Ozone-induced inhibition of kiwifruit ripening is amplified by 1-methylcyclopropene and reversed by exogenous ethylene

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Abstract

Background: Understanding the mechanisms involved in climacteric fruit ripening is key to improve fruit harvest quality and postharvest performance. Kiwifruit (Actinidia deliciosa cv. 'Hayward') ripening involves a series of metabolic changes regulated by ethylene. Although 1-methylcyclopropene (1-MCP, inhibitor of ethylene action) or ozone (O3) exposure suppresses ethylene-related kiwifruit ripening, how these molecules interact during ripening is unknown.

Results: Harvested 'Hayward' kiwifruits were treated with 1-MCP and exposed to ethylene-free cold storage (0 °C, RH 95%) with ambient atmosphere (control) or atmosphere enriched with O3 (0.3 μL L−1) for up to 6 months. Their subsequent ripening performance at 20 °C (90% RH) was characterized. Treatment with either 1-MCP or O3 inhibited endogenous ethylene biosynthesis and delayed fruit ripening at 20 °C. 1-MCP and O3 in combination severely inhibited kiwifruit ripening, significantly extending fruit storage potential. To characterize ethylene sensitivity of kiwifruit following 1-MCP and O3 treatments, fruit were exposed to exogenous ethylene (100 μL L−1, 24 h) upon transfer to 20 °C following 4 and 6 months of cold storage. Exogenous ethylene treatment restored ethylene biosynthesis in fruit previously exposed in an O3-enriched atmosphere. Comparative proteomics analysis showed separate kiwifruit ripening responses, unraveled common 1-MCP- and O3-dependent metabolic pathways and identified specific proteins associated with these different ripening behaviors. Protein components that were differentially expressed following exogenous ethylene exposure after 1-MCP or O3 treatment were identified and their protein-protein interaction networks were determined. The expression of several kiwifruit ripening related genes, such as 1-aminocyclopropane-1-carboxylic acid oxidase (ACO1), ethylene receptor (ETR1), lipoxygenase (LOX1), geranylgeranyl diphosphate synthase (GGP1), and expansin (EXP2), was strongly affected by O3, 1-MCP, their combination, and exogenously applied ethylene.

Conclusions: Our findings suggest that the combination of 1-MCP and O3 functions as a robust repressive modulator of kiwifruit ripening and provide new insight into the metabolic events underlying ethylene-induced and ethylene-independent ripening outcomes.

Keywords: Actinidia deliciosa, Climacteric, Cold storage, Ethylene biosynthesis, Gene expression, Kiwifruit ripening, 1-Methylcyclopropene, Ozone, Postharvest, Proteomics, Softening, Protein-protein interaction

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Background

Fleshy fruits undergo a complex developmental program that ends in the irreversible process of ripening and eventual tissue senescence [1]. Over-ripening and ethylene-induced senescence shorten fruit postharvest storage potential and create huge economic losses [2]. Thus, understanding the regulation of fruit ripening is of considerable agronomic value. Kiwifruit (Actinidia delicosa, cv. ‘Hayward’) is classified as climacteric in which ethylene synthesis, perception and signal transduction play key roles in ripening [3, 4]. In climacteric fruits, pre-climacteric exposure to 1-methylecyclopropene (1-MCP), through its high affinity for binding to ethylene receptors, can inhibit the perception of ethylene in fruit tissues [5], delay ethylene-dependent ripening and senescence and prolong fruit storage life [6, 7]. Many molecular and genetic mechanisms underlying the action of 1-MCP in fruit ripening have been identified. In kiwifruit, 1-MCP application inhibits expression of specific ethylene receptors (Ad-ERS1a, Ad-ETR2 and Ad-ETR3) and several transcription factors (Ad-ERF4, Ad-ERF6, Ad-ERF10 and Ad-ERF14) [8, 9]. In addition, 1-MCP inhibits kiwifruit softening by reducing cell wall related gene expression, such as polygalacturonase (PG) and expansin (EXP) [10–12]. Together, these data on 1-MCP function indicate that ethylene signal transduction is essential not only to initiate climacteric kiwifruit ripening, but also to complete ripening and senescence.

Ozone (O3) can reduce spoilage of fresh fruits and vegetables and delay fruit ripening by directly oxidizing ethylene [13–16]. O3 exerts major residual effects in kiwifruit ripening physiology and both ethylene biosynthesis and cell wall turnover are specifically targeted by O3 after long-term exposure under cold storage conditions. Two or more months of storage in an O3-enriched atmosphere blocked kiwifruit ethylene biosynthesis during ripening at 20°C by inhibiting ACS and ACO gene expression and enzymatic activity [17, 18]. In agreement with its function as a softening repressor, O3 modulated kiwifruit cell wall is remodeled by depressing cell wall swelling, pectin and neutral sugar solubilization and by inhibiting the activity of cell wall-degrading enzymes [18]. Upstream of the ethylene pathway, several transcripts, such as bet v 1 related allergen, 3 hydroxy-3-methylglutaryl CoA reductase and geranylgeranyl diphosphate synthase are regulated by O3 [19]. The reported proteomic analysis determined that down-regulation of protein expression is one of the components related to kiwifruit ripening inhibition by O3, as O3 causes widespread down-regulation of protein abundance of ATP-citrate lyase, kiwellin and pectin acetyltransferase precursor [17, 19]. Further research is necessary to characterize kiwifruit ripening and the specific roles of O3.

This study thoroughly investigates 1-MCP and O3 signaling in kiwifruit ripening physiology through a systematic analysis at the physiological and molecular level. Physiological data collected during ripening when fruits were exposed to either ambient air or exogenous ethylene are combined with proteomic and transcriptional approaches.

Results

Physiological characterization of the effect of 1-MCP, O3 and exogenous ethylene on kiwifruit ripening

Endogenous ethylene emission from untreated kiwifruit was initiated after 10, 6 and 4 days (d) ripening at 20°C, following 2, 4 and 6 months of cold storage, respectively. Treatment with 1-MCP immediately after harvest, exposure to O3 during cold storage and the combination of both treatments (1-MCP + O3) significantly inhibited endogenous ethylene production of kiwifruit during ripening at all three ripening periods (following 2, 4 or 6 months of cold storage) (Fig. 1a-c). To clarify whether the inhibition of endogenous ethylene production observed in kiwifruit exposed to O3 and 1-MCP could be recovered by a short treatment with exogenous ethylene, an intermittent experiment was set up. O3-stored fruit in the absence of 1-MCP produced significant concentrations of ethylene (12.2 or 27.6 μL kg h−1) after 14 or 8 d ripening in response to exogenous ethylene exposure following 4 or 6 months of cold storage, respectively (Fig. 2a, c). 1-MCP-treated fruit stored without O3 exhibited onset of ethylene production (1.7 or 1.1 μL kg h−1) in response to exogenous ethylene after 14 or 8 d ripening at 20°C following 4 or 6 months of cold storage, respectively (Fig. 2a, c). In contrast, 1-MCP-treated fruit stored in O3 atmosphere for up to 4 or 6 months exhibited no detectable ethylene production during ripening following exogenous ethylene treatment (Fig. 2a, c).

