Postharvest Treatment of Hydrogen Sulfide Delays the Softening of Chilean Strawberry Fruit by Downregulating the Expression of Key Genes Involved in Pectin Catabolism

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Abstract: Hydrogen sulfide (H₂S) plays several physiological roles in plants. Despite the evidence, the role of H₂S on cell wall disassembly and its implications on fleshy fruit firmness remains unknown. In this work, the effect of H₂S treatment on the shelf-life, cell wall polymers and cell wall modifying-related gene expression of Chilean strawberry (Fragaria chiloensis) fruit was tested during postharvest storage. The treatment with H₂S prolonged the shelf-life of fruit by an effect of optimal dose. Fruit treated with 0.2 mM H₂S maintained significantly higher fruit firmness than non-treated fruit, reducing its decay and tripling its shelf-life. Additionally, H₂S treatment delays pectin degradation throughout the storage period and significantly downregulated the expression of genes encoding for pectinases, such as polygalacturonase, pectate lyase, and expansin. This evidence suggests that H₂S as a gasotransmitter prolongs the post-harvest shelf-life of the fruit and prevents its fast softening rate by a downregulation of the expression of key pectinase genes, which leads to a decreased pectin degradation.

Keywords: Fragaria chiloensis; strawberry decay; shelf-life; cell wall disassembly; pectinases; gasotransmitter

1. Introduction

The Chilean strawberry [Fragaria chiloensis (L.) Mill.] is a native species from South America and the maternal progenitor of the commercial strawberry Fragaria × ananassa Duch. [1]. The Chilean strawberry is a non-climacteric fruit that possesses remarkable organoleptic properties, such as good taste, aroma, nutritional value and an exotic white fruit appearance, having a great potential to become a new exotic berry fruit for the worldwide market [2–4]. Besides, F. chiloensis fruit is emerging as a new model for studying several ripening-associated processes in strawberries [5,6], such as anthocyanin biosynthesis and plant cell wall disassembly [7,8]. Studies on fruit softening are important as this influences the postharvest life of highly perishable fruit [9].

The ripening-associated softening of fleshy fruit has been largely described as a direct consequence of enzyme-mediated cell wall disassembly [10]. Events such as depolymerization and solubilization of hemicelluloses and pectins within the cell wall...
often occur in many fleshy fruits, including strawberry [7]. Genes encoding for proteins involved in cell wall disassembly, such as polygalacturonase (PG), pectate lyase (PL), pectin methylesterase (PE), endo-β,1,4-glucanase (EG), β-galactosidase and expansins (EXP), increase their expression during strawberry ripening [7,11–14]. Previous reports have shown that Chilean strawberry fruit has a faster-softening rate than F. x ananassa ‘Chandler’ fruit, which is the major disadvantage for the commercialization of this fruit [12]. The higher softening rate of F. chiloensis fruit has been associated with a high expression level of the PG gene during fruit ripening [12]. Several reports indicate that pectin metabolism has a significant role in strawberry fruit firmness [7,13,15,16] rather than hemicellulose or cellulose metabolism [17]. However, hemicellulose modification might play a role in Chilean strawberry fruit softening [18]. Regarding the evidence, it is relevant to gain insights about signal molecules that regulate at the biochemical and molecular level and this biological process during the ripening and postharvest of strawberry fruit.

Hydrogen sulfide (H2S) is a gas molecule traditionally associated with phytotoxins [19]. However, evidence in animal systems has demonstrated its roles as an endogenous signal molecule and it has been proposed as a gaseous regulator of various physiological functions, similar to nitric oxide (NO) and carbon monoxide (CO) [20–22]. Increasing evidence in plants suggests that H2S plays several physiological roles, such as enhancer of photosynthesis, stomatal movement, seed germination, root organogenesis, delay senescence of cut flowers and fresh fruit, and biotic and abiotic stress tolerance alone or through interaction with plant hormones [22–29]. Furthermore, H2S is biosynthesized in plants from sulfite by sulfite reductase, and from cysteine by L/D-cysteine desulphhydrase and β-cyanoalanine synthase [30–32]. Additionally, H2S can be removed enzymatically from plant tissue by the action of O-acetylserylserine (thiol) lyase [33]. Altogether, H2S physiological roles and metabolism evidenced in plants suggest that it might have a role as an endogenous gaseous regulator [32].

