoel and Dong, 2008). Gibberellic 302 acid, auxin indolacetic acid, brassinosteroids (BR), and cytokinines have recently emerged as 303 important modulators of plant defenses against microorganisms, mostly based on vegetative tissue 304 data and on the lifestyle of the infecting pathogen (Robert-Seilaniantz et al., 2011). The SA and JA 305 306 signaling pathways are generally considered as antagonistic and are dependent on NPR1 (Spoel et al.,

2007; Spoel and Dong, 2008; Pieterse et al., 2012). This interplay between SA and JA was suggested 307 to optimize the host response to the pathogen lifestyle (Glazebrook, 2005; Spoel and Dong, 2008; 308 Pieterse et al., 2012; Siciliano et al., 2015). In vegetative tissues, it is commonly postulated that 309 effective responses to biotrophic pathogens are typically mediated by SA and programmed cell death, 310 311 and responses to necrotrophic pathogens, which benefit from host cell death, involve JA and ET signaling (Glazebrook, 2005; Spoel and Dong, 2008). Examples of changes in gene expression or 312 enzyme activities in response to application of natural and synthetic chemicals are reported in Tab. 313 3. 314

315

316 *5.1.1. Salicylic acid and its analogs*

SA participates in the mechanisms of defense, plant development, fruit ripening, and responses to 317 various abiotic factors. SA application to an active necrotrophic infection with Colletotrichum 318 319 gloeosporioides led to programmed cell death and increased susceptibility (Alkan et al., 2012). Similarly, infection with C. gloeosporioides on a ripe NahG tomato fruit mutant that lacked SA 320 responses showed increased tolerance to C. gloeosporioides (Alkan and Fortes, 2015). Also, 321 preharvest and postharvest treatments with SA for latent infection of C. gloeosporioides effectively 322 323 reduced the occurrence of anthracnose of mango (Zainuri et al., 2001). Application of SA at 0.14 mg/mL by preharvest spraying or soaking before storage induced resistance to gray mold in kiwi fruit 324 (Poole et al., 1998). SA at 0.05 mM in combination with ultrasound induced higher disease resistance 325 to blue mold in peach fruit (Yang et al., 2011). Treatment with 0.5 mM SA reduced incidence and 326 severity of decay caused by *P. expansum* on sweet cherries (Chan and Tian, 2006). 327

Treatment with SA and its analogs induces the accumulation of ROS, which can kill pathogen 328 cells (Baker and Orlandi, 1995; Mittler et al., 2011). ROS levels are determined by the activities of 329 enzymes, such as NADPH oxidase, that are designated as 'respiratory burst oxidase homologs', and 330 antioxidants, such as SOD, ascorbate peroxidase (APX), glutathione reductase (GR), and CAT 331 332 (Sharma et al., 2012). The induced host resistance was closely related to the levels of hydrogen peroxide and the activities of antioxidant enzymes (Liu et al., 2005; Ren et al., 2012; Dickman and 333 334 Fluhr, 2013; Ge et al., 2015). On the other hand, activation of NADPH oxidase promotes the SA defense response (Alkan et al., 2012). 335

SA increased the activities of CHT, PAL, GLU and GR, and reduced the activities of CAT
and APX in pears (Cao et al., 2006). A combination of SA and an antagonistic yeast significantly
increased the activities of polyphenol oxidase (PPO), PAL, and GLU activities in cherries (Qin et al.,
2003). Additionally, in cherry, increases in the activities of CAT, POD, APX, and SOD were observed
after application of MeSA (Valverde et al., 2015).

Proteome research has shown that antioxidant and PR proteins, as well as enzymes associated with sugar metabolism, are involved in resistance of peach and sweet cherry fruit treated with SA (Chan et al., 2007). Therefore, the induced resistance by SA in fruit and vegetables activates a global defense response, which includes activation of the phenylpropanoid pathway, induction of accumulation of ROS and antioxidants, and production of PR proteins (Liu et al., 2014; Wang et al., 2015b).

347

348 5.1.2. Benzothiadiazole (BTH)

Benzothiadiazole (benzo(1,2,3)-thiadiazole-7-carbothioicacid S-methyl ester, also known as BTH or 349 ASM) is perhaps the most potent synthetic elicitor discovered to date (Terry and Joyce, 2004; Bi et 350 351 al., 2007). BTH is a light-insensitive functional analog of SA that induces resistance against a broad range of pathogens through activation of SAR in plants. Preharvest and postharvest BTH treatments 352 353 have effectively reduced latent infections and induced resistance to diseases in fruit and vegetables, including strawberries (Terry and Joyce, 2004; Mazaro et al., 2008; Cao et al., 2011; Feliziani et al., 354 355 2015), pears (Cao et al., 2006), peaches (Liu et al., 2005), melons (Ren et al., 2012; Liu et al., 2014; Li et al., 2015b), and potatoes (Bokshi et al., 2003). 356

Preharvest BTH treatment significantly reduced *Alternaria* rot and blue mold of pears during storage (Cao et al., 2005). The same authors reported that the activities of PR proteins such as POD, CHT, and GLU were significantly enhanced in pears treated with BTH. Similar results were also observed in potatoes (Bokshi et al., 2003), peaches (Liu et al., 2005), and melons (Bi et al., 2006a).

Postharvest application of BTH to strawberries induced gene expression and increased activity of a range of enzymes that included several that are linked to biotic stress resistance (Landi et al., 2014). BTH-induced disease resistance enhanced gene expression of PPO and POD, and upregulation of these genes was related to accumulation of total phenolic compounds, in harvested mango fruit (Lin et al., 2011). Postharvest dipping with BTH at 100 mg/L reduced artificial and natural infections in melons, while concentrations greater than 300 mg/L failed to promote resistance and caused phytotoxicity (Bi et al., 2006a).

368

369 5.1.3. 2,6-Dichloronicotinic acid (INA)

2,6-Dichloronicotinic acid (INA) is a synthetic compound that is a structural and functional analog
of SA, and it has been reported to mediate resistance against a broad spectrum of pathogens and its
induced resistance has been suggested to have long-lasting effects (Lucas, 1999).

373 Preharvest foliar spray of INA at 50 mg/L significantly reduced postharvest diseases of melons
374 (Bokshi et al., 2006). The resistance against *C. gloeosporioides* in mango was noticeably enhanced

by preharvest treatment with INA (Santiago et al., 2006). INA at 0.5 g/L also effectively reduced the
disease spot sizes on the peel of banana fruit when inoculated with *C. musae* (Huang et al., 2011b).

377

378 5.1.4. Jasmonic acid (JA) and methyl jasmonate (MeJA)

Application of JA and methyl jasmonate (MeJA) can control decay incidence of several fruit. Postharvest JA treatment at 0.01 mM reduced green mold of grapefruit and orange (Porat et al., 2002). The optimal concentration of JA or MeJA varies for different fruit, and for different diseases on the same fruit. The effective concentration of MeJA to control tomato anthracnose was 0.0448 mM (Tzortzakis, 2007), while it was 10 mM for the control of tomato gray mold (Zhu and Tian, 2012). When applied by infiltration, the concentration of MeJA was reduced to 0.1 mM (Yu et al., 2009).

JA stimulated production of signaling molecules related to resistance and accumulation of 385 antimicrobial compounds, and strengthened the structural barriers that restrict pathogen infection 386 387 (Tian et al., 2007). MeJA treatment promoted higher PAL activity and increased total phenolics, flavonoids, and anthocyanins (Wang et al., 2009b). MeJA treatment promoted early accumulation of 388 389 hydrogen peroxide, and increased gene expression of Cu-Zn SOD, CAT, and APX, at the same time. MeJA treatment also enhanced the contents of ascorbic acid and glutathione, which can scavenge 390 391 excess ROS to alleviate protein oxidation injury (Zhu and Tian, 2012). MeJA treatment induced resistance against *Penicillium citrinum* by priming defense responses, and up-regulated the hydrogen 392 peroxide burst and enhanced translation levels of defense-related proteins and the contents of 393 antimicrobial compounds in Chinese bayberries (Wang et al., 2014). 394

395

396 5.1.5. Other chemical elicitors

Brassinosteroids (BRs) are a group of phyto-steroidal hormones that have crucial roles in a wide spectrum of biochemical, physiological, growth, and developmental processes in plants. A remarkable feature of BRs is their potential to increase resistance to a wide spectrum of stress in plants (Krishna, 2003). BRs at 5 μ M effectively inhibited development of blue mold rot and enhanced the activities of defense-related enzymes in jujube fruit. BRs did not show in vitro antimicrobial activity against *P. expansum* (Zhu et al., 2010).

Although BABA is only rarely found naturally in plants, it has been shown to be a potent inducer of acquired resistance and has a broad spectrum of activity against many pathogens (Conrath et al., 2001). BABA induced resistance of mango fruit to postharvest anthracnose caused by *C*. *gloeosporioides*, and enhanced the activity of fruit defense mechanisms (Zhang et al., 2013). BABA caused short-lasting activation of CHT and POD after a first spray, and a boost after a second spray, although to a lower level than that caused by INA (Bokshi et al., 2006). The application of oxalic acid has been shown to induce systemic resistance against postharvest diseases (Zheng et al., 2005). Oxalic acid induced defense-related enzymes and priming of defense responses, enhanced the contents of antimicrobial compounds and modulated antioxidant enzymes in muskmelon (Deng et al., 2015). Oxalic acid at 5 mM increased antioxidant levels and PPO activity, which was beneficial for delayed ripening and enhanced disease resistance in peach fruit during storage at low temperature (Zheng et al., 2005).

L-arginine induced disease resistance via its effects on nitric oxide (NO) biosynthesis and defensive enzyme activity in tomato fruit (Zheng et al., 2011).

Riboflavin at 1.0 mM inhibited development of *Alternaria* rot, enhanced the activities of
defense-related enzymes, such as PAL, PPO and POD, and increased accumulation of flavonoids,
phenolics, and lignin (Li et al., 2012b).

420 1-Methylcyclopropene induced resistance against postharvest decay in jujube fruit, and
421 increased PAL, PPO, CAT, and SOD activities (Zhang et al., 2012).