Respiration rate (RR) in control fruit immediately upon removal from cold storage following 2, 4 or 6 months was reduced or remained constant after 4 or 2 d ripening, respectively; then RR exhibited a respiratory climacteric increase (Fig. 1d-f). In contrast, the postharvest treatments, with the exception of 1-MCP-treated fruit after 6 months cold storage, reduced RR during ripening following 2, 4 and 6 months of storage. Although RR was unaffected by exogenous ethylene in control fruit during ripening after 6 months of cold storage, it was increased by exogenous ethylene in fruit exposed to O3 or 1-MCP individual treatments after 4 or 6 months of cold storage. In contrast, the 1-MCP plus O3-treated fruit had the lowest RR after exogenous ethylene treatment (Fig. 2b, d).

Analysis of pericarp and core tissue firmness showed that 1-MCP + O3 was the most effective treatment in inhibiting kiwifruit softening during storage (Fig. 1g-l). Interestingly, 1-MCP + O3-treated fruit had high pericarp (28 N) and core (98 N) tissue firmness, that is significantly greater than the acceptable for consumption levels (10–15 N), after 2 months of cold storage plus 14 d ripening. Individual 1-MCP and O3 treatments...
lowered the softening rates below those of control fruit; however, they did not retain firmness comparable to 1-MCP + O₃. 1-MCP alone retained higher levels of core tissue firmness than O₃-enriched atmosphere (Fig. 1j-l), but there were no differences in pericarp firmness between these two treatments. Exogenous ethylene after 8 d ripening following 4 months of storage increased softening rates in all treatments, although there were sharp changes in core tissue firmness in kiwifruit exposed to both 1-MCP treatments. Following 6 months of cold storage, exogenous ethylene reduced pericarp and core firmness in all treatments while in control fruit, no further firmness reduction was observed (Fig. 2e-h). Kiwifruit from control, control-ETH and O₃-ETH treatments, which produced high ethylene rates following 6 months of cold storage (Fig. 2a, c), exhibited internal breakdown symptoms due to over-ripening after 8 d (Fig. 2i).

Fig. 1 Kiwifruit ripening was inhibited by 1-MCP and O₃. Following harvest, kiwifruit (cv. 'Hayward') were treated with or without (control) 1-MCP (0.6 μL L⁻¹, 0 °C, 24 h) and then cold-stored (0 °C, RH 90%) in two separated cold rooms in which ethylene was controlled by catalytic ethylene oxidation and the atmosphere was ambient (control) or enriched with O₃ (0.3 μL L⁻¹). Fruits were removed from cold storage after 2, 4 and 6 months and then transferred to 20 °C (90% RH), where kiwifruit ripening was characterized for up to 14 d. Changes in ethylene emission rate (a, e, i), respiration rate (b, f, j), firmness of outer pericarp (c, g, k) and core tissue (d, h, l) in kiwifruit during ripening at 20 °C were measured. Phenotypes of kiwifruit after 8 d ripening at 20 °C following 6 months cold storage (m). Values represent the mean of three replicates of 10 fruits each that were analyzed at each ripening time point. Vertical bars in figure plates represent the least significant difference (LSD, P = 0.05), which was used for comparisons of means between treatments and ripening time points.
Kiwifruit dry matter content (DMC) remained constant postharvest, irrespective of the experimental conditions (Additional file 1: Figure S1). In addition, soluble solids concentration (SSC) increase was delayed by 1-MCP or O₃ application while 1-MCP + O₃ severely delayed SSC accumulation during ripening following 2 months of storage (Additional file 2: Figure S2). Exogenous ethylene, particularly following 4 months of cold storage plus 8 d ripening, increased SSC regardless of treatment (Additional file 3: Figure S3). Generally, no differences were found in titratable acidity (TA) among treatments (Additional file 2: Figure S2), although exogenous ethylene decreased TA in O₃-treated fruit following 6 months of cold storage plus 8 d ripening (Additional file 3: Figure S3).

Ethylene biosynthesis during kiwifruit ripening is affected by 1-MCP and O₃
To further describe the effect of 1-MCP and O₃ on ethylene biosynthesis, we profiled ethylene biosynthesis intermediates and enzymes (Fig. 3) during ripening at specific time points based on the ripening behavior of control fruit (Fig. 1). These time points included the beginning of ripening immediately upon removal from 2 months of cold storage (0 d at 20 °C), pre-ethylene production (4 d at 20 °C), initiation of ethylene production (8 d at 20 °C), increase of ethylene production (10 d at 20 °C), and peak of ethylene production (14 d at 20 °C). In addition, five similar time points based on the ripening behavior of control fruit (0, 2, 4, 6 and 8 d at 20 °C) were selected following 6 months of cold storage. All postharvest treatments inhibited ACC and MACC concentrations and ACS and ACO enzyme activities during ripening, following 2 and 6 months of cold storage (Fig. 3a-h). Exogenous ethylene after 6 months of cold storage increased ACC concentrations in O₃-stored fruit at 6 d ripening but did not affect ACC concentrations in control or 1-MCP-treated fruit, compared to their untreated with ethylene counterparts (Fig. 3b). The concentration of ACC in O₃-treated fruit increased further after 8 d ripening due to exogenous ethylene (Fig. 3b). Ethylene biosynthesis analysis also showed strong

Fig. 2 Exposure to exogenous ethylene reversed the O₃-mediated ripening inhibition in kiwifruit. Following 4 or 6 months of cold storage plus 8 or 1 d maintenance at 20 °C (indicated with grey arrows), respectively, kiwifruit exposed to 1-MCP and/or O₃ (see Fig. 1) were further treated with exogenous ethylene (100 μL L⁻¹, 20 °C, 90% RH, 24 h) and their ripening was characterized for up to 14 d. Changes during kiwifruit ripening at 20 °C in ethylene emission rate (a, c), respiration rate (b, d), firmness of outer pericarp (e, g) and core tissue (f, h). Kiwifruit phenotypes at 8 d of ripening at 20 °C following 6 months of cold storage (i). Values represent the mean of three replicates of 10 fruits each that were analyzed at each ripening time point. Markers and lines in grey represent kiwifruit untreated with exogenous ethylene as in Fig. 1. Vertical bars in figure plates represent the least significant difference (LSD, №.05), which was used for comparisons of means between treatments (exposed or not to exogenous ethylene) and ripening time points.
inhibition of ACS and ACO activities and ACC accumulation by 1-MCP and O₃ treatments (Fig. 3a, e, g). Exogenous ethylene following 6 months of cold storage rapidly increased ACS and ACO activities in O₃-treated (O₃-ETH) kiwifruit after 8 d ripening (Fig. 3c, g). Exogenous ethylene did not affect ACS and ACO enzyme activities either in control or in 1-MCP-treated fruit with or without O₃ compared to their untreated with ethylene counterparts (Fig. 3f, h).