H2S-fumigated fruit, using several donors of this gasotransmitter, prolongs its postharvest shelf-life and increases the antioxidant capacity of the tissues, reducing the levels of reactive oxygen species (ROS) and ROS-induced damage [29]. Interestingly, it has also been described as the impact of H2S in postharvest physiology of several climacteric (e.g., banana and tomato) and non-climacteric (e.g., grape and strawberry) fruits [34–38]. In banana, H2S treatment sustained fruit chlorophyll content, increased carotenoids, soluble proteins, and the overall antioxidant capacity [34]. Authors suggest that H2S delayed banana fruit ripening and senescence via an antagonizing effect with ethylene, through the alleviation of oxidative stress and inhibition of ethylene signaling [34]. In this sense, H2S application to tomato fruits delayed the acquisition of color and maintained higher chlorophyll and nutritional-related metabolites content during ripening [35]. In the case of non-climacteric fruits, fumigation with H2S of grape berries, prior to postharvest storage, preserved high levels of several quality markers such as firmness, soluble solids, titratable acidity (TA), and relevant metabolites such as ascorbic acid, carotenoids, flavonoids, total phenolics, reducing sugars, and soluble proteins [36]. In the same work, H2S fumigation reduced the accumulation of ROS and malondialdehyde (MDA) in grape pulp, while it increased the activity of antioxidant enzymes [36]. In strawberry (F. x ananassa) fruit, postharvest H2S treatment also increased the activity of antioxidant-related enzymes and maintained fruit firmness mainly by a decrease of the activity of cell wall-modifying enzymes such as PG, PE, and EG prolonging the shelf-life of strawberries [37,38].

Thus, according to recent evidence, we hypothesized that H2S applied at harvest can delay the softening of Chilean strawberry fruit during its postharvest period by affecting cell wall disassembly. In the present work, we study the effect of H2S treatment on the metabolism of cell wall polysaccharides, i.e., pectin and hemicellulose catabolism and the implications on the expression of genes encoding for enzymes involved in cell wall modification during postharvest-associated fruit softening.
2. Results

2.1. Effects of H₂S Treatment on Postharvest Shelf-Life of Chilean Strawberry Fruit

The effect of H₂S gas, produced by its donor NaHS, was studied during the shelf-life of Chilean strawberry fruit at 20 °C. Different NaHS concentrations (0.2, 0.4, 0.8 and 1.2 mM) were assayed. Fruit decay was evaluated during the postharvest period and the limit of shelf-life was set at a decay index of 40 (Figure 1A). H₂S treatment reduces fruit decay and prolongs the postharvest shelf-life of the Chilean strawberry fruit by an effect of optimal dose (Figure 1). Untreated strawberries rapidly decay during shelf-life, reaching the disposal limit after only two days at 20 °C. By contrast, strawberries treated at harvest with NaHS are less prompted to fungal infection and displayed a slower decay.

Fruit exposed to 0.2–1.2 mM NaHS solutions displayed a longer shelf-life than untreated fruit, nevertheless, the maximum effect was reached with 0.2 mM NaHS (Figure 1A). In subsequent experiments, 0.2 mM NaHS was employed, extending the shelf-life period (14 days) (Figure 2). NaHS-treated fruit reached their shelf-life limit after 6 d of storage; while untreated fruit reached this stage after 2 d, thus the shelf-life period of NaHS-treated fruit was nearly tripled (Figure 2A). Interestingly as shown in Figure 2B, after 6 d of storage, untreated fruit evidenced severe signs of fungal infection, while NaHS-treated strawberries were not affected.

![Figure 1](image-url). Effects of different H₂S donor concentrations on rot index and appearance during postharvest shelf-life of Chilean strawberry (*Fragaria chiloensis*) fruit. Strawberries were treated after harvest with H₂S donor NaHS at different concentrations (0, 0.2, 0.4, 0.8 and 1.2 mM). Changes in (A) rot index; (B) Appearance of strawberry fruit at harvest and (C) after exposure to H₂S donor NaHS during 3 d of storage at 20 °C.
2.2. Effects of H₂S Treatment on Fruit Color during the Shelf-Life of Chilean Strawberry Fruit

Changes in the external color of strawberries during storage, both in control and NaHS-treated fruit, were evaluated throughout the shelf-life period (Figure 2C). Untreated fruit showed a significant decrease in lightness (L*), while fruit exposed to 0.2 mM NaHS maintained this color parameter during 4 d of shelf-life (Figure 2C). In addition, control and NaHS-treated fruit displayed a similar decreasing pattern of other color parameters (a* and b* values) during shelf-life (Figure 2D,E). Color measurements were not reliable at longer shelf-life periods due to the presence of fungal development on the surface of control strawberries.