422

423 5.2. Biological elicitors

424 5.2.1. Bacterial effectors

425 5.2.1.1. Harpin

Harpin is an acidic, heat-stable, glycine-rich, 44-kDa protein encoded by the hrpN gene, and it was 426 first described in Erwinia amylovora, which causes fire blight of Rosaceae. Bacterial harpin has been 427 shown to elicit the hypersensitive response and to induce SAR in plants (Baker and Orlandi, 1995). 428 This elicitor has also been shown to induce resistance in some postharvest fruit and vegetables (Bi et 429 al., 2007). Postharvest treatment with harpin at 0.04 mg/L to 0.16 mg/L inhibited incidence of blue 430 mold in apple (De Capdeville et al., 2003). Field spraying with harpin at 50 mg/L reduced latent 431 infections in muskmelons caused by A. alternata and Fusarium spp., with reductions proportional to 432 the levels of applied harpin, up to 90 mg/L (Wang et al., 2011b). However, harpin higher than 90 433 mg/L failed to promote resistance in melons (Bi et al., 2005). Harpin reduced lesion diameter in both 434 treated and untreated halves of the same melon, which indicated that SAR was induced by harpin (Bi 435 et al., 2005). Some of the defense reactions in melons elicited by harpin were identified as alterations 436 in the levels of preformed antifungal substances, such as phenols and flavonoids, accumulation of PR 437 proteins, such as CHT and GLU, induction of enzyme activities, such as PAL and POD, modulation 438 of metabolism of ROS, such as SOD, CAT, and hydrogen peroxide, and reinforcement of cell walls 439 and lignin (Bi et al., 2005; Wang et al., 2011b; Zhu and Zhang, 2016). 440

441

442 5.2.1.2. Oligandrin

Oligandrin is an elicitin-like protein with molecular mass of >10 kDa that has been shown to be secreted by *Pythium oligandrum*. Oligandrin is known to induce resistance against a number of plant diseases. The treatment of tomato fruit with oligandrin at 10 μ g/mL significantly reduced incidence and severity of gray mold, increased the activities of the defense-related enzymes, like PAL, PPO, and POD, and the mRNA levels of the genes encoding PR proteins, and activated the ET-dependent signaling pathway (Wang et al., 2011a).

449

450 5.2.2. Fungal effectors

451 5.2.2.1. Chitosan

Chitosan (poly- β -(1,4)-N-acetyl-d-glucosamine) is a natural biopolymer with wide antimicrobial 452 properties that can elicit defense responses in fruit and vegetables (El Ghaouth et al., 1992; 453 Romanazzi et al., 2002). This chemical and its derivatives, such as oligochitosan and glycol chitosan, 454 455 can be used in solution, as powders, and as edible coatings (Romanazzi et al., 2016b). Preharvest and postharvest treatments with chitosan and its derivatives tend to suppress storage rots in many 456 457 commodities, such as strawberry (El Ghaouth et al., 1992; Reddy et al., 2000), jujube (Yan et al., 2012), sweet cherries (Feliziani et al., 2013a), citrus fruit (Fajardo et al., 1998; Zeng et al., 2010), 458 459 apples (Felipini and Di Piero, 2009), banana (Meng et al., 2012), table grapes (Romanazzi et al., 2002; Meng et al., 2008), and tomatoes (Liu et al., 2007; Badawy and Rabea, 2009), and for many other 460 vegetables (Miranda-Castro, 2016). 461

Several studies have shown that chitosan has multiple mechanisms of action, with direct 462 antimicrobial properties, film-forming activities, and induction of host defenses (Romanazzi et al., 463 2016b). There was a significant increase in CHT and GLU activities in banana and jujube treated with 464 oligochitosan (Meng et al., 2012; Yan et al., 2012). Chitosan and oligochitosan treatments induced 465 significant increases in the activities of PPO (Liu et al., 2007), POD (Liu et al., 2007; Yan et al., 466 2012), and PAL (Romanazzi et al., 2002; Meng et al., 2012; Landi et al., 2014) in several harvested 467 products. Chitosan treatment enhanced the total content of phenolics, flavonoids, and other antifungal 468 substances (El Ghaouth et al., 1992; Yan et al., 2012), and accumulated ROS through regulation of 469 470 the activity of metabolic enzymes, such as SOD, CAT, and APX (Zeng et al., 2010; Yan et al., 2012; Landi et al., 2014). Spraying with chitosan increased the activities of PPO and PAL in table grapes, 471 thus promoting protection from latent infection of *B. cinerea* (Romanazzi et al., 2002; Meng et al., 472 2008). Next-generation sequencing with chitosan-treated avocado has defined more genes as up-473 regulated than down-regulated (Gutiérrez-Martínez et al., 2016). CHT and ROS production in table 474 grape berries varied according to the formulation of chitosan applied (Feliziani et al., 2013b). 475 Structural defense responses, such as preservation of pectin binding sites and the intense and regular 476

- 477 cellulose distribution over host cell walls were observed in the first tissue layers beneath the ruptured478 cells in bell peppers treated with chitosan (El Ghaouth et al., 1997).
- 479

480 5.2.2.2. Burdock fructooligosaccharide

Burdock fructooligosaccharide (BFO) is a fructosan oligomer that effectively inhibited postharvest diseases and reduced incidence of gray mold in tomato. BFO increased mRNA levels of genes encoding PR proteins, such as PR-1a, PR-2a (extracellular GLU), PR-2b (intracellular GLU), PR-3a (extracellular CHT), and PR-3b (intracellular CHT), and induced accumulation of PAL mRNA in tomatoes (Wang et al., 2009a). BFO also effectively controlled postharvest diseases in grapes, apples, banana, kiwi fruit, citrus fruit, strawberries, and pears (Sun et al., 2013).

487

488 *5.2.3. Other biological elicitors*

Peach fruit pretreated with yeast saccharide activated CHT, GLU, PAL, and POD. Moreover, yeast saccharide triggered endogenous NO in peaches during storage (Yu et al., 2012). The flavonoid quercetin significantly reduced blue mold in apples, and genes differentially expressed in quercetintreated apples revealed high similarities with different classes of PR proteins (i.e., RNase-like PR10, PR8), and with proteins expressed under stress conditions (Sanzani et al., 2009b; Sanzani et al., 2010). The resistance inducer protein hydrolysates were effective against green mold of citrus fruit and gray mold of table and wine grapes (Lachhab et al., 2015, 2016).

496

497 5.3. Inorganic elicitors

498 *5.3.1. Silicon (Si)*

Silicon (Si) is the second most abundant element in the Earth lithosphere, and it is as important as phosphorus and magnesium in the biota (Exley, 1998). Si is also considered to be biologically active and to trigger more rapid and extensive deployment of plant natural defenses. Guo et al. (2007) reported that Si oxide and sodium silicate suppressed pink rot in muskmelons. Sodium silicate at 100 mM reduced rots in melons caused by *A. alternata*, *F. semitectum*, and *T. roseum* (Bi et al., 2006b; Li et al., 2012a).

Sodium silicate has been shown to be effective for suppression of pathogen growth and for
induction of resistance to postharvest diseases in fruit and vegetables. Si treatment did not affect the
activities of POD, PPO, PAL, and GLU, or the content of total phenols and flavonoids in potato tuber,
although these were significantly accumulated after a challenge with *F. sulphureum* (Li et al., 2009).
These results indicated that a priming state was induced by Si in potato tubers (Conrath et al., 2001).
In Hami melons, Si treatments caused activation of POD and CHT (Bi et al., 2006b). The effects of

511 Si on postharvest diseases of muskmelons were associated with elicitation of the antioxidant system512 (Li et al., 2012a).

513

514 *5.3.2. Nitric oxide (NO)*

Nitric oxide (NO) is an important bioactive molecule that serves as a signal in plants, in particular for maturation and senescence (Leshem et al., 1998). Lai et al. (2011) reported that NO treatment activated antioxidant enzymes and induced resistance against gray mold in tomato. NO increased accumulation of phytoalexin rishitin in potato tubers (Niritake et al., 1996), and induced resistance against dry rot in potato tubers (Hu et al., 2014a). Moreover, NO treatment modulated anthracnose levels and induced defense-related enzymes in mangoes (Hu et al., 2014b).

521

522 *5.3.3. Sodium carbonate*

523 Sodium carbonate and bicarbonate were effective in the control of green mold of citrus fruit (Youssef 524 et al., 2014). These salts exerted direct antifungal effects on *Penicillium digitatum*, and increased the 525 activities of resistance enzymes, such as GLU, POD, and PAL. Moreover, citrus peel extracts showed 526 increased levels of sugars and phytoalexins, with sucrose and scoparone being the most represented.

527

528 6. Induced resistance by disinfecting agents

529 *6.1. Ozone*

Since its recognition in 1997 by the US Food and Drug Administration as a safe food disinfectant, 530 both gaseous ozone (O₃) and ozonated water have gained particular attention for the control of 531 postharvest diseases of fruit and vegetables. The most recent examples of its use are those reported 532 for blueberries (Crowe et al., 2012), persimmons (Ikeura et al., 2013), papaya (Ali et al., 2014), and 533 table grapes (Feliziani et al., 2014). Ozone is a strong oxidizing agent; however, as the O₃ disinfecting 534 activity is limited to surface-contaminating microflora, and as various microbes show different 535 susceptibilities (Pascual et al., 2007), further modes of action appear to be involved (Feliziani et al., 536 2016). Artes-Hernandez et al. (2007) reported that continuous flow of O₃ increased total flavonol and 537 hydroxycinnamates contents in cold-stored 'Autumn Seedless' table grapes. Moreover, it has been 538 reported that O₃ boosts maintained the total polyphenols, and greatly increased the phytoalexin 539 resveratrol content in cold-stored 'Napoleon' table grapes (Artés-Hernández et al., 2003). Similarly, 540 O₃ at 0.1 µmol/mol increased total phenolic content in red bell peppers (Glowacz et al., 2015). A 541 proteomic analysis conducted on O₃-treated kiwi fruit identified 102 differentially expressed proteins 542 that were mainly involved in energy, protein metabolism, defense, and cell structure (Minas et al., 543 544 2012). A set of candidate kiwifruit proteins was defined as sensitive to protein carbonylation, which

was induced by ripening and depressed by O₃. Moreover, O₃ treatment transiently increased
expression of CHT and PAL in 'Redglobe' and 'Sugraone' table grapes (Duarte-Sierra et al., 2016)
(Tab. 4).

548

549 6.2. Electrolyzed water

Electrolyzed water (EW) is obtained by adding a small amount of an electrolyte to tap water, which 550 is traditionally sodium chloride, and passing an electrical current across an anode and a cathode. EW 551 has high oxidation-reduction potential with strong direct effects against decay causing fungi 552 (Guentzel et al., 2010). However, recently, it has shown resistance inducer potential. In particular, it 553 caused 30% increase in the total phenolic content of broccoli (Navarro-Rico et al., 2014) and 554 maintained fresh-cut cilantro firmness (Hao et al., 2015), thus improving their resistance to pathogen 555 attack. More recently, other electrolytes have been tested to improve EW performance and to avoid 556 production of chlorine by-products. Encouraging results were obtained using NaHCO3 against 557 postharvest rots of citrus fruit, which also induced host defense responses (Fallanaj et al., 2016). In 558 559 particular, up-regulation of defense-related genes that encode CHT, POD, and PAL was observed at 6 h to 12 h post-treatment, with increased activity of the related enzymes and of GLU (Tab. 4). As 560 561 this was observed at 12 h post-treatment, this suggested an early host response against P. digitatum 562 by limiting tissue colonization.