**Kiwifruit protein changes and functional classification**

To characterize the current biological system, large scale analysis of kiwifruit proteome was performed in the outer pericarp tissue (green flesh) of kiwifruit stored for 6 months and subsequently ripened 8 d at 20 °C (Figs. 1 and 2). Following comparative 2DE-analysis (Fig. 4a, b) and nanoLC/MS/MS, we identified 127 proteins that differentially accumulated among treatments, of which 48 were up-accumulated and 79 were down-accumulated (Fig. 4b). Identified proteins were predominantly related to disease/defence (39.2%), followed by energy (20.8%), protein destination/storage (7.5%), cell structure/cell wall (5.8%) and signal transduction (5%). A complete list of the protein identification, including peptide sequences, accession number, subcellular localization and matching criteria is presented in Additional file 4: Table S1. Seventeen proteins were identified in more than one spot, indicating that many of the differentially expressed spots were either subjected to post-translational modification or were members of multi-genic protein families. Among these multi-spot identified proteins were β-D-galactosidase (Fig. 4c, spots: 7908, 7909, 7907, 7619, 7910, 8911), chaperonin CPN60 (Fig. 4c, spots: 2801, 2802), fructose-bisphosphate aldolase (Fig. 4d, spots: 6607, 6506, 7510, 6505, 7509, 7608, 6606, 7507), glycer-aldehyde 3-phosphate dehydrogenase (GAPDHc, Fig. 4d, spots: 7511, 7513, 7613, 7512, 8606, 3505), lactoylglutathion iyase (Fig. 4c, spots: 2501, 2601), malate dehydrogenase (Fig. 4c, spot: 2314; Fig. 4d, spot: 3501; Fig. 4d, spot: 7609), polyphenoloxidase (Fig. 4d, spots: 3505, 3601), natterin (Fig. 4d, spot: 3508; Fig. 4d, spot: 7709; Fig. 4d, spot: 5510), and remorin (Fig. 4c, spot: 8803; Fig. 4d, spot: 7607).

The proteins that exhibited either increased or decreased abundance relative to their respective controls were separated into three groups using a Venn diagram (Fig. 5). The first group is represented by 13 proteins (5 up-accumulated and 8 down-accumulated) that had altered expression with 1-MCP; the second, by 30 proteins (10 up-accumulated and 20 down-accumulated) modulated by O₃; and the third, by 31 proteins (20 up-accumulated and 11 down-accumulated) regulated by the combination of 1-MCP and O₃ (Figs. 5 and 7, Additional file 4: Table S1). The functional classes of kiwi-fruit proteins that were similarly or differentially accumulated in control and 1-MCP/O₃-treated fruit are also presented (Fig. 5). For example, the 31 proteins modulated by 1-MCP + O₃ were related to disease/defence, cell structure/cell wall and energy. A proportion of all identified proteins were common to two or more group types. The overlap among all treatments (1-MCP, O₃, and 1-MCP + O₃) included 6 proteins, while the overlap between 1-MCP and O₃ treatments included only 1 protein, polygalacturonase (PG) (Fig. 5, Additional file 4: Table S1). Other proteins were specifically induced or repressed by particular treatments. Four proteins were affected only by 1-MCP, while 16 proteins...
were exclusively modulated under O₃ or 1-MCP + O₃ conditions, suggesting changes in biological activities specific to O₃ or to the combination of 1-MCP and O₃. The major portion of 1-MCP/O₃-specific proteins included disease/defence proteins (Fig. 5).

To investigate ethylene-related protein changes, the fruit proteome with each chemical treatment and exogenous ethylene was compared to control fruit treated with ethylene (Fig. 6, Additional file 4: Table S1). Among ethylene-affected proteins, 23 were different in 1-MCP-treated fruit, of which 6 proteins were up-accumulated and 17 were down-accumulated. In addition, 12 differentially accumulated proteins were identified in the O₃ treatment, 4 up-accumulated and 8 down-accumulated (Figs. 6 and 8, Additional file 4: Table S1). Fifteen proteins (13 up-accumulated and 2
down-accumulated) were modulated by ethylene in the combined 1-MCP + O$_3$ treatment (Figs. 6 and 8, Additional file 4: Table S1). Four kiwifruit ETH-responsive proteins were commonly detected in kiwifruit exposed to 1-MCP and O$_3$, while 1 protein overlapped between 1-MCP and 1-MCP + O$_3$. There was no overlap between O$_3$ and 1-MCP + O$_3$ treatments. Additionally, 3 proteins were ETH-affected across all treatments. These commonly regulated proteins were involved in disease/defence and signal transduction. A comparison of protein changes in each single treatment (control, 1-MCP, O$_3$, or 1-MCP + O$_3$) with their ethylene-treated counterparts is also presented (Additional file 5: Figure S4 and Additional file 6: Figure S5).

To validate the protein abundance results, transcript expression of PG was assayed by q-RT PCR analysis at various ripening times (Fig. 4e). PG gene transcript was strongly induced by ripening in control fruit, while O$_3$ significantly restricted this PG induction, confirmed by the abundance of spot 3702, identified as PG protein. Following exogenous ethylene application, PG expression was recovered completely in O$_3$-treated fruit (Fig. 4e), supporting the results attained by quantitative 2-DE and LC-MS/MS analysis (Figs. 7 and 8, Additional file 4: Table S1).