Figure 2. Effects of H₂S treatment on decay index and color parameters during postharvest shelf-life of Chilean strawberry (Fragaria chiloensis) fruit. Strawberries were treated after harvest with 0.2 mM H₂S donor NaHS. Changes in (A) decay index; (B) Images of control and H₂S-treated strawberries after 6 d of storage at 20 °C, (C–E) Color parameters showing changes in L, a*, b* values, respectively. Each value is the mean of three replicates and vertical bars represent standard errors. Different letters indicate significant differences at p < 0.05. For details, see Section 4.
2.3. Effects of H$_2$S Treatment on Fruit Softening and Respiration Rate of Chilean Strawberry Fruit during the Shelf-Life Period

Changes in fruit firmness were followed during the storage period in control and NaHS-treated fruit. Untreated Chilean strawberry fruit evidenced a rapid decrease in firmness throughout the shelf-life period; nevertheless, fruit treated with 0.2 mM NaHS remained firmer than untreated fruit (Figure 3A). Major differences in firmness between treatments were recorded after 4 d at 20 °C, with values of 1.61 N for NaHS-treated fruit and 0.76 N for untreated fruit. Firmness values of untreated fruit were not reliable after 6 d of storage.

As H$_2$S is an acid molecule, titratable acidity (TA) was determined to investigate whether exposure to NaHS treatment influences changes in acidity (Figure 3B). TA values increased both in NaHS-treated and control fruit, although with a different pattern. In untreated fruit, TA increases immediately after harvest, while in NaHS-treated fruit, there is a delay as no changes in acidity take place during the first two days. At 6 d of storage, the same acidity level was recorded in both fruit conditions.

Changes in respiration rate were also determined during the shelf-life period (Figure 3C). The production of CO$_2$ of NaHS-treated fruit was maintained at a lower rate during shelf-life, and in contrast, it increased constantly in untreated fruit until 4 d. The respiration rate of untreated fruit was not measured at 6 d of storage due to the presence of fungus on the fruit surface, which also releases CO$_2$, interfering with the determination of the fruit tissue.

Changes in soluble solids content (SSC) were also followed in control and NaHS-treated fruit during shelf-life (Figure 3D). SSC decreased in both fruit groups during shelf-life with no differences.

Figure 3. Effects of H$_2$S treatment on firmness, titratable acidity, respiration rate and soluble solids content during postharvest shelf-life of Chilean strawberry (Fragaria chiloensis) fruit. Strawberries were treated after harvest with 0.2 mM H$_2$S donor NaHS. Changes in (A) fruit firmness; (B) titratable acidity; (C) respiration rate, and (D) soluble solids content. Each value is the mean of three replicates and vertical bars represent standard errors. Different letters indicate significant differences at $p < 0.05$. For details, see Section 4.
2.4. Effects of H$_2$S Treatment on Cell Wall Polymer Solubilization during the Shelf-Life of Chilean Strawberry Fruit

Cell wall fractionation was performed from AIR samples prepared from *F. chiloensis* fruit samples subjected to H$_2$S treatment. Total cell wall yield was subjected to a sequential fractionation procedure to separate several pectin fractions. The fractionation of pectins in WSF, CSF, and NSF fractions corresponds to loosely-, ionically-, and covalently-bound pectins, respectively, while KSF is mainly associated with the hemicellulose fraction. During the postharvest period, significantly higher solubilization of pectins was observed in the control fruit than in H$_2$S-treated fruit (Table S1, Figure 4). The solubilization of pectins advised as the increment in WSF is accompanied by a decrease in CSF and NSF fractions in non-treated fruit (Figure 4B–D). In H$_2$S-treated fruit, pectin solubilization was delayed as WSF did not increase and CSF and NSF did not decrease as in the control fruit. All this evidence suggests that H$_2$S treatment delays pectin degradation in Chilean strawberry fruit during the shelf-life period.

**Figure 4.** Effects of H$_2$S treatment on the content of pectin-related polymers during postharvest shelf-life of Chilean strawberry (*Fragaria chiloensis*) fruit. Strawberries were treated after harvest with 0.2 mM H$_2$S donor NaHS, and sampling was performed at harvest (0 d) and after 2, 4 and 6 d of storage at 20 °C. The content of uronic acids (UA) was determined in the water (WSF), CDTA (CSF)- and Na$_2$CO$_3$ (NSF)-soluble cell wall fractions. Changes in (A) water-soluble pectins; (B) CDTA-soluble pectins; (C) Na$_2$CO$_3$-soluble pectins, and (D) the relative content (%) of several pectin fractions in control and H$_2$S-treated fruits during postharvest. Values correspond to the mean of three independent cell wall extractions per sampling date. Each value is the mean of three replicates and vertical bars represent standard errors. Different lowercase letters indicate significant differences between control and treatment conditions in each day at *p* < 0.05. For details, see Section 4.