563

564 *6.3. Ethanol*

The effects of ethanol dipping as a surface disinfectant that can reduce pathogen populations without 565 impairing product quality are well known (Lichter et al., 2002; Mlikota Gabler et al., 2004; Lee et al., 566 2015). However, its mode of action appears to be much more complex. Ethanol induction of 567 resistance to postharvest anthracnose in loquat fruit was demonstrated by Wang et al. (2015a). 568 Ethanol at 300 µL/L inhibited anthracnose caused by Colletotrichum acutatum, and maintained 569 570 overall quality. Moreover, it increased SOD activity, thus resulting in higher levels of hydrogen peroxide, which can activate disease resistance. Meanwhile, ethanol treatment significantly enhanced 571 the activities of defense-related enzymes, including PAL, POD, PPO, CHT, and GLU (Tab. 4). 572 Recently, ethanol treatments were shown to decelerate the ripening process and down-regulate 573 expression of major lipoxygenase-encoding genes involved in melon fruit ripening, thus contributing 574 to its increased resistance to biotic and abiotic stress (Zhang et al., 2015). 575

576

577 7. Induced resistance by microbial and plant volatile organic compounds (VOCs)

Biotic agents that contribute to plant disease management include PGPR, growth promoting fungi, and fungi that work as biocontrol agents (Lyon, 2007). In addition to the use of microorganisms and chemical compounds in plant protection, there is an important developing area of research in the control of postharvest pathogens in fruit and vegetables based on the use of volatile organic compounds (VOCs), which can be produced by microorganisms (MVOCs) or by plants (PVOCs) (Mari et al., 2016).

585

586 7.1. Induced resistance by microbial volatile organic compounds (MVOCs)

587 Microbial volatile organic compounds (MVOCs) are mixtures of carbon-based compounds that are 588 highly volatile or are vapors (Morath et al., 2012). Fungi, yeasts, and bacteria produce MVOCs as 589 primary and secondary metabolites (Jijakli and Lepoivre, 1998; Korpi et al., 2009). MVOCs have 590 shown applicative potential as biofumigants and have motivated great interest, mainly because they 591 occur naturally without chemical synthesis (Li et al., 2015a). MVOCs can be signaling substances for 592 regulation and control of some physiological actions, which include induction of systemic resistance 593 against pathogens (Ryu et al., 2004).

Some defense mechanisms activated by biocontrol bacteria are induced in fruit and leaves by
MVOCs, including production of phytoalexins, PR proteins, such as CHT and GLU, and protein
inhibitors (Conrath et al., 2006; Li et al., 2012c). *Bacillus* spp. VOCs (2-3-butanediol and lipopetides)
induced over-expression of the surfactin and fengicin genes, which caused metabolic changes in host
tissues (Ryu et al., 2004; Ongena et al., 2007).

599

600 7.2. Induced resistance by plant volatile organic compounds (PVOCs)

The eliciting activities of PVOCs, such as MeSA, were reported in section 5.1.1. Citrus fruit produce 601 VOCs that are actively involved in defense systems before pathogen attacks. The presence and 602 variation of VOCs depend on the type of produce and of its development phase. VOCs occur naturally 603 604 in plant systems and can be associated with the biochemistry of constitutive defense mechanisms (Wightwick et al., 2010). Structures in the petals of citrus flowers, called osmophores, can release 605 more than 60 VOCs, such as phenols, terpenes, and lipophilic compounds, which have been 606 recognized as antifungal agents (Caccioni et al., 1995; Lattanzio et al., 2006). It was reported that 607 limonene and linalool have antifungal actions against C. acutatum and are associated with constitutive 608 biochemical responses and can be used in the control of pathogens (Rodrigues Marques et al., 2015). 609

Essential oils produce their activity through vapor, and thyme and cinnamon essential oils increased PPO, PAL, CHT, GLU activities in peach fruit (Cindi et al., 2016). In tomato, it has been reported that VOCs are induced and modified in defense responses, and ripening processes, and by wounding. In a microarray analysis of tomato fruit at different stages of ripening, more than 4,000
differentially expressed genes have been reported. Genes related to defense mechanisms were
expressed in the early stages of ripening, and genes related to VOCs changed during late maturation.
Overall, more than 40 VOCs were detected, and their profiles were characterized along the fruit
ripening stages (Baldassarre et al., 2015).

618

619 8. Induced resistance perspectives

620

A large amount of data has been generated during the last few years that are related to the triggering 621 of host defenses during postharvest (see as examples, Tables 1, 2, 3 and 4, and Fig. 1). The elicitation 622 623 of host defenses has a central role nowadays in integrated disease management strategies, and this is welcome for the reasons summarized in Tab. 5. The effects on plants can last for weeks or months. 624 625 Induced resistance has a wider spectrum of targets than synthetic fungicides. Induced resistance does not lead to the appearance of resistant strains, due to involvement of various modes of action. Induced 626 627 resistance is usually applied before the appearance of the symptoms, so there are no side effects on nontarget organisms and on humans, in terms of farmers, people working in the packinghouses, 628 629 retailers, or consumers. Moreover, induced resistance can lead to increased levels of phenolic compounds in the plant tissues, which often have antioxidant properties that are highly beneficial to 630 humans. However, we cannot forget the weaker points linked to the application of strategies based on 631 induced resistance, such as possible inconsistent results or difficulties in their implementation in 632 packinghouse practices. Recently, a high number of biostimulants have appeared on the market, and 633 are considered as part of the effective treatments. Usually these biostimulants have combinations of 634 direct activities on pathogens and indirect activities on the host. Increasing interest in this novel 635 approach that is based on the triggering of host defenses satisfies consumer demands and the 636 guidelines on sustainable approaches to plant protection, which in European Union is implemented 637 through Directive 128/2009 on sustainable use of fungicides. New tools, such as the -omics sciences, 638 allow better understanding of the changes in host physiology and provide information on gene 639 640 functions. This information will provide optimization of the application of alternative treatments to control postharvest decay. 641

- 642
- 643

644 Acknowledgements

This work was supported by the EUBerry Project [EU FP7 KBBE 2010-4, Grant Agreement No.265942].

647 **References**

- Ali, A., Ong, M.K., Forney, C.F., 2014. Effect of ozone pre-conditioning on quality and antioxidant
 capacity of papaya fruit during ambient storage. Food Chem. 142, 19-26.
- Alkan, N., Fluhr, R., Prusky, D., 2012. Ammonium secretion during *Colletotrichum coccodes*infection modulates salicylic and jasmonic acid pathways of ripe and unripe tomato fruit. Mol.
 Plant-Microbe Interact. 25, 85-96.
- Alkan, N., Fortes, A.M., 2015. Insights into molecular and metabolic events associated with fruit
 response to postharvest fungal pathogens. Front. Plant Sci. 6.
- Alvarez, M.E., Pennell, R.I., Meijer, P.J., Ishikawa, A., Dixon, R.A., Lamb, C., 1998. Reactive
 oxygen intermediates mediate a systemic signal network in the establishment of plant immunity.
 Cell 92, 773-784.
- Artés-Hernández, F., Artes, F., Tomas-Barberan, F.A., 2003. Quality and enhancement of bioactive
 phenolics in cv. Napoleon table grapes exposed to different postharvest gaseous treatments. J.
 Agric. Food Chem. 51, 5290-5295.
- Artés-Hernández, F., Aguayo, E., Artes, F., Tomas-Barberan, F.A., 2007. Enriched ozone atmosphere
 enhances bioactive phenolics in seedless table grapes after prolonged shelf life. J. Sci. Food Agric.
 87, 824-831.
- Badawy, M.E.I., Rabea, E.I., 2009. Potential of the biopolymer chitosan with different molecular
 weights to control postharvest gray mold of tomato fruit. Postharvest Biol. Technol. 51, 110–117.
- Baker, C.J., Orlandi, E.W., 1995. Active oxygen in plant pathogenesis. Annu. Rev. Phytopathol. 33,
 299–321.
- Baldassarre, V., Cabassi, G., Spadafora, N., Aprile, A., Müller, C., Rogers, H., Ferrante, A., 2015.
 Wounding tomato fruit elicits ripening-stage specific changes in gene expression and production
 of volatile compounds. J. Exp. Bot. 66, 1511–1526.
- Bent, A.F., Mackey, D., 2007. Elicitors, effectors, and R genes: The new paradigm and a lifetime
 supply of questions. Annu. Rev. Phytopathol. 45, 399-436.
- Bi, Y., Tian, S.P., Zhao, J., Ge, Y.H., 2005. Harpin induces local and systemic resistance against
 Trichothecium roseum in harvested Hami melons. Postharvest Biol. Technol. 38, 183–187.
- Bi, Y., Ge, Y.H., Li, Y.C., Wang, J.J., Miao, X.Y., Li, X.W., 2006a. Postharvest acibenzolar-S-methyl
- treatment suppresses decay and induces resistance in Hami melons. Acta Hortic. 712, 393–399.
- Bi, Y., Tian, S.P., Guo, Y.R., Ge, Y.H., Qin, G.Z., 2006b. Sodium silicate reduces postharvest decay
- on Hami melons: induced resistance and fungistatic effects. Plant Dis. 90, 279–283.

- Bi, Y., Li, Y., Ge, Y., 2007. Induced resistance in postharvest fruits and vegetables by chemicals and
 its mechanism. Stewart Postharvest Rev. 3, 1-7.
- Bokshi, A.I., Morris, S.C., Deverall, B.J., 2003. Effects of benzothiadiazole and acetylsalicylic acid
 on β-1,3-glucanase activity and disease resistance in potato. Plant Pathol. 52, 22–27.
- Bokshi, A.I., Morris, S.C., McConchie, R.M., Deverall, B.J., 2006. Pre-harvest application of 2,6dichloroisonicotinic acid, beta-aminobutyric acid or benzothiadiazole to control post-harvest
 storage diseases of melons by inducing systemic acquired resistance (SAR). J. Hortic. Sci.
 Biotechnol. 81, 700–706.
- Buonaurio, R., Iriti, M., Romanazzi, G., 2009. Induced resistance to plant diseases caused by
 Oomycetes and fungi. Petria 19(3), 130-148.
- Burg, S.P., Burg, E.A., 1966. Fruit storage at subatmospheric pressures. Science 153, 314-315.
- Caccioni, D., Deans, L., Ruberto, G., 1995. Inhibitory effect of citrus oil components on *Penicillium italicum* and *P. digitatum*. Petria 5, 177-182.
- Cantos, E., Garcia-Viguera, C., de Pascual-Teresa, S., Tomas-Barberan, F.A., 2000. Effect of
 postharvest irradiation on resveratrol and other phenolics of cv. Napoleon table grapes. J. Agric.
 Food Chem. 48, 4606-4612.
- Cao, J.K., Jiang, W.B., He, H., 2005. Induced resistance in Yali pear (*Pyrus bretschneideri* Rehd.)
 fruit against infection by *Penicillium expansum* by postharvest infiltration of acibenzolar-Smethyl. J. Phytopathol. 153, 640-646.
- Cao, J.K., Zeng K.F., Jiang, W.B., 2006. Enhancement of postharvest disease resistance in Yali pear
 (*Pyrus bretschneideri*) fruit by salicylic acid sprays on the trees during fruit growth. Eur. J. Plant
 Pathol. 114, 363-370.
- Cao, S.F., Hu, Z., Zheng, Y., Yang, Z., Lu, B., 2011. Effect of BTH on antioxidant enzymes, radical scavenging activity and decay in strawberry fruit. Food Chem. 125, 145–149.
- Chan, Z.L., Tian, S.P. 2005. Interaction of antagonistic yeasts against postharvest pathogens of apple
 fruit and possible mode of action. Postharvest Biol. Technol. 36, 215-223.
- Chan, Z.L., Tian, S.P. 2006. Induction of H₂O₂-metabolizing enzymes and total protein synthesis by
 antagonistic yeast and salicylic acid in harvested sweet cherry fruit. Postharvest Biol. Technol.
 39, 314-320.
- 709 Chan, Z.L., Qin, G.Z., Xu, X.B., Li, B.Q., Tian, S.P. 2007. Proteome approach to characterize proteins
- induced by antagonist yeast and salicylic acid in peach fruit. J. Proteome Res. 6, 1677-1688.
- 711 Charles, M.T., Arul, J., 2007. UV treatment of fresh fruits and vegetables for improved quality: a
- status report. Stewart Postharvest Rev. 3, 6.