**Networks of ripening inhibition- and stimulation-responsive kiwifruit proteins**

The experimental structure, with three treatments that inhibited ethylene biosynthesis (1-MCP, O$_3$, and 1-MCP + O$_3$) and an intermittent test using exogenous ethylene exposure that stimulated ripening, provided an interesting biological system in which to study regulatory networks among the identified proteins. PPI networks created through STRING 9.0 [20] predicted functional links among identified kiwifruit proteins that were expressed in response to ripening inhibitors or stimulators. In kiwifruit treated with ripening inhibitors and not exposed to exogenous ethylene, the major clusters of interacting proteins involve proteins related to energy, protein destination/storage, and disease/defence (Fig. 9a). In kiwifruit treated with ripening inhibitors postharvest and exposed to exogenous ethylene post-storage, the major groups of interacting proteins are related to disease/defence, energy, transporters, protein destination/storage, signal transduction and secondary metabolism (Fig. 9b).

BiNGO 2.44 [21] software was used to predict statistically significant categories in over- or under-represented biological pathways and molecular functions of kiwifruit.
undergoing ripening inhibition and stimulation (Fig. 10). The enriched Gene Ontology (GO) lists of biological pathways (Additional file 7: Table S2) and molecular functions (Additional file 8: Table S3) of identified and annotated kiwifruit proteins are provided. The statistically significant over-represented biological pathways in ripening inhibited kiwifruit were the response to stress \( (p = 9.78 \times 10^{-6}) \) and the response to cadmium and metal ion \( (p = 2.53 \times 10^{-5} \text{ and } 6.34 \times 10^{-5}, \text{respectively}) \) (Fig. 10a, Additional file 7: Table S2a). In kiwifruit treated with ethylene in order to stimulate ripening, the major groups were the response to stress \( (1.44 \times 10^{-9}) \) and glucose catabolism \( (3.15 \times 10^{-8}) \), along with monosaccharide catabolism \( (3.41 \times 10^{-8}) \) and hexose catabolism \( (3.41 \times 10^{-8}) \) (Fig. 10c, Additional file 7: Table S2b). Molecular functions that were enriched in kiwifruit undergoing ripening inhibition were lyase activity \( (p = 3.83 \times 10^{-6}) \) and copper ion binding \( (p = 2.26 \times 10^{-5}) \) (Fig. 10b, Additional file 8: Table S3a). On the other hand, lyase activity \( (p = 3.32 \times 10^{-7}) \) was the most highly enriched molecular function in kiwifruit treated with exogenous ethylene (Fig. 10d, Additional file 8: Table S3b).

Ripening-related gene and transcription factor expression in kiwifruit exposed to 1-MCP, \( O_3 \), and exogenous ethylene
To examine the impact of 1-MCP, \( O_3 \), and exogenous ethylene in kiwifruit ripening, we profiled the expression of several ripening-related genes and transcription factors: aminocyclopropane-1-carboxylic acid oxidase 1 (ACO1), ethylene receptor (ETR), lipoxygenase 1 (LOX1), geranylgeranyl diphosphate synthase (GGPS), and expansin 2 (EXP2) (Fig. 11). Real-time RT-PCR analysis showed that \( ACO1 \) expression was constant in the control, but lower
in 1-MCP, O3 and 1-MCP + O3 treatments during ripening following 6 months of storage. Fruit exposed to O3 and 1-MCP + O3 retained less ACO1 expression than control or 1-MCP treatments at 8 d ripening (Fig. 11a, b, c). Exogenous ethylene induced ACO1 transcription in all treatments; however, there was more ACO1 expression in O3-ETH fruit than 1-MCP-ETH and 1-MCP + O3-ETH fruit at 8 d (Fig. 11d, e). Additionally, ETR expression increased in control fruit at 0 and 4 d and in 1-MCP-treated fruit at 8 d, but not in fruit exposed to O3 and 1-MCP + O3, that remained suppressed throughout the ripening period (Fig. 11f, g, h). Exogenous ethylene increased ETR transcription in control-ETH, 1-MCP-ETH and O3-ETH treatments compared to fruit maintained under ambient air (Fig. 11i, j). Meanwhile, EXP2 expression diminished gradually during ripening,
particularly in control and 1-MCP-treated fruits (Fig. 11k, l, m). Exogenous ethylene induced EXP2 expression in all treatments after 8 d ripening compared to fruit maintained under ambient air. Control-ETH followed by O₃-ETH and 1-MCP-ETH conditions exhibited the most EXP2 expression after 8 d ripening, with 1-MCP + O₃-ETH significantly lower (Fig. 11n, o). In all treatments except O₃, GGPS expression was induced at 4 d; however, GGPS declined to basal levels at 8 d (Fig. 11p, q, r). Exogenous ethylene provoked GGPS expression in all treatments at 8 d. Fruit exposed to O₃-ETH exhibited the most GGPS expression after 8 d ripening, followed
by control-ETH and 1-MCP-ETH, with 1-MCP + O₃-ETH significantly less (Fig. 11s, t). Gene expression of \textit{LOX1} in fruit treated with O₃ or 1-MCP + O₃ after 4 and 8 d ripening was less than control and 1-MCP (Fig. 11u, v, w). Exogenous ethylene stimulated \textit{LOX1} expression in all treatments after 8 d ripening. Control-ETH had the most \textit{LOX1} expression at 8 d, followed by O₃-ETH and 1-MCP-ETH, with 1-MCP + O₃-ETH having the least (Fig. 11x, y).

**Discussion**

This study was designed to describe and evaluate physiological and molecular changes caused by O₃ and 1-MCP in kiwifruit. The underlying rationale was to provide data relevant to a comprehensive understanding of regulatory mechanisms governing climacteric fruit ripening.

**Combined 1-MCP and O₃ treatment severely inhibits kiwifruit ripening**

Our results on kiwifruit ripening after 1-MCP and O₃ treatment (Fig. 1a-f) are consistent with previous work indicating that postharvest 1-MCP treatment and cold storage in an atmosphere enriched with O₃ can effectively inhibit ethylene emission rate and RR during kiwifruit ripening [10, 12, 17–19]. Although fruit exposed to 1-MCP or O₃ alone showed similar ripening behavior based on ethylene emission (Fig. 1a-c) and pericarp firmness (Fig. 1g-l), there were substantial differences in RR (Fig. 1d-f) and core tissue firmness (Fig. 1j-l). These differences indicate that 1-MCP or O₃ could regulate kiwifruit ripening through ethylene-dependent and independent pathways. This hypothesis was further supported by the combined 1-MCP + O₃ treatment, which suppressed kiwifruit softening more than the individual treatments alone (Fig. 1g-l). From a practical perspective, treatment with 1-MCP and subsequent cold storage in an O₃-enriched atmosphere could effectively inhibit kiwifruit ripening and softening and successfully extend their storage potential by 1.5 to 2 more months beyond individual 1-MCP or O₃ application. This is the first report of a positive interaction between 1-MCP and O₃ in fruit postharvest physiology and represents an interesting experimental model for fruit ripening syndromes. The severe ripening inhibition by 1-MCP + O₃ could lead to failure of the fruits ability to ripen and soften at eating-ripe firmness, particularly following short-term cold storage (2 months or less). It is recommended, therefore, that this combined 1-MCP + O₃ treatment should be applied only on fruit intended for long-term cold storage (more than 3 months) to avoid possible consumer rejection. Alternatively, 1-MCP + O₃-treated fruits should be exposed to exogenous ethylene if they have to be transported to the market earlier than the three-month storage window.