Regarding the solubilization of hemicelluloses, an increase in its content was observed in cell wall material obtained from control fruit during the shelf-life period;
however, in H₂S-treated fruit, this increase is delayed in time (Table S1). The content of hemicelluloses of control fruit after 2 d of shelf-life was reached after 6 d of storage by H₂S-treated fruit.

2.5. Effects of H₂S Treatment on the Expression of Genes Involved in Pectin Degradation during Strawberry Shelf-Life

To gain molecular insights to explain fruit firmness changes and pectin solubilization in H₂S-treated fruit, the expression of genes involved in cell wall metabolism was analyzed during shelf-life (Figure 5). The expression of genes encoding enzymes involved in pectin solubilization such as polygalacturonase (FcPG1) and pectate lyase (FcPL1), displayed a drastic and fast reduction in response to H₂S treatment (Figure 5A,B). By contrast, untreated fruit displayed a slow reduction in the expression level of both genes. The expression of these genes reached the lowest levels after 6 d of shelf-life in both treatments. Remarkably, a gene encoding an isoform of expansin (FcEXP2) showed a similar expression profile to that of FcPG1 and FcPL1 (Figure 5C). The expression of a gene encoding for the enzyme xyloglucan endotransglycosylase/hydrolase 1 (FcXTH1), involved in molecular modifications of hemicellulose did not change its transcriptional level during the first two shelf-life days either in control or H₂S-treated fruit (Figure 5D). From the fourth day, a strong reduction in FcXTH1 transcript levels was observed in H₂S treated fruit, while in contrast, in non-treated fruit, there is an increment in their levels. Furthermore, the expression of a gene that encodes for endo-β-1,4-glucanase 1 (FcEG1), with cellulase activity, recorded a similar pattern to that of FcXTH1 both in H₂S-treated and controls groups (Figure 5E); however, FcEG1 transcripts were not detected in untreated fruit at 6 d of storage.
Figure 5. Effects of H₂S treatment on the expression of cell wall disassembly- and remodeling-related key genes during postharvest shelf-life of Chilean strawberry (Fragaria chiloensis) fruit. Strawberries were treated after harvest with 0.2 mM H₂S donor NaHS and maintained under storage at 20 °C for up to 6 d. Sampling was performed at harvest (0 d) and after 2, 4 and 6 days. Gene expression profile of (A) polygalacturonase 1 (FcPG1); (B) pectate lyase 1 (FcPL1); (C) expansin 2 (FcEXP2); (D) xyloglucan transglycosylase-hydrolase 1 (FcXTH1), and (E) endo-β-1,4-glucanase 1 (FcEG1). The data were analyzed by Tukey test ($p < 0.05$) per time. Each value represents mean ± SD ($n = 4$). Asterisks indicate significant differences between treatments. For details, see Section 4.

2.6. Principal Components and Correlation Analyses on Studied Parameters in H₂S-Treated Strawberries

To understand the impact of H₂S treatment in strawberry fruit concerning the analyzed parameters and to identify correlations between them, principal component analysis (PCA) and correlation analysis were performed. The responses are mainly explained by the first two main components (PCA 1: 45.22%; PCA 2: 22.18%) revealing that the most significant variables for the principal component 1 (PC1) were those related
to CSF, NSF, and the fruit color (value of L* parameter), which showed negative correlations with treatment time, respiratory rate, and decay index. The principal component (PC2) showed a higher relationship factor for the FcPL1, FcEXP2, FcPG1 genes, and the fruit firmness, showing a negative correlation for the FcXTH1 gene and TA. It is worth noting the strong relationship between the ionically-bound pectin fraction (CSF) and the FcPL1 gene. H2S treatments influenced the fruit responses mainly at 4 d of treatment (Figure 6B). Control fruit showed, at this time, a higher contrasting relationship for the H2S-treated fruit, which experienced major changes in the analyzed variables.

H2S-treated strawberry fruit showed a strong positive correlation amongst firmness, the CSF and covalently-bound pectin (NSF) fractions, and genes involved in pectin solubilization (FcPG1, FcPL1, FcEXP2), exhibiting a higher significance than the control fruit (Figure 6C,D). Furthermore, exclusively in the treated fruit, the decay index was negatively correlated to these studied variables. On the other hand, the decay index, which indicates the fruit decay, evidenced a negative correlation with the FcXHT1 gene in the treated fruit, which showed a positive correlation in the control fruit. Additionally, this gene showed a negative correlation for the loosely-bound pectin fraction (WSF) in treated fruit. However, in the control group, the FcXHT1 gene was positively correlated with the WSF fraction, which is also positively correlated with the respiratory rate (Figure 6C,D).
Figure 6. Principal component (PCA) and correlation analyses of Chilean strawberry (*Fragaria chiloensis*) fruit in response to H$_2$S treatment during postharvest shelf-life: (A) Graph of variables; (B) Graph showing the response of *F. chiloensis* fruit to 0.2 mM NaHS treatment along storage time. Correlation graph of (C) control and (D) H$_2$S-treated fruit. For details, see Section 4.