- Chaturvedi, R., Venables, B., Petros, R.A., Nalam, V., Li, M.Y., Wang, X.M., Takemoto, L.J., Shah,
 J., 2012. An abietane diterpenoid is a potent activator of systemic acquired resistance. Plant J.
 715 71, 161-172.
- Cindi M.D., Sivakumar D., Romanazzi G., Soundy P., 2016. Differential defense responses and
 brown rot control after essential oil fumigation in two *Prunus persica* cultivars during storage.
 Postharvest Biol. Technol. 119, 9-17.
- Conrath, U., Thulke, O., Katz, V., Schwindling, S., Kohler, A., 2001. Priming as a mechanism in induced systemic resistance of plants. Eur. J. Plant Pathol. 107, 113–119.
- Conrath, U., Pieterse, C., Mauch-Mani, B., 2002. Priming in plant–pathogen interactions. Trends
 Plant Sci. 7, 210–216.
- Conrath, U., Beckers, G.J.M., Flors, V., Garcia-Agustin, P., Jakab, G., Mauch, F., Newman, M.A.,
 Pieterse, C.M.J., Poinssot, B., Pozo, M.J., Pugin, A., Schaffrath, U., Ton, J., Wendehenne, D.,
 Zimmerli, L., Mauch-Mani, B., 2006. Priming: getting ready for battle. Mol. Plant-Microbe
 Interact. 19:1062–1071.
- Crowe, K.M., Bushway, A., Davis-Dentici, K., 2012. Impact of postharvest treatments, chlorine and
 ozone, coupled with low-temperature frozen storage on the antimicrobial quality of lowbush
 blueberries (*Vaccinium angustifolium*). LWT Food Sci. Technol. 47, 213-215.
- Da Rocha, A., Hammerschmidt, R., 2005. History and perspectives on the use of disease resistance
 inducers in horticultural crops. HortTechnology 15: 518–529.
- De Capdeville, G., Beer, S.V., Watkins, C.B., Wilson, C.L., Tedeschi, L.O., Aist, J.R., 2003. Preand postharvest harpin treatments of apples induced resistance to blue mold. Plant Dis. 89, 39–
 44.
- Deng, J., Bi, Y., Zhang, Z., Xie, D., Ge, Y., Li, W., Wang, Y., 2015. Postharvest oxalic acid treatment
 induces resistance against pink rot by priming in muskmelon (*Cucumis melo* L.) fruit. Postharvest
 Biol. Technol. 106, 53-61.
- Dickman, M.B., Fluhr, R., 2013. Centrality of host cell death in plant-microbe interactions. Annu.
 Rev. Phytopathol. 51, 543-570.
- Droby, S., Wisniewski, M., Teixidó, N., Spadaro, D., Jijakli, H.M., 2016. The science, development,
 and commercialization of postharvest biocontrol products. Postharvest Biol. Technol. (this
 issue).
- 743 Duarte-Sierra, A., Aispuro-Hernández, E., Vargas-Arispuro, I., Islas-Osuna, M.A., González-
- Aguilar, G.A., Martínez-Téllez, M. Á., 2016. Quality and PR gene expression of table grapes
- treated with ozone and sulfur dioxide to control fungal decay. J. Sci. Food Agric. 96, 2018-2014.
- 746 Durrant, W., Dong, X., 2004. Systemic acquired resistance. Annu. Rev. Phytopathol. 42, 185–209.

- El Ghaouth, A., Arul, J., Grenier, J., Asselin, A., 1992. Antifungal activity of chitosan on two
 postharvest pathogens of strawberry fruits. Phytopathology 82, 398-402.
- El Ghaouth, A., Arul, J., Wilson, C., Benhamou, N., 1997. Biochemical and cytochemical aspects of
 the interactions of chitosan and *Botrytis cinerea* in bell pepper fruit. Postharvest Biol. Technol.
 12, 183–194.
- El Ghaouth, A., Wilson, C.L., Callahan, A.M., 2003a. Induction of chitinase, β-1,3 glucanase, and
 phenylalanine ammonia-lyase in peach fruit by UV-C treatment. Phytopathology 93, 349-355.
- El Ghaouth, A., Wilson, C.L., Wisniewski, M., 2003b. Control of postharvest decay of apple fruit
 with *Candida saitoana* and induction of defense responses. Phytopathology 93, 344-348.
- Exley, C., 1998. Silicon in life: a bioinorganic solution to bioorganic essentiality. J. Inorg. Biochem.
 69, 139-144.
- Fajardo, J.E., McCollum, T.G., McDonald, R.E., Mayer, R.T., 1998. Differential induction of
 proteins in orange flavedo by biologically based elicitors and challenged by *Penicillium digitatum* Sacc. Biol Control 13, 143-151.
- Fallanaj, F., Ippolito, A., Ligorio, A., Garganese, F., Zavanella, C., Sanzani, S.M., 2016. Electrolyzed
 sodium bicarbonate inhibits *Penicillium digitatum* and induces defence responses against green
 mould in citrus fruit. Postharvest Biol. Technol. 115, 18-29.
- Fallik, E., 2004. Prestorage hot water treatments (immersion, rinsing and brushing). Postharvest Biol.
 Technol. 32, 125-134.
- FAO, 2011. Global food losses and food waste extent, causes and prevention. In: Gustavsson, J.,
 Cederberg, C., Sonesson, U., van Otterdijk, R., Meybeck, A. Rome
 (http://www.fao.org/docrep/014/mb060e/mb060e00.pdf).
- Felipini, R.B., Di Piero, R. M., 2009. Reduction of the severity of apple bitter rot by fruit immersion
 in chitosan. Pes. Agropec. Bras. 44, 1591-1597.
- Feliziani, E., Santini, M., Landi, L., Romanazzi, G., 2013a. Pre- and postharvest treatment with
 alternatives to synthetic fungicides to control postharvest decay of sweet cherry. Postharvest
 Biol. Technol. 78, 133-138.
- Feliziani, E., Smilanick, J.L., Margosan, D.A., Mansour, M.F., Romanazzi, G., Gu, H., Gohil, H.L.,
 Rubio Ames, Z., 2013b. Preharvest fungicide, potassium sorbate, or chitosan use on quality and
 storage decay of table grapes. Plant Dis. 97, 307-314.
- 777 Feliziani, E., Romanazzi, G., Smilanick, J.L., 2014. Application of low concentrations of ozone
- during the cold storage of table grapes. Postharvest Biol. Technol. 93, 38-48.

- Feliziani, E., Landi, L., Romanazzi, G., 2015. Preharvest treatments with chitosan and other
 alternatives to conventional fungicides to control postharvest decay of strawberry. Carbohydr.
 Polym. 132, 111-117.
- Feliziani, E., Lichter, A., Smilanick, J.L., Ippolito, A., 2016. Disinfecting agents for controlling fruit
 and vegetable diseases after harvest. Postharvest Biol. Technol. (this issue).
- Fu, Z.Q., Dong, X., 2013. Systemic Acquired Resistance: turning local infection into global defense.
 Annu. Rev. Plant Biol. 64, 839-863.
- Fujita, M., Fujita, Y., Noutoshi, Y., Takahashi, F., Narusaka, Y., Yamaguchi-Shinozaki, K.,
 Shinozaki, K., 2006. Crosstalk between abiotic and biotic stress responses: a current view from
 the points of convergence in the stress signaling networks. Curr. Opin. Plant Biol. 9, 436-442.
- Gapper, N.E., Giovannoni, J.J., Watkins, C.B., 2014. Understanding development and ripening of
 fruit crops in an 'omics' era. Hortic. Res. 1, 14034; doi:10.1038/hortres.2014.34.
- Ge, Y., Deng, H., Bi, Y., Li, C., Liu, Y., Dong, B., 2015. Postharvest ASM dipping and DPI pretreatment regulated reactive oxygen species metabolism in muskmelon (*Cucumis melo* L.)
 fruit. Postharvest Biol. Technol. 99, 160-167.
- Glazebrook, J., 2005. Contrasting mechanisms of defense against biotrophic and necrotrophic
 pathogens. Annu. Rev. Phytopathol. 43, 205-227.
- Glowacz, M., Colgan, R., Rees, D., 2015. Influence of continuous exposure to gaseous ozone on the
 quality of red bell peppers, cucumbers and zucchini. Postharvest Biol. Technol. 99, 1-8.
- Guentzel, J.L., Lam, K.L., Callan, M.A., Emmons, S.A., Dunham, V.L., 2010. Postharvest
 management of gray mold and brown rot on surfaces of peaches and grapes using electrolyzed
 oxidizing water. Int. J. Food Microbiol. 143, 54-60.
- Guo, Y.R., Liu, L., Zhao, J., Bi, Y., 2007. Use of silicon oxide and sodium silicate for controlling
 Trichothecium roseum postharvest rot in Chinese cantaloupe (*Cucumis melo* L.). Int. J. Food Sci.
 Technol. 42, 1012–1018.
- Gutiérrez-Martínez, P., Chacón-López, A., Xoca-Orozco, L.A., Ramos-Guerrero, A., VelázquezEstrada, R., Aguilera-Aguirre, S., 2016. Chitosan and changes in gene expression during fruit–
 pathogen interaction at postharvest stage. In: Bautista-Baños, S., Romanazzi, G., Jiménez-
- Aparicio, A. (Eds.), Chitosan in the preservation of agricultural commodities, Academic Press,
 Elsevier, London, UK, pp. 299-311.
- Hao, J., Li, H., Wan, Y. Liu, H., 2015. Effect of slightly acidic electrolyzed water (SAEW) treatment
- on the microbial reduction and storage quality of fresh-cut cilantro. J. Food Proc. Preserv. 39, 559566.