**Ripening inhibition is reversible by exogenous ethylene in O₃-treated, but not in 1-MCP + O₃-treated, kiwifruit**

The mechanism of 1-MCP action is well-characterized and involves tight binding to ethylene receptors in plant
tissues, blocking ethylene signaling [5]. Although O₃ inhibits ripening [17], the mechanism has not been clearly defined. The critical question raised by this and previous studies [17, 18] is whether the mode of action of O₃ in kiwifruit ripening stems from permanent oxidative damage or from potentially reversible biochemical inhibition. To directly address this question and given that exogenous ethylene directly induces kiwifruit ripening [4, 22], 1-MCP/O₃–treated kiwifruit that could not produce ethylene were exposed to exogenous ethylene (ETH) at specific ripening times. ETH treatment did not induce rapid endogenous ethylene production in fruit treated with 1-MCP, unlike in control-ETH-treated fruit (Fig. 2a, c). This suggests that 1-MCP blocks normal feedback regulation of ethylene production in kiwifruit, as in mature banana fruit [23]. On the other hand of particular interest is that O₃-treated fruit exposed to exogenous ethylene produced more endogenous ethylene (Fig. 2a, c), and can therefore sense and transduce ethylene signalling. The significant ethylene emission levels from O₃-ETH treated fruit suggests that O₃ may block kiwifruit ripening through biochemical inhibition of ACS and ACO activity, rather than through oxidative damage to some ethylene biosynthesis pathway components (Fig. 3).

**Specifically accumulated proteins by postharvest treatments provide insights into understanding kiwifruit ripening**

To further explore fruit ripening inhibition by chemical treatments (Figs. 1, 2 and 3), we used 2DE coupled with MS/MS to characterize 1-MCP and/or O₃ proteome activity. Thirteen or 30 proteins changed in response to 1-MCP or O₃, respectively, while 31 proteins were altered when these treatments were combined (Fig. 5, Additional file 4: Table S1). Nucleoside diphosphate kinase, a possible component of the ethylene signal transduction chain [24], was down-accumulated in response to postharvest 1-MCP treatment in papaya fruit [25]. The down-accumulation of a pathogenesis-related (PR) protein exclusively by 1-MCP suggests that PR may be associated with ethylene-related susceptibility to pathogen infection. A
similar trend in PR abundance was observed in unripe banana fruit exposed to 1-MCP [26]. The observed up-regulation of two kiwelin isoforms by 1-MCP, combined with several other kiwelin isoforms that were up-accumulated by the various 1-MCP/O3 treatments and the reversal of this up-regulation by exogenous ethylene, (Fig. 8, Additional file 4: Table S1) suggests that kiwelin has an important role in kiwifruit ripening. O3 probably acted as a protein repressor, since most proteins exclusively affected by O3 were down-accumulated (Fig. 7). Kiwifruit metabolism is unusual in that carbon is primarily stored as starch and eventually, through the ripening process, is hydrolyzed and converted to CO2 and sugars [27]. To accurately estimate a carbon budget in kiwifruit during ripening, it is necessary to consider CO2 production through respiration and sugar accumulation, which represent the glycolytic and gluconeogenic carbon flux, respectively. Both respiration and the expression of protein-associated gluconeogenesis-glycolysis, such as phosphoenolpyruvate carboxykinase, sucrose synthase and enolase, were depressed by O3 (Fig. 7). This finding suggests that carbon can be diverted in gluconeogenic (sucrose biosynthesis) and at the same time in glycolytic (CO2 production) pathways in kiwifruit undergoing ripening inhibition. This is consistent with the patterns of malate dehydrogenase abundance described below and collectively suggests that additional regulators besides endogenous ethylene may govern kiwifruit ripening inhibition. In kiwifruit, remorins and ‘Viral A-type inclusion protein repeat-containing protein expressed’ were induced by ripening and reduced by O3 (Fig. 7). Because the above-mentioned proteins were specifically depressed by O3 (Fig. 7), down-accumulation of these proteins plays a role in kiwifruit ripening inhibition.

Our proteomic analysis revealed accumulation of several kiwifruit proteins (n = 16) in response to combined 1-MCP and O3 (Figs. 5 and 7, Additional file 4: Table

![Fig. 11. Kiwifruit ripening-related genes and transcription factors expression as affected by 1-MCP, O3 and exogenous ethylene. Gene expression of aminocyclopropane-1-carboxylic acid oxidase 1 (ACO1, a-e), ethylene receptor (ETR, f-j), expansin 2 (EXP2, k-o), geranylgeranyl diphosphate synthase (GGPS, p-t) and lipoxygenase 1 (LOX1, u-y) in kiwifruit ripened under ambient air or exogenous ethylene (ETH) at various ripening time points at 20 °C following 6 months of cold storage. Real-time RT-qPCR was used to analyse the relative mRNA abundance on three biological repeats per treatment. Vertical bars in figure plates represent the least significant difference (LSD, P = 0.05), which was used for comparisons of means between treatments (exposed or not to exogenous ethylene) and ripening time points.](image-url)
expression pattern (Fig. 4e), thus indicating that repression with greater firmness retention (Fig. 1g-l) coneogenesis by inducing malate dehydrogenase. Our re-

of malate dehydrogenase and natterin were regulated by control point to prevent kiwifruit softening. An isoform of cell wall remodeling by these chemicals is a critical propaga-
tion/storage-associated proteins (e.g., HSP70) (Fig. 9, Additional file 4: Table S1) was defined as the central core protein in the created interacting network among the identified proteins of the present study. Enolase, an enzyme which is also called phosphopyruvate hydratase, is responsible for the catalysis of 2-phosphoglycerate (2-PG) conversion to phosphoenolpyruvate (PEP), in the ninth step of glycolysis. The down-accumulation of two enolase isoforms by the two kiwifruit ripening inhibitors O₃ and 1-MCP and their induction by exogenous ethylene (Figs. 7 and 8; Additional file 4: Table S1) reveals a potential association of enolase in kiwifruit climacteric ripening regulation. The above observation of increased enolase accumulation during fruit ripening is in agreement with previous reports of kiwifruit and tomato fruit experiencing ripening as a consequence of exogenous ethylene exposure [4, 38]. In kiwifruit exposed to exogenous ethylene, the major group of energy-associated proteins interacts with five other protein clusters: disease/defence-associated (e.g., lactoylglutathione lyase), protein destination/storage-associated proteins (e.g., HSP70), signal transduction-related (nucleoside diphosphate kinase), and secondary metabolism-associated (terpene synthase), further highlighting the importance of ethylene in kiwifruit ripening (Fig. 9b, Additional file 9: Table S4).