3. Discussion

Until now, most of the research about the role of H$_2$S on the maintenance of fruit quality during postharvest storage has mainly been focused on the antioxidant system and its effect on the ethylene pathway. In this sense, the reduction in the accumulation of ROS and the increase in ascorbic acid, flavonoids, phenolics, and the enzymatic activities of ascorbate peroxidase (APX), catalase (CAT), peroxidase (POD), and superoxide dismutase (SOD) has been reported in fruits such as apple, pear, kiwifruit, grape, strawberry, and mulberry [36,37,39–42]. Regarding interference on ethylene pathway by H$_2$S treatment, it has been reported in tomato, a downregulation of fruit ripening-related genes such as those encoding for ethylene-responsive transcription factors *ERF003* and
In apple and kiwifruit, H$_2$S was linked with the suppression of the expression of genes involved in ethylene biosynthesis and signal transduction [39] and the inhibition of ethylene production [41], thereby supporting the counteractive role of H$_2$S in the ethylene pathway in climacteric fruits. Certainly, less is known about the specific role of H$_2$S on cell wall degradation and related gene expression during postharvest of fleshy fruits. In the present work, we showed evidence about the effect of H$_2$S on the pectin and hemicellulose catabolism and the expression of genes encoding for the main strawberry cell wall-modifying enzymes.

A quick decay, softening, and loss of fruit peel lightness are common processes during the senescence of _Fragaria chiloensis_ fruit [43,44]. In this work, it was found that the shelf-life of Chilean strawberry fruit was extended by H$_2$S treatment by an effect of optimal dose, reaching a most favorable effect with a dose of 0.2 mM NaHS donor, which was evidenced by the largest decrease in decay index and extending the shelf-life limit of treated fruit, around thrice compared to control fruit. The extended strawberry shelf-life was accompanied by a maintained firmness and color lightness, which indicates a significant delay in the senescence process of strawberry fruit tissue. These findings were consistent with previous studies [37] that have reported that H$_2$S prolongs the postharvest shelf-life of strawberry ( _F. × ananassa_ ‘Bao Jiao’) by an antioxidative role in fruit. Interestingly, Hu et al. [37] recorded that the optimal dose of H$_2$S donor NaHS was 0.8 mM, which is four times higher than the 0.2 mM required for the optimum effect in _F. chiloensis_ fruit (Figure 1). This result might be explained by differences in the perception and signaling pathway of H$_2$S of these strawberry species. It is worth noting that _F. chiloensis_ is the maternal parental of _F. × ananassa_ and the above-described differences might be derived from the genetic cross during the mating process between the parental strawberry species, i.e., _F. chiloensis × F. virginiana_ [1]. Alternatively, the lower dosage of H$_2$S required for decay control in _F. chiloensis_ could also be explained by its rusticity, as it is still a native undomesticated species that preserves its natural defensive strategies [4].

The physiological events during the ripening and senescence of strawberry fruit mainly involve softening associated cell wall modification [7]. The results reported in this work have shown that H$_2$S treatment delayed softening on fumigated _F. chiloensis_ fruit, which remained significantly firmer than untreated fruit throughout the monitored days (Figure 3A). Similar results in firmness preservation triggered by H$_2$S applications have been reported in strawberry ‘Bao Jiao’ and ‘Fengxiang’ cultivars [37,38], banana ‘Brazil’ [45,46] and kiwifruit ‘Jinkui’ [41,47]. Besides, in the present research, the respiration rate and titratable acidity were also delayed by the H$_2$S exposure on Chilean strawberry fruit, confirming the role of this molecule as an inhibitor of respiration rate as has been described in other non-climacteric fruits such as _F. × ananassa_ [37,38] and mulberry ( _Morus indica_ ‘Dianmian-1’) [42]. It has been previously reported that the high softening rate of Chilean strawberry fruit contributes to its fast postharvest decay [12]. Furthermore, the respiration rate is an important factor in determining the postharvest deterioration of strawberry fruit [37]. Therefore, a delayed softening and respiration rate, due to the H$_2$S treatment, influences an extended postharvest shelf-life by a decrease in the senescence and deterioration process.

