Physiological and proteomic approaches to address the active role of ozone in kiwifruit post-harvest ripening

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Abstract

Post-harvest ozone application has recently been shown to inhibit the onset of senescence symptoms on fleshy fruit and vegetables; however, the exact mechanism of action is yet unknown. To characterize the impact of ozone on the post-harvest performance of kiwifruit (Actinidia deliciosa cv. ‘Hayward’), fruits were cold stored (0 °C, 95% relative humidity) in a commercial ethylene-free room for 1, 3, or 5 months in the absence (control) or presence of ozone (0.3 ml l⁻¹) and subsequently were allowed to ripen at a higher temperature (20 °C), herein defined as the shelf-life period, for up to 12 days. Ozone blocked ethylene production, delayed ripening, and stimulated antioxidant and anti-radical activities of fruits. Proteomic analysis using 1D-SDS-PAGE and mass spectrometry identified 102 kiwifruit proteins during ripening, which are mainly involved in energy, protein metabolism, defence, and cell structure. Ripening induced protein carbonylation in kiwifruit but this effect was depressed by ozone. A set of candidate kiwifruit proteins that are sensitive to carbonylation was also discovered. Overall, the present data indicate that ozone improved kiwifruit post-harvest behaviour, thus providing a first step towards understanding the active role of this molecule in fruit ripening.

Key words: Actinidia deliciosa, antioxidants, anti-radical activity, ethylene, ozone, post-harvest storage, protein carbonylation.

Introduction

Fruit ripening is a complex, genetically programmed senescence process, and its elucidation relies more on a holistic than on a single process study approach (Ramina et al., 2008). Fleshy fruits have been classified as climacteric or non-climacteric, depending on whether or not a fruit exhibits a peak in respiration and ethylene production during ripening (Giovanoni, 2004). The major area of ripening research involves the post-harvest management, since horticultural industry faces huge economic losses every year as a result of ethylene-induced senescence (Martínez-Romero et al., 2007). Therefore, understanding the ripening programme is crucial to control ripening under post-harvest conditions and thus to reduce massive spoilage of fruits during storage (Michailides and Manganaris, 2009).

Ozone (O₃), also known as triatomic oxygen, is a naturally occurring highly reactive form of oxygen. Ozone signalling in the induction of cell death and defence genes and interaction with other reactive oxygen species and ethylene is well documented in plants (Castagna et al., 2007). In addition, recent studies have demonstrated the benefits of post-harvest ozone application on fleshy fruits and vegetables. However, the specific mode of action of this molecule is not yet fully understood. It has been suggested that ozone exposure prevents microbial spoilage and some diseases, decreases respiratory activity (Zhang et al., 2005), promotes ethylene oxidation in storage rooms (Skog and Chu, 2001), stimulates sucrose degradation and reduces the emission of volatile esters (Perez et al., 1999), induces the production of
antioxidant compounds with health-promoting properties (Allende et al., 2007; Artés-Hernández et al., 2007), and modulates protein and enzymatic profiles (Baur et al., 2004; Zhang et al., 2005).

Proteomics represent a rapidly developing large-scale approach to globally investigate complex biological processes such as fruit ripening. It has recently been used to describe proteome changes during ripening in many fruit species, including grape (Deytieux et al., 2007; Giribaldi et al., 2007, 2010; Negri et al., 2008; Martinez-Esteso et al., 2011), orange (Muccilli et al., 2009), strawberry (Bianco et al., 2009), peach (Nilo et al., 2010; Zhang et al., 2011), pear (Pedreschi et al., 2007; Pedreschi et al., 2008), and tomato (Rocco et al., 2006; Faurobert et al., 2007; Page et al., 2010). Proteins undergo various post-translational modifications that can affect its activity, conformation, folding, distribution, stability, and therefore its function (Rinalducci et al., 2008). Specifically, mild oxidants, which are usually produced during fruit ripening (Pedreschi et al., 2008), signal through chemical reactions with specific atoms of target proteins resulting in covalent protein modifications, including oxidation, tyrosine nitration, S-nitrosylation, glutathionylation, and disulfide formation. Importantly, protein oxidation is expected to be a major component of the oxidative signalling that regulates many physiological outputs (Moller et al., 2007; Wong et al., 2010). Amongst a variety of methods for assessing oxidative-based protein modifications, protein carboxylation has been used (Job et al., 2005). With the employment of redox proteomic approaches, several carboxylated proteins have been identified during the senescence of apple (Qin et al., 2009a) and peach (Qin et al., 2009b). However, there is still a lack of information about the proteomic hallmarks during fruit ripening, especially when taking into consideration the comparatively low number of horticultural crops with their genomes completed (Hertog et al., 2011).

Kiwifruit is an economically important crop being highly appreciated for its high levels of bioactive compounds (i.e. antioxidants) that are closely related to its ability to tolerate prolonged cold storage (Tavarini et al., 2008). Currently, the cultivation of Actinidia and the post-harvest behaviour of kiwifruit are considered to be very important in Greece. The economic importance is due to the fact that Greece is one of the largest producers of kiwifruit in the world and over one-half of Greece’s kiwifruit production is exported. Besides such socioeconomic considerations, kiwifruit provides an excellent experimental model for investigating the post-harvest action of ozone. Indeed, kiwifruit displays non-climacteric behaviour at temperatures ≤10 °C (e.g. during cold storage) and climacteric behaviour at temperatures in the order of 20 °C (e.g. during ripening in most storage/conditioning rooms following cold storage, herein referred to as the shelf life) (Antunes, 2007). In addition, recent evidence suggests that ozone application during cold storage could influence the post-harvest life of kiwifruit by suppressing the development of stem-end rot disease caused by Botrytis cinerea (Barboni et al., 2010; Minas et al., 2010).

On the other hand, a recent microarray analysis combined with transgenic approaches revealed the expression of a surprisingly large number of genes of unknown function in ripened kiwifruit (Atkinson et al., 2011), testifying to the complexity of the kiwifruit ripening process. In addition, the availability of the Actinidia expressed sequence tag (EST) database (Crowhurst et al., 2008) offers an opportunity to identify kiwifruit proteins during ripening, which, along with gene expression analysis, could globally add to the growing understanding of molecular changes underpinning climacteric fruit ripening.

This study characterized the physiological impact of ozone on qualitative and antioxidant-related properties of kiwifruit subjected to shelf-life conditions (20 °C), following short (1 month), intermediate (3 month), and extended (5 month) low-temperature storage (0 °C, 95% relative humidity). Also, it documents, as far as is known for the first time, the proteome of kiwifruit during ripening after removal from cold storage.