- Hashmi, M.S., East, A.R., Palmer, J.S., Heyes, J.A., 2013. Hypobaric treatment stimulates defencerelated enzymes in strawberry. Postharvest Biol. Technol. 85, 77-82.
- Hershkovitz, V., Sela, N., Taha-Salaime, L., Liu, J., Rafael, G., Kessler, C., Aly, R., Wisniewski, M.,
- 815 Droby, S., 2013. De-novo assembly and characterization of the transcriptome of *Metschnikowia*
- *fructicola* reveals differences in gene expression following interaction with *Penicillium digitatum*
- and grapefruit peel. BMC Genomics 14, 168.
- Hu, L., Li, Y., Bi, Y., Li, J., Bao, G., Liu, J., Yu, X., 2014a. Effects of nitric oxide on growth of *Fusarium sulphureum* and its virulence to potato tubers. Eur. Food Res. Technol. 238, 10071014.
- Hu, M., Yang, D., Huber, D. J., Jiang, Y., Li, M., Gao, Z., Zhang, Z., 2014b. Reduction of postharvest
 anthracnose and enhancement of disease resistance in ripening mango fruit by nitric oxide
 treatment. Postharvest Biol. Technol. 97, 115-122.
- Huang, R., Li, G.Q., Zhang, J., Yang, L., Che, H.J., Jiang, D.H., Huang, H.C., 2011a. Control of
 postharvest Botrytis fruit rot of strawberry by volatile organic compounds of *Candida intermedia*. Phytopathology 101, 859-869.
- Huang, X.M., Zhang, C., Pang X.Q., Zhang Z.Q., 2011b. Early changes of reactive oxygen species
 in 2,6-dichloroisonicotianic acid inducing tolerance in postharvest banana fruits. Acta Hortic.
 Sin.38, 265–272.
- Ikeura, H., Hamasaki, S., Tamaki, M., 2013. Effects of ozone microbubble treatment on removal of
 residual pesticides and quality of persimmon leaves. Food Chem. 138, 366-371.
- Ippolito, A., El Ghaouth, A.E., Wilson, C.L., Wisniewski, M., 2000. Control of postharvest decay of
 apple fruit by *Aureobasidium pullulans* and induction of defense responses. Postharvest Biol.
 Technol. 19, 265-272.
- Janisiewicz, W., Bastos Periera, I., Almeida, M.S., Roberts, D.P., Wisniewski, M., Kurtenbach, E.,
 2008. Improved biocontrol of fruit decay fungi with *Pichia pastoris* recombinant strains
 expressing Psd1 antifungal peptide. Postharvest Biol. Technol. 47, 218-225.
- Jijakli, M.H., Lepoivre, P., 1998. Characterization of an exo-β-1,3-glucanase produced by *Pichia anomala* strain K, antagonist of *Botrytis cinerea* on apples. Phytopathology 88, 335-343.
- Jung, H.W., Tschaplinski, T.J., Wang, L., Glazebrook, J., Greenberg, J.T., 2009. Priming in systemic
 plant immunity. Science 324, 89-91.
- Korpi A., Järnberg J., Pasanen A., 2009. Microbial volatile organic compounds. Crit. Rev. Toxicol.
 39, 139–193.
- Krishna, P., 2003. Brassinosteroid-mediated stress responses. J. Plant Growth Reg. 22, 289-297.

- Lachhab, N., Sanzani, S.M., Fallanaj, F., Youssef, K., Nigro, F., Boselli, M., Ippolito, A., 2015.
 Protein hydrolysates as resistance inducers for controlling green mould of citrus fruit. Acta
 Hortic. 1065, 1593-1598.
- Lachhab, N., Sanzani, S.M., Bahouaoui, M.A., Boselli, M., Ippolito, A., 2016. Effect of some protein
 hydrolysates against gray mould of table and wine grapes. Eur. J. Plant Pathol. 144, 821-830.
- Lai, T.F., Wang, Y.Y., Li, B.Q., Qin, G.Z., Tian, S.P., 2011. Defensive responses of tomato fruit to
 exogenous nitric oxide during postharvest storage. Postharvest Biol. Technol. 62, 127–132.
- Landi, L., Feliziani, E., Romanazzi, G., 2014. Expression of defense genes in strawberry fruit treated
 with different resistance inducers. J. Agric. Food Chem. 62, 3047-3056.
- Lattanzio, V., Cardinali, A., Lattanzio, V., 2006. Role of phenolics in the resistance mechanisms of
 plants against fungal pathogens and insects. In: Imperato, F. (Ed), Phytochemistry: advances in
 research. Research Signpost, Kerala, India, pp. 23-67.
- Lee, J.-S., Kaplunov, T., Zutahy, Y., Daus, A., Alkan, N., Lichter, A., 2015. The significance of
 postharvest disinfection for prevention of internal decay of table grapes after storage. Scientia
 Hortic. 192, 346-349.
- Leshem, Y.A.Y., Wills, R.B.H., Ku, V.V.V., 1998. Evidence for the function of the free radical gas
 nitric oxide (NO•) -as an endogenous maturation and senescence regulating factor in higher
 plants. Plant Physiol. Biochem. 36, 825–833.
- Li, Y.C., Bi, Y., Ge, Y.H., Sun, X.J., Wang, Y., 2009. Antifungal activity of sodium silicate on *Fusarium sulphureum* and its effect on dry rot of potato tubers. J. Food Sci. 74, 213–218.
- Li, W.H., Bi, Y., Ge, Y.H., Li, Y.C., Wang, J.J. Wang, Y., 2012a. Effects of postharvest sodium
 silicate treatment on pink rot disease and oxidative stress-antioxidative system in muskmelon
 fruit. Eur. Food Res. Technol. 234, 137–145.
- Li, Y.C., Yin, Y., Bi, Y., 2012b. Effect of riboflavin on postharvest disease of Asia pear and the
 possible mechanisms involved. Phytoparasitica 40, 261-268.
- Li, Q.L., Ning, P., Zheng, L., Huang, J.B., Li, G.Q., Hsiang, T., 2012c. Effects of volatile substances
 of *Streptomyces globisporus* JK-1 on control of *Botrytis cinerea* on tomato fruit. Biol. Control
 61, 113-120.
- Li, Q., Wu L., Hao J., Luo L., Cao Y., Li J., 2015a. Biofumigation on post-harvest diseases of fruits
 using a new volatile-producing fungus of *Ceratocystis fimbriata*. PLoS ONE 10(7).
- 875 Li, X., Bi, Y., Wang, J.J., Dong, B., Li, H., Gong, D., Zhao, Y., Tang, Y., Yu, X., Shang, Q., 2015b.
- BTH treatment caused physiological, biochemical and proteomic changes of muskmelon
 (*Cucumis melo* L.) fruit during ripening. J. Prot. 120, 179–193.

- Lichter, A., Zutkhy, Y., Sonego, L., Dvir, O., Kaplunov, T., Sarig, P., Ben-Arie, R., 2002. Ethanol
 controls postharvest decay of table grapes. Postharvest Biol. Technol. 24, 301–308.
- Lin, J.H., Gong, D.Q., Zhu, S.J., Zhang, L.J., Zhang, L.B., 2011. Expression of PPO and POD genes
 and contents of polyphenolic compounds in harvested mango fruits in relation to
 benzothiadiazole-induced defense against anthracnose. Scientia Hortic. 130, 85–89.
- Liplap, P., Charlebois, D., Charles, M.T., Toivonen, P., Vigneault, C., Raghavan, G.S.V., 2013.
 Tomato shelf-life extension at room temperature by hyperbaric pressure treatment. Postharvest
 Biol. Technol. 86, 45-52.
- Liu, H.X., Jiang, W.B., Bi, Y., Luo, Y.B., 2005. Postharvest BTH treatment induces resistance of
 peach (*Prunus persica* L. cv. Jiubao) fruit to infection by *Penicillium expansum* and enhances
 activity of fruit defense mechanisms. Postharvest Biol. Technol. 35, 263–269.
- Liu, J., Tian, S.P., Meng, X.H., Xu Y., 2007. Effects of chitosan on control of postharvest diseases
 and physiological responses of tomato fruit. Postharvest Biol. Technol. 44, 300–306.
- Liu, Y.Y., Ge, Y.H., Bi, Y., Li, Y.C., Deng, H.W., Hu, L.G., Dong, B., 2014. Effect of postharvest
 acibenzolar-S-methyl dipping on phenylpropanoid pathway metabolism in muskmelon (*Cucumis melo* L.) fruits. Scientia Hortic. 168, 113–119.
- Lloyd, A.J., William Allwood, J., Winder, C.L., Dunn, W.B., Heald, J.K., Cristescu, S.M.,
 Sivakumaran, A., Harren, F.J., Mulema, J., Denby, K., 2011. Metabolomic approaches reveal
 that cell wall modifications play a major role in ethylene mediated resistance against *Botrytis cinerea*. Plant J. 67, 852-868.
- Lougheed, E.C., Murr, D.P., Berard, L., 1978. Low pressure storage for horticultural crops.
 HortScience 13, 21-27.
- Lucas, J.A., 1999. Plant immunisation: from myth to SAR. Pesticide Sci. 55, 193–196.
- Lurie, S., Pedreschi, R., 2014. Fundamental aspects of postharvest heat treatments. Horticulture Res.
 1, 14030.
- Lyon, G., 2007. Agents that can elicit induced resistance. In: Walters, D., Newton, A., Lyon, G. (Eds),
 Induced resistance for plant disease control: a sustainable approach to crop protection. Blackwell
 Publishing, Oxford, pp. 9–29.
- Mari, M., Sivakumar, D., Bautista-Baños S., 2016. The role of microbial and plant volatile organic
 compounds in decay control in the postharvest system. Postharvest Biol. Technol. (This issue).
- Mazaro, S.M., Deschamps, C., de Mio, L.L., Biasi, L.A., de Gouvea, A., Sautter, C.K., 2008.
 Postharvest behavior of strawberry fruits after pre harvest treatment with chitosan and acibenzolar-S-methyl. Rev. Brasil. Frutic. 30, 185–190.