Regarding changes in cell wall-associated polymers, the effect of H$_2$S treatment on cell wall polymer solubilization during the shelf-life of Chilean strawberry fruit was investigated. Pectin solubilization of H$_2$S-treated fruit was delayed compared to control fruit as loosely-bound pectin fraction (WSF) did not increase as much as in control fruit up to the fourth day, and consequently, ionically (CSF)- and covalently (NSF)-bound pectin fractions did not decrease as in control fruit (Figure 4A–D). All this evidence suggests that H$_2$S fumigation delays pectin degradation in Chilean strawberry fruit during its postharvest shelf-life period, which is related to the evidenced delay in softening rate in H$_2$S-treated strawberry fruit. Interestingly, differences between control and H$_2$S-treated fruit were also observed in hemicellulose-related fraction (KSF) in all days of treatment (Table S1), supporting the idea that H$_2$S could affect hemicellulose metabolism during...
postharvest storage. NaHS treatment has been recently observed altering the contents of cellulose and hemicellulose in alfalfa [48]. As far as we know, these results are the first report of the effect of H$_2$S treatment on the cell wall polymer contents in fleshy fruit. Therefore, the effect of H$_2$S on cell wall polymers especially hemicellulose and cellulose in fleshy fruit needs further characterization.

Plant cell wall disassembly during softening is a direct result of specific enzymatic activities, where pectin degradation plays a major role [10]. Previous reports have evidenced that decreased pectin depolymerization, by an antisense knockdown of a key pectinase gene (pectate lyase), leads to a reduction in the softening rate of strawberry fruit [13]. Thus, to gain molecular insights about how H$_2$S delays the cell wall polymer solubilization, the relative expression of key genes involved in pectin and hemicelluloses degradation during strawberry postharvest shelf-life was studied. The results evidenced that H$_2$S downregulates the expression of genes involved in pectin and hemicellulose metabolism. Genes encoding key enzymes involved in pectin solubilization such as polygalacturonase (FcPG1), pectate lyase (FcPL1), and an isoform of expansin (FcEXP2), a non-enzymatic protein associated with Chilean strawberry softening [49], exhibited a drastic and fast reduction in response to H$_2$S treatment (Figure 5A–C). It has been reported that the softening rate of the Chilean strawberry fruit reflects the expression of polygalacturonase and pectate lyase genes [12]. Additionally, the expression of expansin genes in strawberry varieties is highly related to contrasting fruit firmness [11,14]. Furthermore, a higher relationship factor for the FcPL1, FcEXP2, FcPG1 genes and the fruit firmness was observed (Figure 6A). Additionally, H$_2$S-treated strawberry fruit showed a strong positive correlation between firmness, the ionically- and covalently-bound pectin fractions (CSF and NSF), and genes involved in pectin solubilization (FcPG1, FcPL1, FcEXP2), exhibiting a higher significance than in control fruit (Figure 6C,D). Therefore, the effects displayed by H$_2$S treatment on decreasing the softening rate, pectin solubilization and the expression of genes involved in pectin depolymerization are highly associated with this previous evidence. Moreover, it has been reported that PG and PE activities are decreased in H$_2$S-treated strawberry fruits, prolonging their shelf-life [37,38]. Furthermore, the expression of a gene encoding for an isoform of xyloglucan endotransglycosylase/hydrolase (FcXTH1), involved in molecular modifications of hemicellulose, and a gene that encodes endo-$\beta$-1,4-glucanase (FcEG1), with cellulase activity, did not change its transcriptional level until the fourth shelf-life day of H$_2$S-treatment, exhibiting a significant increase and a strong reduction, respectively (Figure 5D,E). H$_2$S treatment could probably affect the synthesis of the hemicellulosic polymers by an unknown mechanism that needs further characterization.

4. Materials and Methods

4.1. Plant Material and Treatments

Ripe F. chiloensis fruit were harvested from a commercial orchard at Purén, the Araucanía Region, Chile (latitude 38°04’S; longitude 73°14’ W). The collected strawberries were immediately transported to the laboratory. Fruit of similar size and without external damage and microbial infection symptoms were selected. A total of 540 fruits were used for the following experiments.