Materials and methods

Fruit material, experimental system, and sampling procedure

Kiwifruits (cv. ‘Hayward’), grown under standard cultural practices, were harvested from a commercial orchard (Meliki, Northern Greece) at the physiologically mature stage (Antunes, 2007) (mean weight: 95 ± 5 g, tissue firmness: 60.5 ± 2.3 N, soluble solids content: 7.9 ± 0.1% w/v). Subsequently, fruits were randomly divided into 19 lots of 30 fruits each. One lot was analysed at the time of harvest. The other lots (9 + 9) were subjected to cold storage (0 °C, 95% relative humidity) in the absence (control) or presence of continuously supplied gaseous ozone (0.3 µl l⁻¹) (ozone treatment). In both cases, a Swingtherm catalytic reactor (model BS 500, Fruit Control Equipments, Milano, Italy) was used to control ethylene concentration at a desired level in the air circulating inside the cold storage room. Fruit were removed from cold storage rooms after 1, 3, or 5 months and subsequently were transferred to shelf-life conditions (20 °C) for up to 12 days (d).

Each lot was randomly divided into three 10-fruit sub-lots. From every sub-lot, after removal of the peel, four transverse and cylindrical pieces of flesh (~4.0 g), were obtained from the equatorial region of each fruit, resulting in triplicate bulk samples per treatment. The first piece was used directly for assessment of quality parameters, the second and the third piece for the determination of fruit antioxidant profile and the fourth piece for the proteomic analysis, as described below.

Ethylene production and respiration rate were evaluated every 2 d of shelf life whilst physicochemical (tissue firmness, soluble solids content, and titratable acidity) and antioxidant properties (total antioxidant activity, ascorbic acid, and phenolic contents) were analysed after 1, 6, and 12 d of shelf life. Based on the antioxidant profiles, the anti-radical activity of kiwifruit extracts was tested at 1-d shelf life following 1, 3, or 5 months of cold storage. Finally, a proteomic approach and a Western blot (Oxyblot) analysis were used to characterize kiwifruit proteins during 1, 6, or 12 d of shelf life after 5 months of cold storage from control and ozone-treated fruit with contrasting ethylene production patterns. A scheme summarizing the experimental system and the sampling procedure is depicted in Supplementary Fig. S1 (available at JXB online).

Fruit quality and ripening parameters

Fruit weight was monitored in 20 independent fruits at harvest as well as after 1, 3, and 5 months of cold storage plus 1 d at shelf
life, and weight loss was expressed as percentage of the initial weight at harvest. Tissue firmness was recorded by puncture measurements with a Chatillon penetrometer (Chatillon and Sons, New York, NY, USA) fitted with a flat 8-mm diameter probe. After the removal of a 1-mm-thick disc of skin at the fruit equator, the tip was inserted in two opposite sides and tissue firmness was expressed in Newtons (N). Soluble solids content was assessed in juice using a refractometer (Atago PR-1, Atago Co Ltd., Tokyo, Japan) and titratable acidity by titration of 5 g juice with 10 mM NaOH to pH 8.2.

**Ethylene and respiration rate**

Ethylene production and respiration rate were monitored at 1, 2, 4, 6, 8, 10, and 12 d of shelf life after removal from 1, 3, and 5 months of cold storage, respectively. Ethylene measurements were performed with a gas chromatograph (model 3300, Varian Analytical Instruments, Walnut Creek, CA, USA), equipped with a stainless-steel column filled with Porapak (length 100 cm, diameter 0.32 cm) at 50 °C and a flame-ionization detector at 120 °C, as described (Manganaris et al., 2007). Respiration rate was calculated from CO2 production in the gas phase of the jars, measured by an infrared gas analyser (Comboy 280, David Bishop Instruments, UK).

**Phytochemical analyses**

Fruit material was extracted according to Minas et al. (2010) and the extract was used for the determination of phenol content and antioxidant activity, with the ferric ion reducing antioxidant power (FRAP) and the 1,1-diphenyl-2-picryl hydrazyl (DPPH) methods in a UV-Vis spectrophotometer (model UV-1700, Shimadzu, Kyoto, Japan). The amount of total phenolics was determined according to the Folin-Ciocalteu’s procedure (Scalbert et al., 1989; Skerget et al., 2005). Briefly, 0.5 ml extract was mixed with 2.5 ml of 1:10 diluted Folin-Ciocalteu’s phenol reagent, followed by 2.0 ml of 7.5% (w/v) Na2CO3. After 5 min at 50 °C, the absorbance was measured at 760 nm. Phenol content was estimated from a standard curve of gallic acid and results were expressed as μmol of gallic acid equivalents (GAE) of 100 g fresh weight (GAE, μmol 100 g−1 FW). For FRAP assay, a sample containing 3 ml freshly prepared FRAP solution (0.3 M acetate buffer (pH 3.6) containing 10 mM FeCl3·6H2O) and 100 μl extract was incubated at 37 °C for 4 min and the absorbance was measured at 593 nm (Benzie and Strain, 1996). The absorbance change was converted into a FRAP value by multiplying the absorbance at 593 nm of the test sample to that of the standard solution of l-ascorbic acid and the results were expressed as μmol of ascorbic acid equivalent antioxidant capacity (AEAC) of 100 g FW (AEAC, μmol 100 g−1 FW). DPPH radical scavenging activity was determined according to a modified method of Brand-Williams et al. (1995). Briefly, 50 μl extract was added into 2.95 ml of 100 μmol l−1 DPPH methanolic solution, agitated, and kept in the dark at 20 °C. The decrease in absorbance was measured at 517 nm after 30 min and the results were expressed as μmol of ascorbic acid equivalent antioxidant capacity (AEAC) of 100 g FW (AEAC, μmol 100 g−1 FW). Ascorbic acid was measured with the Merck RQflex reflectometer set (Merck, Darmstadt, Germany) (Pantelidis et al., 2007) according to the protocol for kiwifruit (Ascorbic Acid in Kiwifruit, Merck) and the results were expressed as μmol of ascorbic acid of 100 g FW (μmol 100 g−1 FW).

**Anti-radical activity**

The protective effect of kiwifruit phenols against hydroxyl radicals (HO) generated by Fenton’s reagent was estimated with the DNA nicking assay (Hu and Kitts, 2001). In brief, in a total volume of 20 μl, 0.5 μg pBR322 plasmid DNA was mixed with 4 μl of 180 μM FeCl3·6H2O, 4 μl of 18 μM ascorbic acid, 4 μl of 81 mM H2O2, and phenolic extract (0.002 μmol GAE). The reaction system was incubated in a water bath at 37 °C for 1 h. Then, the whole mixture was loaded onto a 1.5% (w/v) agarose gel and evaluated as described below.