Bioinformatic analysis using BiNGO [21] was able to predict the major molecular functions of the identified proteins that are altered in kiwifruit experiencing ripening inhibition or induction, including lyase activity (5 S1). Accumulation of several intracellular chaperones, such as HSP 70 and chaperonin CPN60, in fruit exposed to 1-MCP + O₃ (Fig. 7) may be associated with its improved postharvest performance after long cold storage, since chaperones promote fruit cell survival following cold stress [28]. A receptor-like protein kinase (RLK) and a major latex-like protein (MLP) were specifically up-accumulated by 1-MCP + O₃ and this response was reversed by exogenous ethylene (Fig. 8, Additional file 4: Table S1). Plant RLKs are transmembrane proteins that perceive signals of environmental conditions and developmental status through their extracellular domains and propagate the signals via their intracellular kinase do-

mains [29]. MLP proteins belong to the Bet v 1 family which act through binding ligands such as cytokinins, brassinolides or secondary metabolites, and trigger downstream signal transduction [30]. The observed changes in abundance of specific isoforms of Bet v 1-related allergen caused by either O₃ or 1-MCP + O₃ (Fig. 7, Additional file 4: Table S1) provide further insights for a potential role of Bet v 1-related allergen during kiwifruit ripening, as was previously reported [4]. 1-MCP + O₃ induced two isoforms of lactoylglutathione lyase (Fig. 7, Additional file 4: Table S1) that were de-

pressed by subsequent exposure to exogenous ethylene (Fig. 8, Additional file 4: Table S1). Lactoylglutathione lyase participates in glutathione-based detoxification of methylglyoxal (methylglyoxal pathway) and was identified via proteomic analysis in various plant systems [31]. This finding and the fact that glyceraldehyde 3-phosphate dehydrogenase, also involved in the methyl-
glyoxal pathway, was affected by exogenous ethylene (Fig. 8, Additional file 4: Table S1) are consistent with reports that involve this pathway with fruit ripening [32, 33], as a potential mechanism to control methylglyoxal levels [34]. Although the two single treatments (1-MCP and O₃) induced similar inhibition of ethylene biosynthesis, there was little overlap in protein signatures among them (Fig. 5), demonstrating that these substances may inhibit kiwifruit ripening through different pathways. However, there is also evidence that they use similar mechanisms to inhibit ripening. Accumulation of polygalacturonase (PG) and beta-D-galactosidase decreased in kiwifruit exposed to either 1-MCP or O₃ (Fig. 7), which is consistent with greater firmness retention (Fig. 1g-l) and the PG expression pattern (Fig. 4e), thus indicating that repression of cell wall remodeling by these chemicals is a critical control point to prevent kiwifruit softening. An isoform of malate dehydrogenase and natterin were regulated by all postharvest treatments (Fig. 7). Malate dehydrogenase is involved in gluconeogenesis in fruit [35] and our results indicate that ripening inhibition may promote glu-
coneogenesis by inducing malate dehydrogenase. Natterin-like proteins are pore-forming, highly toxic complexes, that may be associated with the defense mechanism of specific animals [36]. Although the active role of natterin in higher plants is unknown, our previous study found this protein up-regulated in kiwifruit undergoing artificial ripening [4]. However, the mechanisms underlying the function of natterin during kiwi-
fruit ripening remains unclear.
proteins) and copper ion binding (4 proteins) in the former but only copper ion binding (6 proteins) in the latter (Fig. 10b, d; Additional file 8: Table S3). Bioinformatic analysis also indicated that stress response was the most affected biological pathway in kiwifruit treated with ripening inhibitors (Fig. 10c, Additional file 7: Table S2). Thirteen proteins were classified in the general category of response to stress, including phosphoenolpyruvate carboxykinase, pyruvate decarboxylase, malate dehydrogenase, HSP 70, fructose-bisphosphate aldolase, nucleoside diphosphate kinase, enolase, pathogenesis-related protein, and others (Fig. 10c, Additional file 7: Table S2a). Three other biological pathways that were over-expressed in kiwifruit experiencing ripening inhibition were the response to cadmium ion (6 proteins), response to metal ion (6 proteins), and response to stimulus (13 proteins). In kiwifruit experiencing ripening as a result of exogenous ethylene, the biological pathways over-represented included stress response (16 proteins), glucose catabolism (5 proteins), monosaccharide catabolism (5 proteins), and hexose catabolism (5 proteins) (Fig. 10d, Additional file 7: Table S2b).

Both 1-MCP and O₃ regulate expression of ripening-related gene and transcription factors

In addition to the protein changes described above, a crucial set of genes actively involved in kiwifruit ripening [3, 9, 10] was investigated. Notably, 1-MCP and O₃ treated fruits showed differences in expression of several ripening-related genes (Fig. 11), suggesting that these endogenous ethylene inhibitors act largely independently. The GGPS reaction produces geranylgeranyl diphosphate (GGPP), a common precursor for the synthesis of phytol, tocopherols, plastoquinones, chlorophylls, carotenoids, gibberellins, and other hormones [39]. GGPS (Fig. 10p-r, s, t), ACO1 (Fig. 10a-c, d, e) and ETR (Fig. 11f-h, i, j) expression was strongly depressed by O₃ and stimulated by exogenous ethylene. This suggests that exogenous ethylene not only acts downstream of ethylene synthesis and signaling, but also as a regulator of various isoprene-containing ripening compounds in O₃-treated fruits. EXP2, which is involved in cell wall-loosening [40], exhibited a pattern of induced expression similar to PG in O₃-treated fruit exposed to exogenous ethylene (Fig. 4e). Ethylene-induced EXP2 expression might affect access of hydrolases to cell wall polymers and promote cell wall disassembly and the subsequent fruit softening [41], thereby acting with PG to fine-tune cell wall metabolism in ripening O₃-treated kiwifruit. In contrast, expression of ETR, EXP2, GGP and LOX1 under exogenous ethylene remained lower in fruit exposed to 1-MCP + O₃ (Fig. 10), consistent with the displayed ripening inhibition (Figs. 1 and 2), most likely as a consequence of decreased ethylene sensitivity under such conditions (Fig. 3). The differences in 1-MCP- and O₃-dependent ripening inhibition are further supported by the contrasting LOX1 pattern during kiwifruit ripening in ambient air (Fig. 10). LOX1, which catalyzes hydroperoxidation of polyunsaturated fatty acids, regulates fruit ripening through ethylene-dependent and -independent pathways [42, 43]. These data suggest that ACO, ETR, LOX1, GGP, PG and EXP2 are associated with kiwifruit ripening regulation and provide new perspective on understanding 1-MCP- and O₃-mediated ripening inhibition.