- Meng, X.H., Li, B.Q., Liu, J., Tian, S.P., 2008. Physiological responses and quality attributes of table
 grape fruit to chitosan preharvest spray and postharvest coating during storage. Food Chem. 106,
 501–508.
- Meng, X.H., Tian, S.P., 2009. Effects of preharvest application of antagonistic yeast combined with
 chitosan on decay and quality of harvested table grape fruit. J. Sci. Food Agric. 89, 1838-1842.
- 916 Meng, X.C., Tang, Y.X., Zhang, A.Y., Huang, X.M., Zhang, Z.Q., 2012. Effect of oligochitosan on
- 917 development of *Colletotrichum musae in vitro* and *in situ* and its role in protection of banana
 918 fruits. Fruits 67, 147–155.
- Mercier, J., Arul, J., Cohen, C., 1993. Effect of UV-C on phytoalexin accumulation and resistance to *Botrytis cinerea* in stored carrots. J. Phytopathol. 137, 44-54.
- 921 Minas, I. S., Tanou, G., Belghazi, M., Job, D., Manganaris, G.A., Molassiotis, A., Vasilakakis, M.,
- 2012. Physiological and proteomic approaches to address the active role of ozone in kiwifruit postharvest ripening. J Exp. Bot. 63, 2449-2464.
- Miranda-Castro, S.P., 2016. Application of chitosan in fresh and minimally processed fruits and
 vegetables. In: Bautista-Baños, S., Romanazzi, G., Jiménez-Aparicio, A. (Eds.), Chitosan in the
 Preservation of Agricultural Commodities, Academic Press, Elsevier, London, UK, pp. 67-113.
- 927 Mittler, R., 2002. Oxidative stress, antioxidants and stress tolerance. Trends Plant Sci. 7, 405-410.
- Mittler, R., Vanderauwera, S., Suzuki, N., Miller, G., Tognetti, V.B., Vandepoele, K., Gollery, M.,
 Shulaev, V., Van Breusegem, F., 2011. ROS signaling: the new wave? Trends Plant Sci. 16, 300–
 309.
- Mlikota Gabler, F., Mansour, M., Smilanick, J., Mackey, B., 2004. Survival of spores of *Rhizopus stolonifer*, *Aspergillus niger*, *Botrytis cinerea* and *Alternaria alternata* after exposure to ethanol
 solutions at various temperatures. J. Appl. Microbiol. 96, 1354–1360.
- Morath, S., Hung, R., Bennett, J., 2012. Fungal volatile organic compounds: a review with emphasis
 on their biotechnological potential. Fungal Biol. Rev. 26, 73–83.
- 936 Navarro-Rico, J., Artés-Hernández, F., Gómez, P.A., Núñez-Sánchez, M.Á., Artés, F., Martínez-
- Hernández, G.B., 2014. Neutral and acidic electrolysed water kept microbial quality and health
 promoting compounds of fresh-cut broccoli throughout shelf life. Innov. Food Sci. Emerg.
 Technol. 21, 74-81.
- Nigro, F., Ippolito, A., Lattanzio, V., Di Venere, D., Salerno, M., 2000. Effect of ultraviolet-C light
 on postharvest decay of strawberry. J. Plant Pathol. 82, 29-37.
- 942 Niritake, T., Kawakita, K., Doke, N., 1996. Nitric oxide induces phytoalexin accumulation in potato
 943 tuber tissues. Plant Cell Physiol. 37, 113–116.

- OECD, 2014. Market and trade impacts of food loss and waste reduction. Okawa K. (Ed.). Paris, 944 France 945
- (http://www.oecd.org/officialdocuments/publicdisplaydocumentpdf/?cote=TAD/CA/APM/WP(946 2014)35/FINAL&docLanguage=En) 947
- Ongena, M., Jourdan, E., Adam, A., Paquot, M., Brans, A., Joris, B., Arpigny, J.L., Thonart, P., 2007. 948 Surfactin and fengycin lipopeptides of *Bacillus subtilis* as elicitors of induced systemic resistance 949 950 in plants. Environ. Microbiol. 9, 1084–1090.
- Park, S.W., Kaimoyo, E., Kumar, D., Mosher, S., Klessig, D.F., 2007. Methyl salicylate is a critical 951 mobile signal for plant systemic acquired resistance. Science 318, 113-116. 952
- Pascual, A., Llorca, I., Canut, A., 2007. Use of ozone in food industries for reducing the 953 environmental impact of cleaning and disinfection activities. Trends Food Sci. Technol. 18, 29-954 35. 955
- Pieterse, C.M.J., Leon-Reyes, A., Van Der Ent, S., Van Wees, S.C.M., 2009. Networking by small-956 molecule hormones in plant immunity. Nat. Chem. Biol. 5, 308-316. 957
- Pieterse, C.M.J., Van der Does, D., Zamioudis, C., Leon-Reves, A., Van Wees, S.C.M., 2012. 958 Hormonal modulation of plant immunity. Annu. Rev. Cell Dev. Biol. 28, 489-521. 959
- 960 Pieterse, C.M.J., Zamioudis, C., Berendsen, R.L., Weller, D.M., Van Wees, S.C.M., Bakker, P.A.H.M., 2014. Induced systemic resistance by beneficial microbes. Annu. Rev. Phytopathol. 961 962 52:347-75.
- Poole, P.R., McLeod, L.C., Whitmore, K.J., Whitaker, G., 1998. Postharvest control of Botrytis 963 cinerea rots in stored kiwifruit. Acta Hortic. 464, 71-76. 964
- Porat, R., McCollum, T.G., Vinokur, V., Droby, S., 2002. Effects of various elicitors on the 965 transcription of a β -1,3-endoglucanase gene in citrus fruit. J. Phytopathol. 150, 70–75. 966
- Qin, G.Z., Tian, S.P., Xu, Y., Wan, Y.K., 2003. Enhancement of biocontrol efficacy of antagonistic 967 yeasts by salicylic acid in sweet cherry fruit. Physiol. Mol. Plant Pathol. 62, 147–154. 968
- 969 Reddy, B.M.V., Belkacemi, K., Corcuff, R., Castaigne, F., Arul, J., 2000. Effect of pre-harvest
- chitosan sprays on post-harvest infection by Botrytis cinerea and quality of strawberry fruit. 970 Postharvest Biol. Technol. 20, 39–51. 971
- Ren, Y.L., Wang, Y.F., Bi, Y., Ge, Y.H., Wang, Y., Fan, C.F., Li, D.Q., Deng, H.W., 2012. 972 Postharvest BTH treatment induced disease resistance and enhanced reactive oxygen species 973 974
- metabolism in muskmelon (Cucumis melo L.) fruit. Eur. Food Res. Technol. 234, 963-971.
- Robert-Seilaniantz, A., Grant, M., Jones, J.D., 2011. Hormone crosstalk in plant disease and defense: 975
- more than just jasmonate-salicylate antagonism. Annu. Rev. Phytopathol. 49, 317-343. 976

- 877 Rodrigues Marques, J.P., Amorim, L., Silva-Junior, G.J., Bellato Sposito, M., Appezzato-da Gloria,
 878 B., 2015. Structural and biochemical characteristics of citrus flowers associated with defence
 879 against a fungal pathogen. AoB PLANTS 7, plu090.
- Romanazzi, G., Nigro, F., Ippolito, A., Salerno, M., 2001. Effect of short hypobaric treatments on
 postharvest rots of sweet cherries, strawberries and table grapes. Postharvest Biol. Technol. 22,
 1-6.
- Romanazzi, G., Nigro, F., Ippolito, A., Di Venere, D., Salerno, M., 2002. Effects of pre and
 postharvest chitosan treatments to control storage grey mold of table grapes. J. Food Sci. 67,
 1862-1867.
- Romanazzi, G., Mlikota Gabler, F., Smilanick, J.L., 2006. Preharvest chitosan and postharvest UV
 irradiation treatments suppress gray mold of table grapes. Plant Dis. 90, 445-450.
- Romanazzi, G., Nigro, F., Ippolito, A., 2008. Effectiveness of short hyperbaric treatment to control
 postharvest decay of sweet cherries and table grapes. Postharvest Biol. Technol. 49, 440–442.
- Romanazzi, G., Smilanick, J.L., Feliziani, E., Droby, S., 2016a. Integrated management of
 postharvest gray mold on fruit crops. Postharvest Biol. Technol. 113, 69-76.
- Romanazzi, G., Feliziani, E., Bautista-Baños, S., Sivakumar, D., 2016b. Shelf life extension of fresh
 fruit and vegetables by chitosan treatment. Crit. Rev. Food Sci. Nutr. (in press, doi:
 10.1080/10408398.2014.900474).
- Ruan, J., Li, M., Jin, H., Sun, L., Zhu, Y., Xu, M., Dong, J., 2015. UV-B irradiation alleviates the
 deterioration of cold-stored mangoes by enhancing endogenous nitric oxide levels. Food Chem.
 169, 417-423.
- Ryals, J.A., Neuenschwander, U.H., Willits, M.G., Molina, A., Steiner, H., Hunt, M.D., 1996.
 Systemic acquired resistance. Plant Cell 8, 1809-1819.
- Ryu, C., Farag, A., Hu, C., Reddy, M., Kloepper, J., Paré, P., 2004. Bacterial volatiles induce systemic
 resistance in *Arabidopsis*. Plant Physiol. 134, 1017–1026.
- Santiago, J.A., Rivera-Vargas, L.I., Rodriguez, R.d.P., Macchiavelli, R., 2006. Resistance-inducing
 chemicals against *Colletotrichum gloeosporioides* in mango. J. Agric. Univ. Puerto Rico, 90,
 221–235.
- Sanzani, S.M., Nigro, F., Mari, M., Ippolito, A., 2009a. Innovations in the control of postharvest
 diseases of fresh fruits and vegetables. Arab J. Plant Prot. 27, 240-244.
- 1007 Sanzani, S. M., De Girolamo, A., Schena, L., Solfrizzo, M., Ippolito, A., Visconti, A., 2009b. Control
- 1008 of *Penicillium expansum* and patulin accumulation on apples by quercetin and umbelliferone.
- 1009 Eur. Food Res. Technol. 228, 381-389.

- Sanzani, S. M., Schena, L., De Girolamo, A., Ippolito, A., González-Candelas, L., 2010.
 Characterization of genes associated with induced resistance against *Penicillium expansum* in apple fruit treated with quercetin. Postharvest Biol. Technol. 56, 1-11.
- Sharma, P., Jha, A.B., Dubey, R.S., Pessarakli, M., 2012. Reactive oxygen species, oxidative damage,
 and antioxidative defense mechanism in plants under stressful conditions. J. Bot., Article ID
 217037, doi:10.1155/2012/217037.
- Shoresh, M., Harman, G.E., Mastouri, F., 2010. Induced systemic resistance and plant responses to
 fungal biocontrol agents. Annu. Rev. Phytopathol. 48, 21-43.
- Siciliano, I., Amaral Carneiro, A., Spadaro, D., Garibaldi, A., Gullino, M.L., 2015, Jasmonic acid,
 abscisic acid and salicylic acid are involved in the phytoalexin responses of rice to *Fusarium fujikuroi*, a high gibberellin producer pathogen. J. Agric. Food Chem. 63, 8134-8142.
- Sivankalyani, V., Feygenberg, O., Diskin, S., Wright, B., Alkan, N., 2016. Increased anthocyanin and
 flavonoids in mango fruit peel are associated with cold and pathogen resistance. Postharvest Biol.
- 1023Technol. 111, 132-139.
- Spadaro, D., Droby, S., 2016. Development of biocontrol products for postharvest diseases of fruit:
 the importance of elucidating the mechanisms of action of yeast antagonists. Trends Food Sci.
 Technol. 47, 39-49.
- Spadoni, A., Guidarelli, M., Sanzani, S.M., Ippolito, A., Mari, M., 2014. Influence of hot water
 treatment on brown rot of peach and rapid fruit response to heat stress. Postharvest Biol. Technol.
 94, 66-73.
- Spadoni, A., Guidarelli, M., Phillips, J., Mari, M., Wisniewski, M., 2015. Transcriptional profiling of
 apple fruit in response to heat treatment: involvement of a defense response during *Penicillium expansum*. Postharvest Biol. Technol. 101, 37-48.
- Spoel, S.H., Dong, X., 2008. Making sense of hormone crosstalk during plant immune responses.
 Cell Host Microbe 3, 348-351.
- Stevens, C., Liu, J., Khan, V.A., Lu, J.Y., Kabwe, M.K., Wilson, C.L., Igwegbe, E.C.K., Chalutz, E.,
 Droby, S., 2004. The effect of low-dose ultraviolet light-C treatment on polygalacturonase
 activity, delay ripening and Rhizopus soft rot development of tomatoes. Crop Prot. 23, 551-554.
- Sun, F., Zhang, P., Guo, M., 2013. Burdock fructooligosaccharide induces fungal resistance in
 postharvest Kyoho grapes by activating the salicylic acid-dependent pathway and inhibiting
 browning. Food Chem. 138, 539-546.
- Tada, Y., Spoel, S.H., Pajerowska-Mukhtar, K., Mou, Z., Song, J., Wang, C., Zuo, J., Dong, X., 2008.
 Plant immunity requires conformational charges of NPR1 via S-nitrosylation and thioredoxins.
- 1043 Science 321, 952-956.