Following the synthesis and measurement of peroxynitrite (ONOO−, 302 nm, ε = 1670 – 1 cm−1) (Uppu and Pryor, 1996), the protective activity of kiwifruit extracts toward ONOO−-induced DNA damage was performed according to Barr and Gedamu (2003). In a total volume of 25 μl, the ONOO− was added in a reaction mixture containing 50 mM sodium phosphate buffer (pH 7.0) 10 mM NaCl, 0.1 mM diethylenetriaminepentaaetic acid, 0.5 μg intact pBR322 plasmid DNA and phenolic extract (0.0025 μmol GAE). After incubation (5 min), the DNA bands were separated on a 1.5% (w/v) agarose gel.

The inhibitory effect of phenolic extracts on DNA breakage induced by peroxyl radical (ROO·) was evaluated as described by Lim et al. (2001). Here, 0.5 μg pBR322 plasmid DNA was mixed with 0.0025 μmol GAE of phenol extract in phosphate-buffered saline. An aliquot (2 μl) of 2,2’-azobis(2-amidinopropane) dihydrochloride were added to initiate the reaction (total volume 25 μl). The mixture was incubated for 2 h at 37 °C before being separated on a 1.5% (w/v) agarose gel.

In all cases, ethidium bromide-stained DNA bands were visualized under UV light, and the Image J software (National Institutes of Health; http://rsb.info.nih.gov/ij/) was employed to quantify DNA strand breaks. A correction factor of 1.4 applied to the quantitation of the signal with the supercoiled (form I) plasmid due to its lower affinity for ethidium bromide (Dong et al., 2006). All assays were run in triplicate and averaged.

**Preparation of protein extracts**

Fruit tissues without seeds (~10 g) were ground in liquid nitrogen using a mortar and pestle. Total proteins were extracted from the resulting powder at 4 °C in 10 ml buffer containing 500 mM potassium phosphate, 1 mM EDTA, 1 M NaCl, 0.5 mM Triton X-100, Complete Mini protease inhibitor cocktail tablets (Roche Molecular Biochemicals), 60 U DNase I (Roche Diagnostics), and 6 U RNase A (Sigma). After 10 min at 4 °C, 20 mM dithiothreitol (DTT, Sigma) was added and the protein extracts were stirred (20 min at 4 °C) and centrifuged (5000 g for 5 min at 4°C). Cold ethanol was added to the supernatant until 20% (v/v) final concentration, the mixtures vortexed and stored at −20 °C for 2 h. The extracts were then centrifuged (15,000 g, 15 min, 4 °C). After removal of the pellet each supernatant was clarified by boiling for 2 min then centrifuged (15,000 g, 15 min, 4 °C). The protein precipitated and protein pellet was washed three times with ice-cold acetone containing 10 mM DTT, washed once with an ice-cold solution of ethanol/ethyl acetate (1:1 v/v), and subsequently was dried under N2 flow. The proteins were resolubilized with the thiourea/urea lysis buffer as described (Harder et al., 1999; Catusse et al., 2008) containing 4% (v/v) 3-[3-Cholamidopropyl]dimethylammonio]-1-propanesulfonate, 2% (v/v) Triton X-100, and 20 mM DTT. Protein concentrations were measured according to Bradford (1976) using a Bio-Rad protein assay kit (Kit II, cat. n. 500-0002). Bovine serum albumin (Sigma) was used as a standard.

**1D-SDS-PAGE and data analysis**

Protein samples were separated by 11.5% one-dimensional sodium dodecyl sulfate-polyacrylamide gel electrophoresis (1D-SDS-PAGE) as described by Laemmli (1970) and stained with Coomassie Brilliant Blue R250 solution (Bio-Rad). Triplicate gels were used for each sample. Stained gels were scanned with a GS-800 Calibrated densitometer (Bio-Rad) and analysed with the Quantity One software (version 4.7, Bio-Rad). Protein zones were auto detected (sensitivity, 150.00: lane width, 0.8 cm: minimum density, 0.10%: noise filter, 4.00: shoulder sensitivity, 1.10: size density, 0.10%: noise filter, 4.00: shoulder sensitivity, 1.10: size
scale, 5) and, when needed, protein zones were identified manually, background subtracted and matched between biological and technical repeats, and normalized quantity determination of the zone volumes was performed (mode, relative quantity of zones in lane) along with statistical analysis.

### Western blot analysis of carbonylated proteins

Electroblotting and membrane treatments were carried out as described (Tanou et al., 2009). The membranes were then treated with 0.5 mM 2,4-dinitrophenylhydrazine (DNPH) and probed using a primary antibody against DNPH (AbD Serotec, Oxford, UK) (1:3000). Immunoreactive bands of carbonylated proteins were detected using the WesternDot 625 Western Blot Kit (Invitrogen, Molecular Probes, Eugene, Oregon, USA). The relative quantitative UV-fluorescence signal of each lane of the Western blots was estimated with the Quantity One software.

Protein oxidation levels were also quantified based on the detection of the reaction of DNPH with protein carbonyls to form protein hydrazones using a Shimadzu UV-1601 spectrophotometer according to Levine et al. (1990). Three biological replicates were performed for each assay.

### Protein identification by mass spectrometry and database analysis

Protein zones were excised from gels and digested with trypsin as described (Tanou et al., 2010). Gel pieces were rinsed with water and acetonitrile, reduced with DTT, alkylated with iodoacetamide (Sigma), and incubated in a microtube overnight at 37 °C in the presence of 3 μl of 0.5 pmol/μl trypsin (sequencing grade, Roche) in 25 mM NH₄HCO₃. The tryptic fragments were extracted, dried, and reconstituted with 0.1% (v/v) formic acid. The peptides were loaded on a C18 column (Atlantis dC18, 3 μm, 75 μm × 150 mm Nano Ease, Waters) and eluted with a 5–60% linear gradient with water/acetonitrile (98:2, v/v) containing 0.1% (v/v) formic acid (buffer A) and water/acetonitrile (20:80, v/v) containing 0.1% (v/v) formic acid (buffer B) over 30 min at a flow rate of 200 nl min⁻¹. Tryptic peptides were analysed by nano-liquid chromatography tandem mass spectrometry (LC-MS/MS, Q-TOF-Ultima Global equipped with a nano-ESI source coupled with a Cap LC nanoHPLC, Waters Micromass Waters, Saint Quentin en Yvelines, France) in the data dependent acquisition mode, allowing the selection of three precursor ions per survey scan. Only doubly and triply charged ions were selected for fragmentation over a mass range of 400–1300 m/z. A spray voltage of 3.2 kV was applied. The MS/MS raw data were processed (smooth 3/2 Savitzky Golay and deisotoping) using the ProteinLynx Global Server 2.05 software (Waters) and peak lists were exported in the micromass pkl format. Peak lists of precursor and fragment ions were matched automatically to proteins in the National Center for Biotechnology Information non-redundant database (version 2008.05.10 (6,512,701 sequences; 2,221,612,072 residues; taxonomy: Viridiplantae, 481,095 sequences)) and the GenBank viridiplantae (EST Viridiplantae sequences; 2,221,612,072 residues; taxonomy: Viridiplantae, 481,095 sequences)) and the GenBank (91,083,810 sequences), using a local version 2008.03.14 (91,083,810 sequences; 15,967,802,572 residues; taxonomy: Viridiplantae, 91,083,810 sequences), respectively.) The individual ion scores for protein or EST databanks, respectively), precursor and fragment ions. To validate protein identification, only oxidation of methionine, and 0.2 Da mass tolerance on both specificity, one missed cleavage, carbamidomethyl cysteine and 95%.