Conclusions

This is the first study that shows the combination of 1-MCP and O₃ severely inhibits ethylene production and reduces softening rates in kiwifruit, leading to depressed ripening even under exogenous ethylene exposure. Endogenous ethylene biosynthesis inhibition in kiwifruit by long-term exposure to O₃-enriched cold storage is reversible by post-storage exogenous ethylene exposure. Protein and gene expression analysis showed that 1-MCP and O₃ have both common and (mostly) unique roles in kiwifruit ripening. Taken together, our results provide a physiological basis for future research on the implications of both 1-MCP and O₃ in climacteric fruit ripening.

Methods

Fruit material, postharvest treatments and experimental approach

‘Hayward’ kiwifruit grown in a commercial orchard (Naousa, Central Macedonia, Greece) were harvested at the stage of physiological maturity (fresh weight: 93.4 ± 2 g; pericarp tissue firmness: 63.8 ± 1.9 N; core tissue firmness: 144.5 ± 5.4 N; SSC: 7.6 ± 0.2%; TA (citric acid, %): 1.9 ± 0.1%; and DMC: 17.4 ± 0.5%). Fruits were divided randomly into lots (113 lots) of 30 fruits. The initial quality of kiwifruit was analyzed in one lot immediately after harvest and the rest were split into two groups (56 + 56) and treated with or without 1-MCP (0.6 µL L⁻¹ SmartFresh, AgroFresh Inc., Rohm and Haas, Spring House, PA, USA) for 24 h at 0 °C using a 4000-L airtight tent in the cold room, according to manufacturer’s instructions. Immediately after 1-MCP treatment, fruits of both groups were cold-stored (0 °C, 95% RH) in two separate cold rooms in which ethylene was oxidized by catalytic ethylene oxidation (Swintherm model BS 500, Fruit Control Equipments s.r.l., Milano, Italy). Cold room atmosphere was ambient (control) or enriched with 0.3 µL L⁻¹ ozone through a dedicated system of continuous ozone generation and monitoring (oxygen generator model SEP-100, ozone generator model COM-AD-04 and ozone analyser model MP-6060, Anseros Klaus Nonnenmacher GmbH, Tübingen,
Germany). Ethylene concentration in the cold storage rooms was monitored and was below the accepted threshold concentration for commercial kiwifruit storage (10 nL L\(^{-1}\)) and not significantly different from each other (data not shown). After 2, 4 and 6 months, fruits were removed from cold storage and held at 20 °C (90% RH), where kiwifruit ripening was analyzed either immediately, or after 2, 4, 6, 8, 10, 12 and 14 d.

Postharvest treatments were segregated based on their ability to inhibit endogenous ethylene biosynthesis. Experimental kiwifruit were further examined to determine their ability to recover from ripening inhibition by a short treatment with exogenous ethylene. Kiwifruit cold-stored for 4 months and maintained for 8 d at 20 °C were treated with 100 μL L\(^{-1}\) exogenous ethylene (20 °C, 90% RH, 24 h). Exogenous ethylene treatment was repeated after 6 months of cold storage plus 1 day of ripening in all treatments to validate the results obtained following 4 months of cold storage. Exogenous ethylene was applied in an airtight tank (100 L) with an internal fan to circulate air. Carbon dioxide was absorbed in a 500 mL solution of 4 M NaOH. Following the 24-h exposure, the ethylene concentration in the tank was 102 and 98 μL L\(^{-1}\) and the CO\(_2\) concentration was 0.45 and 0.39% after 4 and 6 months of cold storage, respectively.

Kiwifruit ripening following cold storage was characterized using ethylene emission rate; CO\(_2\) emission rate (respiration rate, RR); pericarp and core tissue firmness; DMC, SSC and TA, as described [4]. Additionally, pericarp tissue was sampled from each biological replicate per treatment (3 replicates of pericarp flesh tissue content of 10 fruits each), frozen immediately in liquid nitrogen, and finally stored for further analysis (−80 °C). In summary, kiwifruits in the present study were exposed to 4 postharvest treatments, namely, control, 1-MCP, O\(_3\) and 1-MCP + O\(_3\). In addition, a separate treatment with exogenous ethylene (ETH) exposure to postharvest treatments was applied after 4 and 6 months of cold storage, as presented schematically in Additional file 10: Figure S6.

Kiwifruit ripening physico-chemical changes

Firmness of pericarp and core tissue was determined using a fruit texture analyzer (model 53,205, T.R. Turoni srl, Forlì, Italy) and expressed as newtons (N) following the methodology previously described [4]. Soluble solids concentration and TA were assessed as described [4]. Dry matter content was measured in 2 mm thin cylindrical slices from 3 biological replications of 10 fruits as described [44]. Mean values of physico-chemical data (3 biological repetitions) were subjected to analysis of variance (ANOVA) and least significant differences (LSD) (\(P = 0.05\)) calculated using the statistics package SPSS 22.0 for Mac OS X (SPSS, Chicago, IL, USA).

Ethylene emission and respiration rates

Ethylene emission and respiration rates (RR) in 3 repetitions of 3 kiwifruit per treatment during ripening were analyzed using a gas chromatograph system and an infrared gas analyzer, respectively, as previously described [17]. Statistical analysis was as described above.

Analysis of ethylene biosynthesis intermediates and enzyme activities

1-Aminocyclopropane-1-carboxylic acid (ACC) and 1-malonyl-aminocyclo-propane-1-carboxylic acid (MACC) concentrations and ACC synthase (ACS) and ACC oxidase (ACO) enzyme activities were analyzed as previously described [45]. Statistical analysis was as described above.

Kiwifruit soluble protein extraction, 2D-gel electrophoresis and image quantification

Soluble proteins of kiwifruit pericarp tissue were extracted as previously described [46]. Protein concentrations were measured according to Bradford’s method [47]. First and second dimension separation of protein extracts (50 μg) was performed in three biological replications per treatment as described [4]. 2D-gels staining and scanning, and image quantification and analysis performed as previously described [4, 48]. Quantitative protein spot abundance mean values comparisons were performed by one-way analysis of variance using Student’s t-test (\(P = 0.05\)). The statistically different means were further subjected to a quantitative criterion of 1.5-fold change for significant differences determination (Additional file 11: Table S5).