Hydrogen sulfide (H$_2$S) fumigation was performed as described by Hu et al. [37] with some modifications in sealed chambers (5 L) using sodium hydrosulfide (NaHS) as an H$_2$S donor. Each chamber contained six perforated clamshells, which contained six strawberries, respectively. Initially, 200 mL of NaHS aqueous solutions at 0.2, 0.4, 0.8 and 1.2 mM were placed inside independent chambers to fumigate the fruit at 20 °C for up to 5 d. It is worth noting that the H$_2$S-donor (NaHS) and control (water) solutions (200 mL) and the chamber atmosphere were renewed each 24 h, by opening the lid of treatment and control chambers. For the control treatment, water was used instead of NaHS solution. Longer treatments (up to 14 d) were performed at 0.2 mM H$_2$S.
4.2. Evaluation of Decay Index

Decay index was determined as described by Ayala-Zavala et al. [50] and Hu et al. [37] with modifications. Fifty strawberry fruit per treatment were selected for the assessment of the decay index. Each fruit was classified into four ranks according to decay area: 0, no decay; 1, decay surface less than 10%; 2, decay surface between 10% and 30%; 3, decay surface between 30% and 50%; 4, decay surface more than 50%. The decay examination was recorded every day using the whole set of fruit per condition. The decay index was calculated by the following equation: decay index = \[ \sum (\text{rank} \times \text{fruit quantity per rank}) / \text{number of fruit} \times \text{higher rank} \] \times 100%. The experiment was repeated in two different harvest seasons.

4.3. Analysis of Fruit Chromaticity

The external color of individual strawberry fruit was analyzed with a colorimeter (model CR-200, Minolta), which measure L*, a*, and b* values, where L* indicates lightness, a* indicates chromaticity on a green (−) to red (+) axis, and b* indicates chromaticity on a blue (−) to yellow (+) axis. Two measurements on each equatorial side were performed per fruit. Values reported corresponded to the mean ± SE of three fruit per experimental condition. The initial fruit L*, a*, and b* values were a mean of 61.0, 6.1, and 17.0, respectively.

4.4. Analysis of Fruit Respiration Rate

CO₂ production rate (expressed in μmol CO₂ kg⁻¹ s⁻¹) was determined in three independent experiments by a CO₂ and O₂ analyzer (BRIDGE Analyzers, Inc., Bedford Heights, OH, USA). CO₂ percentage was measured after 1 h of incubation of three strawberry fruit in hermetic flasks. The initial fruit respiration rate had a mean of 1.0 μmol CO₂ kg⁻¹ s⁻¹.

4.5. Fruit Firmness Measurement

Firmness (N) was measured using the FirmTech II (BioWorks, Wamego, KS, USA) provided with a flat tip of 2 cm. Samples of six strawberry fruit were evaluated per each experimental condition. Two measurements on each equatorial side were performed per fruit with a penetration depth of 1 mm. The values reported corresponds to the mean ± SE of six fruit per condition. After these analyses, the peduncle and calyx of each fruit were removed, and the fruit was cut into pieces, frozen under liquid nitrogen and stored at −80 °C for further determinations. The fruit from each experimental condition was mixed to provide a bulk of fruit samples. The initial fruit firmness had a mean of 1.7 N.

4.6. Determination of Soluble Solids, pH and Titratable Acidity

Two grams of frozen fruit tissue were homogenized in water with a disperser T25 digital Ultra-turrax® (IKA, Staufen, Germany) and adjusted to 25 mL final volume. The mixture was filtrated through miracloth, and the juice was analyzed for soluble solids content (SSC), pH, and titratable acidity (TA). SSC (expressed as %) was measured at 20 °C using a hand-held temperature compensated refractometer (Atago Co., Tokyo, Japan). TA (expressed as m Eq 0.1 kg⁻¹ of fresh weight (FW)) was determined by titration using a pH meter and automatic titrator PH-Burette 24 (Crisons, Barcelona, Spain) of an aliquot of 5 mL of strawberry juice with 20 mM NaOH until reaching pH 8.2. The pH of the juice was recorded per each replicate. Three independent fruit extractions were prepared from each biological sample, and values correspond to mean ± SE.

4.7. Cell Wall Extraction and Cell Wall Fractionation

Cell wall material was extracted according to Figueroa et al. [44] with some modifications. Five grams of ground frozen fruit tissue were homogenized in 40 mL of 95 % ethanol and boiled for 45 min. The insoluble material was filtered through miracloth
and sequentially washed with 15 mL of boiling ethanol, 15 mL of chloroform/methanol (1:1, \(v/v\)) and 15 mL of acetone. The residue (Alcohol Insoluble Residue, AIR) was dried overnight at 37 °C and weighed. The results of three replicates per treatment were expressed as mg AIR per g\(^1\) FW.