### Results

#### The impact of ozone on the ripening characteristics of kiwifruit

Kiwifruits exposed to ozone generally exhibited higher firmness retention compared with control fruits during shelf life (20 °C) after removal from 1, 3, and especially 5 months of cold storage (0 °C, 95% relative humidity) (Fig. 1A–C). An ozone-enriched cold storage atmosphere generally led to lower soluble solid contents in kiwifruits compared with control fruits (Fig. 1D–F), while titratable acidity generally remained unaffected by ozone (Fig. 1G–I). Also, ozone-treated kiwifruits exhibited lower weight loss compared with control fruits after 3 or 5 months of cold storage (Supplementary Fig. S2).

To investigate whether the ripening changes observed in response to ozone (Fig. 1) was ethylene-regulated, the production of this phytohormone was monitored in kiwifruits during shelf-life period. Ethylene production was not affected after 1 month of cold storage conducted in an ozone-enriched atmosphere compared with control fruits (Fig. 2A). However, ozone-treated fruits subjected to 3 months of cold storage exhibited a 2-d delay in the appearance of ethylene climacteric rise compared with untreated fruits (Fig. 2B). Intriguingly, ethylene remained at basal levels in ozone-treated kiwifruits exposed to a temperature of 20 °C following 5 months of cold storage, without exhibiting the typical climacteric rise observed in control fruits (Fig. 2C), indicating that ozone blocked ethylene production in these conditions. It is noted that ethylene levels within the cold storage rooms were below the generally accepted levels for long-term kiwifruit storage (10 nl l⁻¹), without differences among the cold storage conditions applied (data not shown).

A higher respiration rate was observed in ozone-treated fruits initially subjected to 1 month of cold storage and then to 1 or 2 d at 20 °C (Fig. 2D), as well as in fruits exposed to 3 months of cold storage followed by 1 d at 20 °C (Fig. 2E). In contrast, after 5 months of cold storage, ozone treatment
caused a reduction in respiration rate compared with control fruits (Fig. 2F).

The impact of ozone on antioxidant and anti-radical ability of kiwifruit during ripening

Fruits, as a multifunctional food, contain a large diversity of natural antioxidants (Sacchetti et al., 2005). In the present study, two different assay systems, namely the FRAP and DPPH assays, were employed to determine kiwifruit total antioxidant capacity (Fig. 3). The FRAP assay can measure only hydrophilic antioxidants. Therefore, compounds such as lipophilic carotenoids cannot be detected. On the other hand, the DPPH assay can detect antioxidants soluble in organic solvents such as alcohols or hydro-alcohol mixtures. However, this assay suffers from steric inaccessibility and a narrow linear range of absorbance versus concentration (Apak et al., 2007). The antioxidant activity, as evaluated with the FRAP assay, increased in kiwifruits after 12 d of shelf life following 1 month of cold storage (Fig. 3A). Also, the DPPH assay showed that fruit stored in an ozone-enriched cold atmosphere for 1 or 5 months and then exposed at 20 °C for 6 d exhibited a higher antioxidant activity than control fruits (Fig. 3D,F). Both assays indicated that the ozone treatment stimulated antioxidant capacity after 3 months of cold storage plus 12 d of shelf life (Fig. 3B,E). Notably, both antioxidant assays revealed that ozone treatment stimulated antioxidant synthesis after 1 d of shelf life (Fig. 3A–F). Furthermore, ozone-treated fruits generally exhibited higher levels of ascorbic acid content compared with control fruits, particularly after 1 d of shelf life (Fig. 4A–C). In addition, the total phenolic content was increased throughout the shelf-life period in fruits subjected to 1-month cold storage in an ozone-enriched atmosphere (Fig. 4D), as well as in fruits subjected to a 3-month exposure to ozone in cold conditions followed by 12 d of shelf life (Fig. 4E). As noted for total antioxidant capacity (Fig. 3A–F) and ascorbic acid content (Fig. 4A–C), ozone exposure for 1, 3, or 5 months in cold conditions and an additional 1 d of shelf life caused an induction in phenolic content (Fig. 4D–F).

To strengthen the validity of the present results concerning the increased phenolic content in kiwifruit initially exposed to ozone and then subjected to 1 d of shelf life, the DNA nicking assay was employed to monitor the protective effect of kiwifruit’s phenolic extracts against injury by HO, ONOO⁻, and ROO (Fig. 5). Control experiments showed that incubation of intact plasmid DNA (Fig. 5A,C,E, Intact DNA, Form I) with HO produced by Fenton’s reaction mixture (Fe⁴⁺H₂O₂/ascorbic acid), authentic ONOO⁻, or ROO generated under thermal decomposition of dihydrochloride without kiwifruit extract caused...
open circular DNA (after a single-strand break) and linear DNA (after a double-strand break) (Fig. 5A,C,E, oxidized DNA, Forms II and III, respectively). The addition of extracts obtained from control fruits, which were stored for 1, 3, or 5 months in cold conditions plus 1 d of shelf-life period, substantially reduced radical-driven DNA strand breaking compared with positive controls (Fig. 5A,C,E, L1–L3). Both in-gel and quantitative analyses further showed that the corresponding phenolic-derived samples from ozone-treated kiwifruits (Fig. 5A,C,E, L4–L6) displayed higher ability to inhibit DNA damage induced by HO (Fig. 5A) or ONOO− (Fig. 5C), as inferred from a reduction in the intensity of oxidized Forms II (Fig. 5B) and III (Fig. 5D), respectively.