Protein in-gel digestion and identification by mass spectrometry

Selected spots of interest on the 2D-gels were stained, isolated and trypsin digested as described [4, 19, 49]. Database searches were conducted against the Cornell University kiwifruit protein database (http://bioinfo.bti.cornell.edu/cgi-bin/kiwi/download.cgi) containing 39,004 protein sequences and the National Center for Biotechnology Information (NCBI) databases using BLASTp analyses and MASCOT software as described [4, 19]. Validation of significant differences through the two-way hierarchical clustering was done with Permut Matrix software. Zero-mean and unit-standard deviation was used for the row-by-row normalization of data. Analysis was performed using Pearson’s distance and Ward’s algorithm. Identification criteria, among the positive matches, included at least 2 different peptide sequences of over 6 amino acids with an individual score above 20
RNA isolation and reverse transcription quantitative real-time PCR (RT-qPCR) analysis

Total RNA was isolated and cDNA synthesis was performed as described [19, 51]. Target cDNAs amplification was performed with gene-specific primers (Additional file 12: Table S6) designed as described [19]. Quantitative RT-PCR reactions were performed as described [19]. The specificity of the primers was determined by the dissociation kinetics for the PCR products at the end of each run. Actin (A. deliciosa) was used as reference gene. Relative transcription of the gene of interest and PCR efficiency were calculated as described [19, 52]. Three biological replicates of each treatment were performed and used for gene expression experiments.

Protein-protein interaction network bioinformatic analysis

A protein-protein interaction network (PPI) analysis and prediction of the potential biological processes and molecular functions was performed to shed light on interaction functions of identified proteins as described [4, 20, 21]. Search parameters and statistical analysis of the annotated protein entries for PPI and biological processes and molecular functions analysis were obtained by blasting identified proteins against TAIR10 (The Arabidopsis Information Resource). Annotated proteins with the highest score and lowest E-value were considered relevant for each identified protein (Additional file 9: Table S4) [4, 20, 21].

Additional files

**Additional file 1: Figure S1.** Changes in dry matter content (DMC) in kiwifruit during ripening at 20 °C following 2 or 6 months of cold storage (0 °C, RH 90%). Vertical lines indicate LSD (P = 0.05) of three replicate samples, each consisting of 10 cylindrical slices coming from 10 separate fruit. (PPTX 85 kb)

**Additional file 2: Figure S2.** Soluble solids concentration (SSC, a, c, e) and titratable acidity (TA, b, d, f) in 1-MCP/O3-treated kiwifruit during ripening at 20 °C following 0 or 6 months of cold storage (0 °C, RH 90%). Vertical lines indicate LSD (P = 0.05) of three replicate samples, each consisting of 10 fruit. (PPTX 110 kb)

**Additional file 3: Figure S3.** Soluble solids concentration (SSC, a, c) and titratable acidity (TA, b, d) in 1-MCP/O3/ETH-treated kiwifruit during ripening at 20 °C following 4 or 6 months of cold storage (0 °C, RH 90%). Vertical lines indicate LSD (P = 0.05) of three replicate samples, each consisting of 10 fruit. Markers and lines in grey represent kiwifruit untreated with exogenous ethylene as in Additional file 2: Figure S2. (PPTX 114 kb)

**Additional file 4: Table S1.** Information data and responsiveness to 1-MCP, O3, and exogenous ethylene treatments of identified kiwifruit proteins analyzed by 2DE-PAGE and nanoLC-MS/MS. (XLSX 29 kb)

**Additional file 5: Figure S4.** Reference maps for kiwifruit proteins per treatment. (PPTX 463 kb)

**Additional file 6: Figure S5.** Venn diagram showing the overlapping and unique kiwifruit proteins quantified in each single treatment exposed to exogenous ethylene (control-ETH, 1-MCP-ETH, O3-ETH, 1-MCP + O3-ETH) compared to their counterparts untreated with ethylene. (PPTX 149 kb)

**Additional file 7: Table S2.** (a) Biological pathways and networks analysis generated by BiNGO for proteins with altered abundance in kiwifruit treated with ripening inhibitors but not with exogenous ethylene. (b) Biological pathways and networks analysis generated by BiNGO for proteins with altered abundance in kiwifruit undergoing ripening with exogenous ethylene. (XLSX 15 kb)

**Additional file 8: Table S3.** (a) Molecular functions and networks analysis generated by BiNGO for proteins with altered abundance in kiwifruit treated with ripening inhibitors but not with exogenous ethylene. (b) Molecular functions and networks analysis generated by BiNGO for proteins with altered abundance in kiwifruit undergoing ripening with exogenous ethylene. (XLSX 12 kb)

**Additional file 9: Table S4.** Identified proteins of kiwifruit that Blasted against the TAIR database and their STRING ID. (XLSX 21 kb)

**Additional file 10: Figure S6.** Experimental design. (PPTX 46 kb)

**Additional file 11: Table S5.** Quantitative data of protein spot volumes in kiwifruit on 2DE-PAGE gels. (XLSX 94 kb)

**Additional file 12: Table S6.** Primers used to perform q-RT PCR analysis. (DOCX 20 kb)

**Abbreviations**

1-MCP: 1-methylcyclopropene; ACC: 1-aminocyclopropane-1-carboxylic acid; ACO: ACC oxidase; ACO1: 1-aminocyclopropane-1-carboxylic acid oxidase; ACS: ACC synthase; DMC: Dry matter content; ETRI: Ethylene receptor; EXP: Expansin; GGP1: Geranylgeranyl diphasphate synthase; LOX1: Lipoxygenase; MACC: 1-malonyl-aminocyclo-propane-1-carboxylic acid; MS/MS: Mass spectrometry; NCBI: National Center for Biotechnology Information; O3: Ozone; PG: Polygalacturonase; qRT-PCR: Quantitative reverse transcriptase polymerase chain reaction; SSC: Soluble solids concentration; TA: Titratable acidity

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**Availability of data and materials**

All data generated or analysed during this study are included in this published article and its supplementary information files.
Authors’ contributions
ISM, AM, MV and KKP designed the study. ISM, GT, EK, AK and MB carried out the experimental work and data analysis. ISM interpreted data and performed graph artwork and wrote the first draft of the manuscript. AM, GT, MV, MB, EK and KKP edited the other versions. All authors have read and approved the manuscript.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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