- Terry, L.A., Joyce, D.C., 2004. Elicitors of induced disease resistance in postharvest horticultural
 crops: a brief review. Postharvest Biol. Technol. 32, 1–13.
- Tian, S.P., Wan, Y.K., Qin, G.Z., Xu, Y., 2006. Induction of defense responses against *Alternaria* rot
 by different elicitors in harvested pear fruit. Appl. Microbiol. Biotechnol. 70,729-734.
- Tian, S.P., Yao, H.J., Deng, X., Xu, X.B., Qin, G.Z., Chan, Z.L., 2007. Characterization and
 expression of β-1,3-glucanase genes in jujube fruit induced by the biocontrol microbial agent,
 Cryptococcus laurentii. Phytopathology 97, 260-268.
- Ton, J., Mauch-Mani, B., 2004. Beta-amino-butyric acid-induced resistance against necrotrophic
 pathogens is based on ABA-dependent priming for callose. Plant J. 38, 119-130.
- Tzortzakis, N.G., 2007. Methyl jasmonate-induced suppression of anthracnose rot in tomato fruit.
 Crop Prot. 26, 1507–1513.
- Usall, J., Ippolito, A., Sisquella, M., Neri, F., 2016. Physical treatments to control postharvest
 pathogens. Postharvest Biol. Technol. (this issue).
- USDA, 2014. The estimated amount, value, and calories of postharvest food losses at the retail and
 consumer levels in the United States, by Buzby, J.C., Wells, H.F., Hyman, J. EIB-121, US
 Department of Agriculture, Economic Research Service
 (http://www.ers.usda.gov/publications/eib-economic-information-bulletin/eib-xxx.aspx).
- Vallad, G., Goodman, R., 2004. Systemic acquired resistance and induced systemic resistance in con ventional agriculture. Crop Sci. 44, 1920–1934.
- Valverde, J., Giménez, M., Guillén, F., Valero, D., Martínez-Romero, D., Serrano, M., 2015. Methyl
 salicylate treatments of sweet cherry trees increase antioxidant systems in fruit at harvest and
 during storage. Postharvest Biol. Technol. 109, 106–113.
- Van Wees, S.M., Luijendijk, M., Smoorenburg, I., van Loon, L., Pieterse, C.J., 1999. Rhizobacteriamediated induced systemic resistance (ISR) in Arabidopsis is not associated with a direct effect
 on expression of known defense-related genes but stimulates the expression of the jasmonateinducible gene Atvsp upon challenge. Plant Mol. Biol. 41, 537-549.
- 1070 Verhage, A., Saskia, C.M., van Wees, S., Pieterse, C.M.J., 2010. Plant immunity: it's the hormones
 1071 talking, but what do they say? Plant Physiol. 154, 536-540.
- Walters, D., Fountaine, J., 2009. Practical application of induced resistance to plant diseases: an
 appraisal of effectiveness under field conditions. J. Agric. Sci. 147, 523–535.
- Walters, D.R., Ratsep, J., Havis, N.D., 2013. Controlling crop diseases using induced resistance:
 challenges for the future. J. Exp. Bot. 64, 1263-1280.
- 1076 Wang, F.D., Feng, G.H., Chen, K.S., 2009a. Defense responses of harvested tomato fruit to burdock
 1077 fructooligosaccharide, a novel potential elicitor. Postharvest Biol. Technol. 52, 110-116.

- Wang, K., Jin, P., Cao, S., Shang, H., Yang, Z., Zheng, Y., 2009b. Methyl jasmonate reduces decay
 and enhances antioxidant capacity in Chinese bayberries. J. Agric. Food Chem. 57, 5809–5815.
- Wang, A.Y., Lou, B.G., Xu, T., 2011a. Defense responses in tomato fruit induced by oligandrin
 against *Botrytis cinerea*. Afr. J. Biotechnol. 10, 4596-4601.
- Wang, J.J., Bi, Y., Zhang, Z.K., Zhang, H.Y., Ge, Y.H., 2011b. Reducing latent infection and
 enhancing disease resistance in muskmelon by preharvest application of harpin. J. Agric. Food
 Chem. 59, 12527–12533.
- Wang, K., Jin, P., Han, L., Shang, H., Tang, S., Rui, H., Zheng, Y., 2014. Methyl jasmonate induces
 resistance against *Penicillium citrinum* in Chinese bayberry by priming of defense
 responses. Postharvest Biol. Technol. 98, 90-97.
- Wang, K., Cao, S., Di, Y., Liao, Y., Zheng, Y., 2015a. Effect of ethanol treatment on disease
 resistance against anthracnose rot in postharvest loquat fruit. Scientia Hortic. 188, 115-121.
- Wang, K., Liao, Y., Cao, S., Di, H., Zheng, Y., 2015b. Effects of benzothiadiazole on disease
 resistance and soluble sugar accumulation in grape berries and its possible cellular mechanisms
 involved. Postharvest Biol. Technol. 102, 51-60.
- Wightwick, A., Walters, R., Allinson, G., Reichman, S., Menzies N., 2010. Environmental risks of
 fungicides used in horticultural production systems. In: Carisse O. (Ed.) Fungicides. In-Tech,
 Croatia, pp. 273-304.
- Wilson, C.L., El Ghaouth, A., Chalutz, E., Droby, S., Stevens, C., Lu, J.Y., Khan, V., Arul, J., 1994.
 Potential of induced resistance to control postharvest diseases of fruits and vegetables. Plant Dis.
 78, 837-844.
- Xu, X.B., Chan, Z.L., Xu, Y., Tian, S.P., 2008a. Synergistic effect of antagonist yeast and SA on
 controlling brown rot in peach fruit and its mechanism. J. Sci. Food Agric. 88, 1786-1793.
- 1101 Xu, X.B., Qin, G., Tian, S., 2008b. Effect of microbial biocontrol agents on alleviating oxidative
 1102 damage of peach fruit subjected to fungal pathogen. Int. J. Food Microbiol. 126, 153-158.
- Yan, J.Q., Cao, J.K., Jiang, W.B., Zhao, Y.M., 2012. Effects of preharvest oligochitosan sprays on
 postharvest fungal diseases, storage quality, and defensive responses in jujube (*Zizyphus jujube*Mill. cv. Dongzao) fruit. Scientia Hortic. 142, 196–204.
- Yang, Z.F., Cao, S.F., Cai, Y.T., Zheng, Y.H., 2011. Combination of salicylic acid and ultrasound to
 control postharvest blue mold caused by *Penicillium expansum* in peach fruit. Innov. Food Sci.
 Emerg. Technol. 12, 310-314.
- Youssef, K., Sanzani, S. M., Ligorio, A., Ippolito, A., Terry, L. A., 2014. Sodium carbonate and
 bicarbonate treatments induce resistance to postharvest green mould on citrus fruit. Postharvest
- 1111 Biol. Technol. 87, 61-69.

- Yu, M.M., Shen, L., Fan, B., Zhao, D.Y., Zheng, Y., Sheng, J.P., 2009. The effect of MeJA on
 ethylene biosynthesis and induced disease resistance to *Botrytis cinerea* in tomato. Postharvest
 Biol. Technol. 54, 153–158.
- Yu, Q., Chen, Q., Chen, Z.W., 2012. Activating defense responses and reducing postharvest blue
 mold decay caused by *Penicillium expansum* in peach fruit by yeast saccharide. Postharvest Biol.
 Technol. 74, 100-107.
- 1118 Zainuri, J., Joyce, D.C., Wearing, A.H., Coates, L., Terry, L., 2001. Effects of phosphonate and
- salicylic acid treatments on anthracnose disease development and ripening of 'Kensington Pride'
 mango fruit. Austral. J. Exp. Agric. 41, 805–813.
- Zeng, K.F., Deng, Y.Y., Ming, J., Deng, L.L., 2010. Induction of disease resistance and ROS
 metabolism in navel oranges by chitosan. Scientia Hortic. 126, 223-228.
- Thang, Z.Q., Tian, S.P., Zhu, Z., 2012. Effects of 1-methylcyclopropene (1-MCP) on ripening and
 resistance of jujube (*Zizyphus jujuba* cv. Huping) fruit against postharvest disease. LWT-Food
 Sci. Technol. 45, 13-19.
- Zhang, Z., Yang, D., Yang, B., Gao, Z., Li, M., Jiang, Y., Hu, M., 2013. β-Aminobutyric acid induces
 resistance of mango fruit to postharvest anthracnose caused by *Colletotrichum gloeosporioides*and enhances activity of fruit defense mechanisms. Scientia Hortic. 160, 78-84.
- Zhang, C., Shao, Q., Cao, S.X., Tang, Y.F., Liu, J.Y., Jin, Y.Z., Qi, H.Y., 2015. Effects of postharvest
 treatments on expression of three lipoxygenase genes in oriental melon (*Cucumis melo* var.
 makuwa Makino). Postharvest Biol. Technol. 110, 229-238.
- Zhao, Y., Tu, K., Shao, X., Jing, W., Su, Z., 2008. Effects of the yeast *Pichia guilliermondii* against
 Rhizopus nigricans on tomato fruit. Postharvest Biol. Technol. 49, 113-120.
- Zheng, X.L., Tian, S.P., Li, B.Q., Xu, Y., 2005. Changes in antioxidant systems and polyphenol
 oxidase activity in peach fruit treated with exogenous oxalic acid during storage at low
 temperature. Acta Hortic. Sin. 32, 788–792.
- 1137 Zheng, Y., Sheng, J.P., Zhao, R.R., 2011. Preharvest L-arginine treatment induced postharvest
 1138 disease resistance to *Botrytis cinerea* in tomato fruits. J. Agric. Food Chem. 59, 6543-6549.
- Zhu, Z., Zhang, Z.Q., Qin, G.Z., 2010. Effects of brassinosteroids on postharvest disease and
 senescence of jujube fruit in storage. Postharvest Biol. Technol. 56, 50-55.
- 1141 Zhu, Z., Tian, S.P., 2012. Resistant responses of tomato fruit treated with exogenous methyl
 1142 jasmonate to *Botrytis cinerea* infection. Scientia Hortic. 142, 38–43.
- 1143 Zhu, Z., Zhang, X., 2016. Effect of harpin on control of postharvest decay and resistant responses of
 1144 tomato fruit. Postharvest Biol. Technol. 112, 241-246.