**Nano-LC-MS/MS-based proteomic analysis in kiwifruit during ripening**

To complement this physiological study on the impact of ozone in kiwifruit post-harvest behaviour (Figs. 1–5), 1D-SDS-PAGE along with nano-LC-MS/MS was used to characterize the kiwifruit proteome during ripening at 20 °C. Experimental material from three independent biological replicates was collected at 1, 6, or 12 d shelf-life period following 5 months of cold storage in the absence or presence of ozone. This protocol allowed investigating the proteome of kiwifruits exhibiting contrasting ethylene patterns (Fig. 2C). A total of 31 protein zones were resolved by 1D-SDS-PAGE (1–31; marked in Fig. 6A) and variations in protein abundance within these zones upon comparing the fruit samples are shown in Supplementary Table S1. Interestingly, zones 4, 14, 15, 18, and 26 (Fig. 6A) were generally induced during ripening in both control and ozone-treated kiwifruits, whereas zones 4 and 14 showed decreased intensity of protein bands for fruit exposed to ozone compared with control fruit (Supplementary Table S1). In addition, protein zone 9 (Fig. 6A) remained unaffected during the ripening processes but was temporally and selectively modified by ozone (Supplementary Table S1). Protein zones (1–31,Fig. 6A) were excised and digested with trypsin and the resulting peptides were subjected to mass spectrometry analysis. Through this approach, a total of 102 different proteins were characterized following MASCOT searching, including two unidentified proteins (protein zones 3 and 30 in Fig. 6A). Table 1 summarizes the kiwifruit proteins identified by mass spectrometry analysis and a detailed list of these proteins is presented in Supplementary Table S2. An obvious limitation to the interpretation of the present results is the limited genomic background available in kiwifruit as only 19% of the present proteins were identified in Actinidia genus (Supplementary Fig. 2).

**Fig. 2.** Ethylene production (A–C) and respiration rate (D–F) of kiwifruits (cv. ‘Hayward’) during shelf life (20 °C) for 1, 2, 4, 6, 8, 10, and 12 days after 1 (A,D), 3 (B,E), and 5 (C,F) months of cold storage (0 °C, 95% relative humidity) in the absence (control) or presence of ozone (0.3 μl l−1). Values are means ± SE (n = 5). Asterisks indicate values that differ significantly at each time point (pairwise Student’s t-test, P < 0.05).

**Fig. 3.** Antioxidant activity, evaluated with the ferric ion reducing antioxidant power assay (A–C) and the 1,1-diphenyl-2-picryl hydrazyl assay (D–F), of kiwifruits (cv. ‘Hayward’) during shelf life (20 °C) for 1, 6, and 12 days after 1 (A,D), 3 (B,E), and 5 (C,F) months of cold storage (0 °C, 95% relative humidity) in the absence (control) or presence of ozone (0.3 μl l−1). Values are means ± SE (n = 6). Bars for each shelf-life period with different letters are statistically significant (Duncan’s multiple range test, P < 0.05).
Meanwhile, some of the identified proteins were present in more than one zone (Supplementary Table S2), indicating that these proteins are either products from different genes or arise from different post-translational protein modifications. The subcellular localization of these proteins was also evaluated based on database searches (Giribaldi et al., 2007). The largest portions of the identified kiwifruit proteins were localized in the cytosol (29%), chloroplast (17%), apoplast (14%), and mitochondria (9%) (Supplementary Table S2).

The identified kiwifruit proteins were then classified into functional categories (Fig. 6B), using gene classification resources proposed by Bevan et al. (1998). Identified kiwifruit proteins were mainly classified into the general categories of energy (31%), protein metabolism/modification (19%), plant defence (13%), and cell structure/cell wall (8%) (Fig. 6B). Several proteins associated with metabolism/ amino acid (4%), metabolism/sugars and polysaccharides (4%), cell growth/division/growth regulators (3%), and transporters/transport ATPases (3%) were also identified. An over-representation of proteins specifically involved in glycolysis (14%), the tricarboxylic acid pathway (8%), photosynthesis (5%), and electron-transport (3%) was recorded in the general set of energy-related proteins, whereas the most common in the protein metabolism/modification category were proteins involved in protein destination and storage/folding and stability (8%) and protein destination and storage/proteolysis (7%). Identified proteins that belonged to plant defence include proteins involved in disease/defence/defence-related (6%) and disease/defence/detoxification (6%). The representative kiwifruit proteins in each functional class are listed in Table 1 and are discussed individually below.

**Protein carbonylation profile in kiwifruit during ripening**

The strong impact of ripening and ozone on the antioxidant properties of kiwifruit extracts (Figs. 3–5) led to the investigation of whether the different conditions used to ripen the kiwifruits could be associated with specific features in protein oxidation patterns. To this end, kiwifruit proteins were sampled at the previously mentioned time points (Fig. 6A) and analysed in Western blot experiments based upon the specific detection of carbonylated proteins with anti-DNP antibodies. Analysis of kiwifruit proteins by 1D-SDS-PAGE followed by immunological detection using the Oxyblots revealed five carbonyl-reactive protein zones (Fig. 7A), indicating the involvement of oxidative stress in the kiwifruit ripening process. The carbonylated protein zones exhibited low molecular masses, ranging from ≈17.0 to 27.0 kDa. By matching immunopositive polypeptides (Fig. 7A) to corresponding zones on Coomassie Blue stained gels (Fig. 6A; zones 23, 24, 25, 26, and 28), 14 proteins were tentatively proposed as representing candidate carbonylated proteins (Supplementary Table S2). Quantitation of the immunoblot results by densitometry revealed an overall increased level of protein carbonylation during kiwifruit ripening (Supplementary Fig. S3). In agreement with this, quantitative measurements of protein carbonyl group content (Fig. 7B) confirmed that the extent of protein carbonylation increased during the ripening process. However, the protein carbonyl level at 6 and 12 d of shelf life following 5 months of cold storage was lower in ozone-treated fruits as compared with that of control fruits.

**Discussion**

**Ozone delayed ripening and blocked ethylene production in kiwifruit**

Fruit ripening is a unique cellular process and provides an important contribution to human nutrition (Martínez-Romero et al., 2007). In the present study, kiwifruit ripening was monitored during shelf life at 20 °C, following cold storage (0 °C, 95% relative humidity) in the absence (control) or presence of ozone (0.3 l l⁻¹). Ascorbic acid (A–C) and total phenolic contents (D–F) of kiwifruits (cv. ‘Hayward’) during shelf life (20 °C) for 1, 6, and 12 days after 1 (A,D), 3 (B,E), and 5 (C,F) months of cold storage (0 °C, 95% relative humidity) in the absence (control) or presence of ozone (0.3 l l⁻¹). Values are means ± SE (n = 6). Bars for each shelf-life period with different letters are statistically significant (Duncan’s multiple range test, P < 0.05).
temperatures (20 °C), which is consistent with the characteristic softening behaviour of this fruit (Koukounaras and Sfakiotakis, 2007). However, it should be noted that ozone-treated kiwifruits displayed higher firmness retention, especially after 5 months of cold storage (Fig. 1A–C), than control fruits. This observation along with previous data (Tzortzakis et al., 2007a; Rodoni et al., 2010) suggests that ozone is efficient in promoting higher firmness retention. In addition, ozone-treated kiwifruits displayed lower soluble solids contents than control fruits. (Fig. 1D–F). Collectively, these data suggest that ozone is effective at suppressing ripening and senescence process, and therefore can extend kiwifruit post-harvest life.