The fractionation of cell wall material was performed using a sequential chemical treatment of AIR as previously described [44]. The water-soluble, the 50 mM trans-1,2-diaminocyclohexane-N,N,N',N''-tetraacetic acid (CDTA)-soluble, the 50 mM NaCO\(_3\)-soluble, and the 4 M KOH-soluble fractions were obtained and named as WSF, CSF, NSF, and KSF fractions, respectively. Two independent extractions were obtained from each experimental replicate. The concentration of uronic acid (UA) and neutral sugars (NS) in the different cell wall fractions were determined colorimetrically as previously described [51,52]. The results were calculated using standard curves prepared for galacturonic acid and glucose for UA and NS, respectively. Measurements were performed in triplicate, and the results were expressed as mg of galacturonic acid (UA) or glucose (NS) per g of AIR.

4.8. Analysis of Gene Expression

Total RNA was isolated from 2 g of frozen fruit samples using a modified CTAB method [53]. Three independent RNA extractions were prepared from each biological sample. One microgram of total RNA was treated with DNase (Turbo DNA-free kit, Ambion®, Life Technologies) to eliminate genomic DNA. The RNA quantity and purity were estimated at 260/280 nm by NanoDrop™ 1000 Spectrophotometer (Thermo Scientific, Waltham, MA, USA). The RNA integrity was visualized in electrophoresis in 1.5 % (w/v) agarose gel stained with GelRedTM (Biotium, Hayward, CA, USA). The cDNA was synthesized with a first-strand cDNA synthesis kit (Thermo Scientific, Waltham, MA, USA). Quantitative reverse transcription PCR (RT-qPCR) was performed in a Stratagene MX3000P (Agilent Technologies, Santa Clara, CA, USA). Each reaction consisted of 20 μL containing 2 μL of (1:10 diluted) cDNA, 1 μL of primer mix 10 μM, 10 μL of Sybr Green PCR Master Mix 2X (Stratagene®, Agilent Technologies), and nuclease-free water to reach the final reaction volume. The cycling conditions were 1 cycle of denaturation at 95 °C for 5 min, followed by 40 cycles (two-segment) of amplification (95 °C/15 s, 60 °C/45 s) and a final melting cycle (95 °C/1 min, 55 °C/30 s and 95 °C/30 s). Each RT-qPCR reaction was performed in three technical replicates and the mean was used for further analysis. A control without template was included in each RT-qPCR run. Fluorescence was measured at the end of each extension step. The specific primer sequences for the genes encoding polygalacturonase 1 (FcPG1), pectate lyase (FcPL), endo-\(\beta\)-1,4-glucanase 1 (FcEG1), expansin 2 (FcEXP2), xyloglucan endotransglycosylase/hydrolase 1 (FcXTH1), and glyceraldehyde 3-phosphate dehydrogenase (FcGAPDH) from F. chiloensis were obtained from a previous report [44]. The relative expression levels were normalized by \(2^{-\Delta \Delta C T}\) method [54] using GAPDH as the reference gene. The results were expressed in arbitrary units assigning the value of one unit to time zero. The data were analyzed by Tukey test \((p < 0.05)\) per time. Each value represents mean ± SD \((n = 4)\).

4.9. Statistical Analysis

The results were compared by two-way analysis of variance (ANOVA) (time, treatment) and Duncan’s multiple range test at the 5 % level of significance. Shapiro–Wilk and Levene’s tests were used to verify normality and homoscedasticity, respectively. Principal component analysis and a Pearson correlation analysis were performed to evaluate the differences between the treatments (control and treated with H\(_2\)S) using R 1.3.1093. The data were presented with their means and standard errors.

5. Conclusions

In this work, we demonstrated a novel role of H\(_2\)S as a gasotransmitter prolonging the postharvest shelf-life of the Chilean strawberry fruit by preventing its decay and its
fast softening rate through a decreased pectin degradation, which reflects the downregulation effect on the expression of key pectinase-related genes. These pieces of evidence provide useful information about the biological function of H2S acting as a plant gasotransmitter and transforms the Chilean strawberry fruit as an emergent model for studying the quick softening rate and decay in non-climacteric fruit. In addition, this biological system could provide, in the future, valuable information about the role of H2S as a cell wall modifier in plants, adding new insights into the regulation of postharvest physiology of fruit and vegetables during storage. Furthermore, the use of this elicitor emerges as a potent tool for the exogenous application of horticultural products for storage and shelf-life preservation, albeit the approval for use of H2S gas on fresh foods is still pending at the global level.

Supplementary Materials: The following are available online at www.mdpi.com/article/10.3390/ijms221810008/s1, Table S1: Changes in total cell wall material (AIR) content and pectins and hemicellulose fractions during the shelf-life period of Chilean strawberry fruit subjected to H2S treatment.

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