Zimmerli, L., Jakab, C., Metraux, J.P., Mauch-Mani, B., 2000. Potentiation of pathogen specific
defense mechanisms in Arabidopsis by beta-aminobutyric acid. Proc. Natl. Acad. Sci. USA 97,
12920-12925.

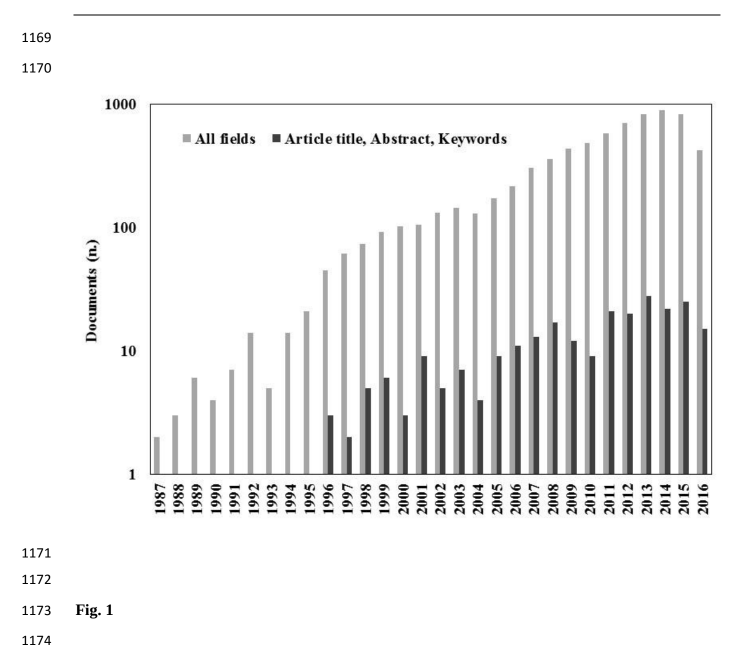
1149 **Figure captions**

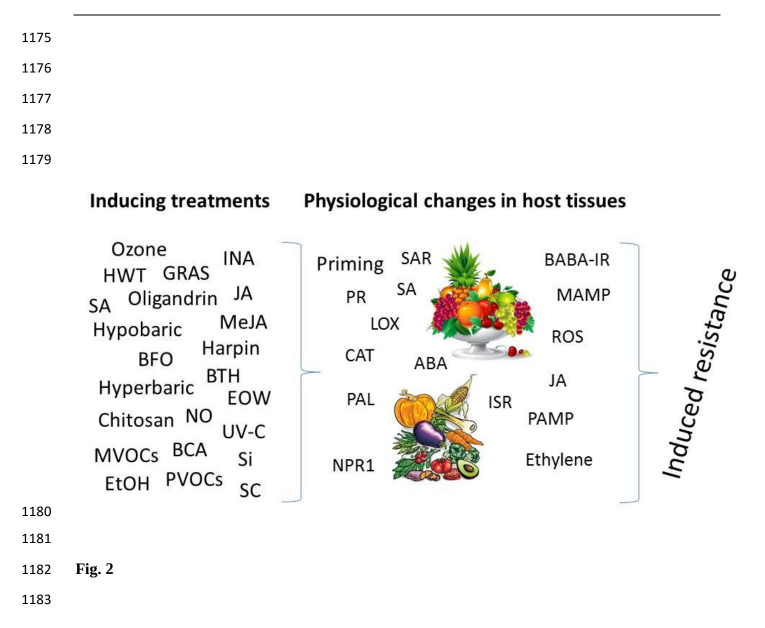
1150

Figure 1. Number of articles available through Scopus over the last 30 years using the search

- keywords of "induced resistance postharvest" (accessed on June 16, 2016).
- 1153
- **Figure 2.** Treatments that can induce resistance in fresh fruit and vegetables, and the mechanisms
- 1155 or enzymes involved. Images from http://cliparts.co/free-fruit-pictures.
- 1156 INA, 2,6-Dichloronicotinic acid; HWT, Hot water treatment; GRAS, generally recognized as safe;
- 1157 JA, jasmonic acid; SA, salicylic acid; MeJa, methyl jasmonate; BFO, Burdock
- 1158 fructooligosaccharide; BTH, benzothiadiazole; EOW, Electrolyzed oxidizing water; NO, nitric
- 1159 oxide; UV-C, ultraviolet C irradiation; BCA, biocontrol agents; MVOCs, microbial volatile organic
- 1160 compounds; Si, silicon; PVOCs, plant volatile organic compounds; EtOH, ethanol; SC, sodium
- 1161 carbonate; SAR, systemic acquired resistance; PG, polygalacturonase; BABA, β -aminobutyric acid;
- 1162 GLU, β -1,3-glucanase; PR, pathogenesis related proteins; MAMP, microbe-associated molecular
- 1163 pattern; LOX, lipoxygenase; ROS, reactive oxygen species; CHT, chitinase; CAT, catalase; ABA,
- abscisic acid; PPO, polyphenol oxidase; PAL, phenylalanine ammonia lyase; ISR, induced systemic
- resistance; PAMP, pathogen-associated molecular pattern; SOD, superoxide dismutase; NPR1,
- 1166 nonexpressor of pathogenesis-related genes 1; hsp, heat shock protein.

1167





Treatment	Genes and/or enzymes										Reference
	PAL*	CHT	SOD	CAT	SA	LOX	GLU	SURFACTIN	FENGICIN	POD	
Bacillus subtilis						+/++**		+/+++	+/+++		Ongena et al., 2007
Candida saitoana		+/++					+				El Ghaouth et al., 2003b
Cryptococcus laurentii							+				Tian et al., 2007
Pichia membranifaciens, Candida		+		+/+++			+/+++			+/++	Xu et al., 2008b
guilliermondii Rhodotorula glutinis											
Pichia guilliermondii	+	+	+	-/+			+			-/+	Zhao et al., 2008
Aureobasidium pullulans		+					+			+	Ippolito et al., 2000

Table 1. Examples of differential gene expression or enzyme activities in response to the application of biocontrol agents.

*PAL, phenylalanine ammonia lyase; CHT, chitinase; SOD, superoxide dismutase; CAT, catalase; SA, salicylic; LOX, lipoxygenase; GLU, β-1,3glucanase; POD, peroxidase

**+, overexpressed up to 3-fold; ++, overexpressed from 4-fold to 10-fold; +++, overexpressed more than 10-fold; -, down-regulated up to 3-fold

Table 2. Examples of differential gene expression or enzyme activities in response to application of physical means.

Treatment			Genes and/or	Reference			
	PAL*	CHT	GLU POD	Trans-	Catechin	-	
				resveratrol			
UV-C irradiation	+**	++/+++	+	++/+++	++	Cantos et al., 2000; Nigro et al., 2000; El	
						Ghaouth et al., 2003a; Romanazzi et al.,	
						2006	
Heat treatment	+/-	-				Spadoni et al., 2014, 2015	
Hypobaric treatment	+	+	+			Hashmi et al., 2013	

*PAL, phenylalanine ammonia lyase; CHT, chitinase; GLU, β-1,3-glucanase; POD, peroxidase

**+, overexpressed up to 3-fold; ++, overexpressed from 4-fold to 10-fold; +++, overexpressed more than 10-fold; -, down-regulated up to 3-fold

Table 3 Examples of differential gene expression or enzyme activities in response to application of natural and synthetic chemicals.

Treatment				Ge	enes and/o	or enzymes			Reference
	SOD	CAT	POD	APX	СНТ	PAL	GLU	PPO	_
Salicylic acid		_**	++	-	+	++	+		Cao et al., 2006
Methyl salicylic acid	+	++	++	+					Valverde et al., 2015
Benzothiadiazole	+	+	++		+	+	++		Cao et al., 2005
β-aminobutyric acid			+		+				Bokshi et al., 2006
Riboflavin			++			+		+	Li et al., 2012b
1-Methylcyclopropene	+	+				+		+	Zhang et al., 2012
Harpin			+++		+++				Bi et al., 2005
Oligandrin			+			+		+	Wang et al., 2011a
Chitosan			+	+/++	+/++	+/+++	+/+++	+	Romanazzi et al., 2002; Liu et al., 2007; Meng et al., 2012; Yan et al., 2012; Feliziani et al., 2013b; Landi et al., 2014
Yeast saccharide			++		+	+++	+++		Yu et al., 2012
Silicon			+++		+++				Bi et al., 2006b
Sodium carbonate			++		-	++	++		Youssef et al., 2014

SOD, superoxide dismutase; CAT, catalase; POD, peroxidase; APX, ascorbate peroxidase; CHT, chitinase; PAL, phenylalanine ammonia lyase; GLU, β-1,3-glucanase; PPO, polyphenol oxidase

**+, overexpressed up to 3-fold; ++, overexpressed from 4-fold to 10-fold; +++, overexpressed more than 10-fold; -, down-regulated up to 3-fold

Treatment		Gene	es and/	Reference			
	PPO*	GLU	PAL	LOX	POD	CHT	-
Ozone			+**			+	Duarte-Sierra et al., 2016
Electrolyzed water			+		++	+++	Fallanaj et al., 2016
Ethanol				-			Zhang et al., 2015

Table 4. Examples of differential gene expression or enzyme activities in response to application of disinfecting agents.

*PPO, polyphenol oxidase; $\overline{\text{GLU}}$, β -1,3-glucanase; PAL, phenylalanine ammonia lyase; LOX, lipoxygenase; POD, peroxidase; CHT, chitinase **+, overexpressed up to 3-fold; ++, overexpressed from 4-fold to 10-fold; +++, overexpressed more than 10-fold; -, down-regulated up to 3-fold

Negative sides	Positive sides
Complete effect is not always reproducible	Long-lasting effects
Does not provide a complete control of decay	Broad range of targets
Not easy to implement as part of farmer and	Do not cause appearance of resistant isolates of
packinghouse practices	the pathogen
Investigation methods are not standardized	Increasing number of biostimulants on the market
	Low side effects
	Reduction of pesticide use
	Promoted by EU Directive n. 128/2009
	«Sustainable Use of Pesticides» and following
	National Action Plans
	Increased amounts of beneficial antioxidant
	compounds

Table 5. Aspects related to the induction of resistance to postharvest diseases of fruit and vegetables