Kiwifruit has a typical climacteric behaviour with ethylene playing a major role in the ripening process (Antunes and Sfakiotakis, 2002). Control fruits showed a climacteric behaviour by starting autocatalytic production of ethylene when placed at 20 °C, though the exact starting time may depend on the duration of cold storage (Fig. 2A–C). Results regarding the impact of ozone on ethylene production during fruit ripening are contradictory and limited (Skog and Chu, 2001; Palou et al., 2002; Tzortzakis et al., 2007a). The present study found that ozone delayed climacteric ethylene rise and totally blocked ethylene production in kiwifruits after intermediate (3 month) or extended (5 month) cold storage, respectively. Since the ethylene climacteric rise is tightly linked with kiwifruit softening (Antunes and Sfakiotakis, 2002), the present data revealed a key post-harvest function of ozone that could be the main aetiological reason for the markedly retarded ripening process (Fig. 1). This regulatory ozone-driven mechanism may provide an excellent model for further studies addressing the role of ethylene signal transduction pathway and ripening phenomena in climacteric fruits. The present data also demonstrated that kiwifruit are characterized by a time lag between the increase in respiration rate and the induction of ethylene production, confirming previous observations (Sfakiotakis et al., 1997). In addition, respiration rates are enhanced by ozone treatment after short (Fig. 2D) or intermediate (Fig. 2E) cold storage and subsequent transfer to 20 °C. This fact, which was also observed by Forney et al. (2007) in cold-stored vegetables, may indicate that kiwifruits experience physiological injury as a result of the oxidative stress caused by ozone exposure. However, the respiration rate is reduced in ozone-treated kiwifruits following extended cold storage compared with control fruits (Fig. 2F). This specific reduction in respiration may be related to the absence of climacteric peak of ethylene production in fruits exposed to ozone for 5 months (Fig. 2C), as previously suggested (Antunes and Sfakiotakis, 2002).
Ozone acts as an antioxidant and anti-radical elicitor in kiwifruits during ripening

As kiwifruit is a good source of phytochemicals (Tavarini et al., 2008), the present study evaluated the antioxidant potency of kiwifruit extracts during shelf-life period. An induction of antioxidant activity, as measured by both FRAP and DPPH assays, was observed in ozone-treated kiwifruits, particularly after 1 d of shelf life (Fig. 3). These data indicated that the exposure of kiwifruits to ozone stimulated their antioxidant activity, presumably because of its action as a signal molecule for the activation of antioxidant-stress responses in kiwifruit, which possibly experienced an oxidative stress during the cold storage period. Although a variety of negative and positive interactions have been reported between ozone with either ascorbic acid (Allende et al., 2007; Tavarini et al., 2008) or phenolic compounds (Allende et al., 2007; Artès-Hernández et al., 2007; Tzortzakis et al., 2007b) in fruits undergoing post-harvest ripening, the present study further showed that ozone application may increase ascorbic acid and phenol content in kiwifruits, especially after 1 d of shelf life (Fig. 4).

Because of the differential patterns of the antioxidant-related parameters between ozone-treated and untreated kiwifruits ripened for 1 d at 20 °C after 1, 3, or 5 months cold storage (Figs. 3 and 4), this study investigated the relative phenolic-dependent anti-radical activity of kiwifruit extracts with respect to their protective abilities to counteract pBR322 DNA strand scission against HO, ONOO⁻, or ROO⁻. Phenolics extracted from untreated fruits were quite effective at protecting radical-derived DNA scission (Fig. 5A,C), thus confirming earlier observations that kiwifruit extracts can protect DNA from oxidation (Collins et al., 2001). Both in-gel assay (Fig. 5A,C) and quantitative analysis (Fig. 5B,D) revealed that the phenolic extracts of ozone-treated kiwifruits were very potent in inhibiting DNA breakage induced by HO or ONOO⁻, thereby lending evidence that ozone acts as a potent anti-radical elicitor against commonly encountered oxidant and nitrosant damaging agents.
Characterization of kiwifruit proteome during ripening

Despite increasing interest in plant proteomics (Agrawal et al., 2011; Job et al., 2011), no systematic analysis of the kiwifruit proteome has been published. To fill this gap, 1D-PAGE coupled with mass spectrometry analysis was applied to characterize the reference proteomic map in kiwifruit during ripening at 20°C. This allowed the characterization of 102 kiwifruit proteins (Table 1; see also Supplementary Table S2), which is a substantial increase because on 25 August 2011 only 17 reviewed kiwi proteins were listed in the Uniprot/Swiss-prot database, of which only three were from kiwifruit (http://www.uniprot.org/uniprot/?query=kiwi&sort=score). The largest functional group of identified kiwifruit proteins is related to energy (Fig. 6B). This is a diverse group of proteins that includes proteins playing a role mainly in glycolysis and the tricarboxylic acid cycle. These proteins are expected to provide the necessary energy metabolites for the ongoing ripening process, as previously suggested in peach fruit (Qin et al., 2009b). The present analysis also showed the presence of various forms of malate dehydrogenase (Fig. 6A, protein zones 6, 14–17), enolase (Fig. 6A, protein zones 8–13), isocitrate dehydrogenase (Fig. 6A, protein zone 11), transaldolase (Fig. 6A, protein zone 15), glyceraldehyde-3-phosphate dehydrogenase (Fig. 6A, protein zones 15 and 17), fructose-bisphosphate aldolase (Fig. 6A, protein zones 15–20), NADPH quinone oxidoreductase (Fig. 6A, protein zone 18), and oxidoreductase NAD-binding (Fig. 6A, protein zones 22 and 23), suggesting that during ripening, specific energy-related isoforms of kiwifruit proteins are presented. Another enzyme that

### Table 1. Identified kiwifruit proteins

Proteins are ontologically classified into functional categories (01–30) proposed by Bevan et al. (1998). Detailed information of protein identification is provided in Supplementary Table S2.

<table>
<thead>
<tr>
<th>Functional category</th>
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<tr>
<td>01 Metabolism (11)</td>
<td>Ferredoxin-dependent glutamate synthase</td>
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<tr>
<td>01.01</td>
<td>β-Galactosidase</td>
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<tr>
<td>01.01</td>
<td>Alanine aminotransferase</td>
</tr>
<tr>
<td>01.01</td>
<td>Glutamate dehydrogenase</td>
</tr>
<tr>
<td>01.05</td>
<td>α-Galactosidase</td>
</tr>
<tr>
<td>01.06</td>
<td>Acetyl-Coenzyme A oxidase (2)</td>
</tr>
<tr>
<td>01.01</td>
<td>Methylcrotonoyl-CoA carboxylase</td>
</tr>
<tr>
<td>01.05</td>
<td>UDP-glucuronate decarboxylase</td>
</tr>
<tr>
<td>01.05</td>
<td>UDP-glucose-1-phosphate uridylyltransferase</td>
</tr>
<tr>
<td>01.03</td>
<td>Nucleoside diphosphate kinase</td>
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<tr>
<td>02.10</td>
<td>Enolase (6)</td>
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<tr>
<td>02.10</td>
<td>Isocitrate dehydrogenase</td>
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<tr>
<td>02.07</td>
<td>Transaldolase (2)</td>
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<tr>
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<td>Malate dehydrogenase (2)</td>
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<td>02.01</td>
<td>Glyceraldehyde-3-phosphate dehydrogenase</td>
</tr>
<tr>
<td>02.01</td>
<td>Fructose-bisphosphate aldolase (5)</td>
</tr>
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<td>Glyceroldehyde-3-phosphate dehydrogenase</td>
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<td>02.01</td>
<td>Triosephosphatase isomerase (2)</td>
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<td>03 Cell Growth (3)</td>
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<td>03.26</td>
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<td>05.01</td>
<td>Eukaryotic translation initiation factor</td>
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<tr>
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<td>Endo-β-1,4-glucanase</td>
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<td>Pectinmethylesterase inhibitor (2)</td>
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<td>10.04.10</td>
<td>Small GTP-binding protein</td>
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<tr>
<td>11 Disease/defence (13)</td>
<td>Glutathione reductase</td>
</tr>
<tr>
<td>11.06</td>
<td>Polygalacturonase inhibitor</td>
</tr>
<tr>
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<td>Peroxidase</td>
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<tr>
<td>11.06</td>
<td>Lactoylglutathione lyase (3)</td>
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<td>Kiwelin</td>
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<td>11.02</td>
<td>2-Cys peroxiredoxin</td>
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<td>11.02</td>
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</tr>
<tr>
<td>11.02</td>
<td>Bet v 1 related allergen</td>
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<tr>
<td>11.05</td>
<td>Glycine-rich RNA binding protein</td>
</tr>
<tr>
<td>12 Unclear classification (1)</td>
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</tr>
<tr>
<td>30-Unknown (9)</td>
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</tr>
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<td>30</td>
<td>Hypothetical protein (6)</td>
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may be linked to the energy group is the mitochondrial ATP synthase (F1) subunit \( \beta \), which was also identified (Fig. 6A, protein zones 8 and 9), possibly due to the increased demand for ATP during starch–sugar conversion in ripening kiwifruits. An interesting detection was the identification of maturase (Fig. 6A, protein zone 7). Chloroplast maturases evolved into general group II intron splicing factors, potentially mirroring a key step in the evolution of spliceosomal introns. The presence of the maturase in the kiwifruit proteome is indicative of ‘generation of precursor metabolites and energy’ as well as ‘metabolic compound salvage’ linking the splicing organelar introns to global signals that regulate gene expression in response to energy state (Mohr and Lambowitz, 2003), thus suggesting a role for this maturase as an effector protein during kiwifruit ripening.

The second major functional group consisted of proteins involved in protein metabolism and modification (Fig. 6B), including proteins with chaperone function such as HSP70 (Fig. 6A, protein zone 4). The accumulation of heat-shock proteins (HSPs) seems to be a common ripening regulatory mechanism in fruits (Giribaldi et al., 2007; Bianco et al., 2009; Muccilli et al., 2009; Manganaris et al., 2011) as these proteins play pivotal role in the degradation of damaged or misfolded peptides as occurs during fruit ripening (Faurobert et al., 2007). Also, the identification of 60S ribosomal protein (Fig. 6A, protein zones 21 and 31) and eukaryotic translation initiation factor 5A (Fig. 6A, protein zone 28) in kiwifruits during ripening is in accordance with the characterized role of these proteins in the regulation of pineapple fruit senescence (Moyle et al., 2005). Additionally, many proteins were identified, such as actinidin (Fig. 6A, protein zone 9), cysteine protease (Fig. 6A, protein zone 14), actinidin chain A (Fig. 6A, protein zone 21), chymopapain (Fig. 6A, protein zone 21), leucine aminopeptidase (Fig. 6A, protein zone 28), and cystatin (Fig. 6A, protein zone 31), which are responsible for proteolysis and nitrogen recycling and could be involved in the degradation of damaged or misfolded peptides during ripening (Faurobert et al., 2007). Finally, the present study identified the biotin-dependent methylcrotonoyl-CoA carboxylase, an enzyme involved in leucine catabolism in all living cells and which in plants is strongly induced during leaf senescence (Alban et al., 2000). Therefore, this enzyme might be involved in maintaining leucine homeostasis during kiwifruit ripening.

A third major functional group consisted of proteins playing a role in plant defence. Considering that both low-temperature storage and fruit ripening are associated with oxidative processes (Pedreschi et al., 2008), the presence of reactive oxygen species-scavenging proteins in kiwifruits, such as glutathione reductase (Fig. 6A, protein zone 10) and peroxidase (Fig. 6A, protein zone 15), could be required for satisfying the cellular oxidative conditions favourable for ripening. In addition, 2-Cys peroxiredoxin (Fig. 6A, protein zone 25) and small GTP-binding protein (Fig. 6A, protein zone 26) were also identified. 2-Cys peroxiredoxins constitute an ubiquitous group of peroxidases that exhibit dual physiological functions both as antioxidant and molecular chaperone in response to oxidative stimuli (Aran et al., 2009), whereas small GTP-binding protein acts as a molecular switch in oxidative transduction cascades (Shichrur and Yalovsky, 2006). The detection of such proteins in the present study supports their role in oxidative signalling during kiwifruit ripening. Further, the identification of nuclear glycine-rich RNA binding protein (Fig. 6A, protein zone 31) in kiwifruit is in agreement with the observation that this protein performs a RNA chaperone function.
during cold adaptation (Kim et al., 2010). Since methylglyoxal (a metabolite deriving from glycolysis) is highly cytotoxic, it is also interesting to note the present identification of various forms of lactoylglutathione lyase in kiwifruit (Fig. 6A, protein zones 17–19). This enzyme, also called glyoxalase I, catalyses the first step of the glyoxal pathway, in which methylglyoxal and glutathione are converted into S-lactoylglutathione, which is further converted into lactic acid (Jia et al., 2006). Thus the glyoxalase activity along with the identification of triosephosphate isomerase (Fig. 6A, protein zone 23), which is directly involved in the generation of methylglyoxal, likely reflects a mechanism to control the level of toxic methylglyoxal in kiwifruit, as previously proposed in pear fruit stored under long-term controlled-atmosphere conditions (Pedreschi et al., 2007).

Proteins belonging to the category of cell wall-modifying and structural proteins, such as pectinesterase (Fig. 6A, protein zones 9 and 10), endo-β-1,4-glucanase (Fig. 6A, protein zone 10), xyloglucan endotransglycosylase (Fig. 6A, protein zone 19), α-expansin (Fig. 6A, protein zones 24 and 25), and pectinmethylesterase inhibitor (Fig. 6A, protein zones 27 and 28), were also identified. The identification of cell wall-loosening proteins supports the pivotal role of cell-wall extensibility required for the softening process during kiwifruit ripening (Fig. 1C). Particularly relevant is also the identification of proteins related to metabolism of sugars/poly saccharides, such as β-galactosidase (Fig. 6A, protein zone 4) and α-galactosidase (Fig. 6A, protein zone 14). Both enzymes are able to release α- or β-linked D-galactose from polymers, oligosaccharides, or secondary metabolites and are responsible for most of cell-wall degradation and softening in kiwifruit (Ross et al., 1993). In addition, kiwifruit proteins involved in the regulation of growth, such as β-1,3-glucanase (Fig. 6A, protein zone 19) and agglutinin (Fig. 6A, protein zones 8 and 9), may be also involved in the softening process as previously suggested (Bogoeva et al., 2004; Deytieux et al., 2007).

It is noted that several of the presently identified kiwifruit proteins, such as β-1,3-glucanase, malate dehydrogenase, eukaryotic translation initiation factor 5A, enolase, expansin, glyceraldehyde-3-phosphate, glycine-rich RNA binding protein, small GTP-binding protein, HSPs, isocitrate dehydrogenase, malate dehydrogenase, pectinesterase, and triosephosphate isomerase (Table 1; see also Supplementary Table S2), have also been identified in a number of fruit species during their ripening (Rocco et al., 2004; Deytieux et al., 2009, 2011; Zhang et al., 2011), thus highlighting their general importance for the ripening process (Fig. 1C). Particularly relevant is also the identification of triosephosphate isomerase (Fig. 6A, protein zones 17–19). This enzyme, also called glyoxalase I, catalyses the first step of the glyoxal pathway, in which methylglyoxal and glutathione are converted into S-lactoylglutathione, which is further converted into lactic acid (Jia et al., 2006). Thus the glyoxalase activity along with the identification of triosephosphate isomerase (Fig. 6A, protein zone 23), which is directly involved in the generation of methylglyoxal, likely reflects a mechanism to control the level of toxic methylglyoxal in kiwifruit, as previously proposed in pear fruit stored under long-term controlled-atmosphere conditions (Pedreschi et al., 2007).

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Protein carbonylation is increased during kiwifruit ripening

The remarkable antioxidant and anti-radical responses observed in ozone-treated fruits (Figs. 3–5) prompted this investigation of whether ozone treatment can modify the protein carbonylation status in kiwifruit during ripening. Quantitative evaluation of the immunoblot results (Supplementary Fig. S3) together with the measurement of protein carbonyl group content (Fig. 7B) showed that protein carbonylation was progressively induced during ripening, thus confirming that protein oxidation is linked to fruit senescence (Qin et al., 2009a,b). The observed lower levels of protein carbonyls in ozone-treated fruits during ripening compared with control fruits (Fig. 7B) may be a consequence of the inhibition of ethylene biosynthesis (Fig. 2C) and the relatively high antioxidant and anti-radical activity in ozone-treated fruits (Figs. 3 and 4C,F and Fig. 5A–D, respectively) and may be associated with delayed senescence features in kiwifruit (Fig. 1). Since previous studies have indicated that protein oxidation is not necessarily a deleterious phenomenon (Job et al., 2005; Oracz et al., 2007) but has been involved in cellular signalling (Wong et al., 2010), the overall pattern of protein carbonylation in kiwifruit (Fig. 7) raises the hypothesis that this redox modification is an important signalling mechanism governing fruit activity during ripening.

In summary, the use of an ozone-enriched cold atmosphere markedly delayed kiwifruit ripening at 20 °C compared with conventional storage. The present data also underscores that the delay/block of climacteric rise of ethylene production is a key mechanism targeted by ozone
in kiwifruit upon ripening. Furthermore, this study provides evidence that ozone application during cold storage acts as an antioxidant and anti-radical elicitor in kiwifruit. This report is the first draft of the Actinidia proteome during ripening using 1D-SDS-PAGE and mass spectrometry techniques. A number of proteins that are related to energy, protein metabolism, defence, cell structure, and amino acid or sugar/poly saccharide metabolism are identified. In addition, this investigation shows that ripening and ozone specifically regulate kiwifruit protein carbonylation. These results provide insights into kiwifruit metabolism during post-harvest life and also a starting point to develop ozone-based treatments strategies for controlling fruit ripening and quality in practice.

Supplementary information

Supplementary data are available at JXB online.

Supplementary Table S1. Quantification of protein zones from 1D-gels of kiwifruit proteins.

Supplementary Table S2. Detailed information of the identified kiwifruit proteins.

Supplementary Fig. S1. Scheme summarizing the experimental set up.

Supplementary Fig. S2. Weight loss of kiwifruits (cv. Hayward) after cold storage in the absence or presence of ozone for 1, 3, or 5 months plus 1 day of shelf life.

Supplementary Fig. S3. Relative abundance (arbitrary units) of carbonyl-reactive signal of the immunoblot results in Fig. 7A.

Supporting information

Active Link 1. Mascot results for xyloglucan endotransglycosylase (protein zone 19) identifications based on one peptide sequences: http://139.124.36.211/mascot/cgi/peptide_view.pl?file=/data/20100517/F064815.dat&query=235&hit=1&index=gi%7c950299&px=1&section=5&ave_thresh=45.

Active Link 2. Mascot results for α-expansin (protein zone 25) identifications based on one peptide sequences: http://139.124.36.211/mascot/cgi/peptide_view.pl?file=/data/20100517/F064843.dat&query=31&hit=1&index=gi%7c1815681&px=1&section=5&ave_thresh=45.

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References


