RNA-seq analysis reveals different transcriptomic responses to GA3 in early- and mid-season varieties before ripening initiation in sweet cherry (Prunus avium L.) fruits

Nathalie Kuhn  
Pontifical Catholic University of Valparaiso

Jonathan Maldonado  
University of Chile

Claudio Ponce  
University of Chile

Macarena Arellano  
University of Chile

Alson Time  
University of Chile

Salvatore Multari  
Fondazione Edmund Mach

Stefan Martens  
Fondazione Edmund Mach

Esther Carrera  
Universitat Politècnica de València

José Manuel Donoso  
Instituto de Investigaciones Agropecuarias

Boris Sagredo  
Instituto de Investigaciones Agropecuarias

Lee Ann Meisel (meisel@inta.uchile.cl)  
University of Chile

Research Article

Keywords: Gibberellin, GA3, IAD, non-climacteric, ripening, RNA-seq, sweet cherry, transcriptome

DOI: https://doi.org/10.21203/rs.3.rs-189281/v1

License: © This work is licensed under a Creative Commons Attribution 4.0 International License. Read Full License
Abstract

Gibberellin (GA) negatively affects color evolution and other ripening-related processes in non-climacteric fruits. The bioactive GA, gibberellic acid (GA$_3$), is commonly applied at the light green-to-straw yellow transition to increase firmness and delay ripening in sweet cherry (*Prunus avium* L.), though causing different effects depending on the variety. Recently, we reported that GA$_3$ delayed the IAD parameter (a ripening index) in a mid-season variety, whereas GA$_3$ did not delay IAD but reduced it at ripeness in an early-season variety. To further explore this contrasting behavior between varieties, we analyzed the transcriptomic responses to GA$_3$ applied on two sweet cherry varieties with different maturity time phenotypes. At harvest, GA$_3$ produced fruits with less color in both varieties. Similar to our previous report, GA$_3$ delayed fruit color initiation and IAD only in the mid-season variety, and reduced IAD at harvest only in the early-season variety. RNA-seq analysis of control- and GA$_3$-treated fruits of both varieties revealed that ripening related categories were overrepresented in the early-season variety, including 'photosynthesis' and 'auxin response.' GA$_3$ also changed the expression of carotenoid and abscisic acid (ABA) biosynthetic genes in this variety. In contrast, overrepresented categories in the mid-season variety were related mainly to metabolic processes. In this variety, some *PP2Cs* putative genes were positively regulated by GA$_3$, which are negative regulators of ABA responses. This was accompanied with downregulation of *MYB44*-like genes, putative repressors of *PP2Cs* expression. These results show that GA$_3$ differentially modulates the transcriptome at the onset of ripening in a variety-dependent manner and suggest that GA$_3$ impairs ripening through the modification of ripening associated gene expression only in the early-season variety; whereas in the mid-season variety, possibly a control of the ripening timing occurs through the *PP2C* gene expression regulation. This work contributes to the understanding of the role of GA in non-climacteric fruit ripening.

Introduction

Fruit ripening is a complex process that involves changes in the cell wall composition, accumulation of sugars and pigments in the peel (exocarp), cell enlargement, and the decrease in organic acid content. In non-climacteric fruits, which do not depend on an ethylene rise for ripening initiation, many hormones control this process, with abscisic acid (ABA) being the most important$^{1,2}$. Exogenous application of ABA triggers the ripening process in non-climacteric species such as grapevine$^3$ (*Vitis vinifera*), strawberry$^2$ (*Fragaria* spp.) and sweet cherry$^4$ (*Prunus avium*). Other hormones are involved in this process, but their participation is poorly characterized. ABA's most characteristic feature is its sharp increase a few days before the anthocyanin accumulation$^{3,5}$. However, ABA is not the only hormone that is present during this period. In sweet cherry, auxin, gibberellin (GAs), and cytokinins are present$^5$. These hormones may modulate the ripening process together with ABA. For example, underripe grapevine berries have higher seed content and higher auxin levels in *veraison* bunches$^6$, and NAA (α-naphthalene acetic acid) application delays ripening in this non-climacteric species$^7$. In sweet cherry, indole-3-acetic acid (IAA) downregulates the ABA signaling pathway genes$^8$.

Regarding GA, mainly GA$_4$ is detected after the grapevine berry set$^9$. In strawberry, GA$_4$ decreases from the white to the red stage of the fruits$^{10}$. In sweet cherry, GA$_4$ significantly and inversely correlates with ripening parameters$^5$. Taking these results together, GA possibly exerts a negative role in the ripening process. At the molecular level, at *veraison*, some GAs oxidase genes change their expression compared to the previous stage in grapevine fruits$^{11}$. GA$_3$ treatment at *veraison* delays ripening and affects the transcript levels of putative *PP2Cs*, which are negative regulators of ABA signaling$^{12}$. In sweet cherry, GA$_3$ application delays ABA accumulation and reduces anthocyanin levels$^{13}$. 
Additionally, GA$_3$ delays fruit size increase, ripening, and color development when applied at the degreening stage$^{14,15,16}$. Choi et al.$^{14}$ reported that the mid-season varieties had a delayed fruit size increase and polygalacturonase and Cx-cellulase activities after the GA$_3$ treatment, whereas GA$_3$ did not affect fruit growth in the early-season varieties. Kuhn et al.$^{17}$ reported that the Index of Absorbance Difference (IAD), a ripening index, was delayed after the GA$_3$ treatment only in a mid-season variety, whereas the treatment reduced the IAD only at harvest in an early-season variety; interestingly, both varieties presented modified expression of ABA pathway related genes. Despite the available information on the participation of GA in the fruit ripening process, the effect of this hormone has been barely explored at the whole gene expression level during the sweet cherry fruit ripening.

There are a few transcriptomic analyses in sweet cherry. These works explore the transcriptomic features underlying fruit development$^{18}$, fruit coloring$^{19}$, and light-dependent anthocyanin accumulation in sweet cherry fruits$^{20}$. Guo and co-workers$^{20}$ found that some ABA and GA pathway genes co-expressed with light-regulated genes and suggested that both hormones may play crucial roles in light-dependent anthocyanin biosynthesis. Therefore, further exploration of the interplay of GA with ABA and other hormones during the ripening process of non-climacteric fruits is required. In grapevine, some transcriptomic studies of the effect of GA$_3$ treatment during the initial fruit development have been performed$^{21}$. In loquat (Eriobotrya japonica), GA$_3$ applied at bloom changed the expression of ABA and auxin biosynthesis-related genes$^{22}$. Finally, RNA-seq performed in Chinese sour cherry (Cerasus pseudocerasus) identified transcriptomic changes in response to GA$_3$-induced parthenocarpy$^{23}$. These studies focus on initial fruit development in which the GA participation is quite clear; however, the GAs role in the ripening process at the transcriptomic level has not been addressed to date.

Here we analyze the effect of GA$_3$ at the physiological and global gene expression level in early- and mid-season sweet cherry varieties. This work aims to contribute to the understanding of the participation of GA in the fruit ripening process in non-climacteric fruits.

**Results**

‘Celeste’ and ‘Bing’ are described as early- and mid-season varieties$^{24}$. Both varieties were characterized for fruit growth and phenology in two seasons (Fig. 1A and Fig. S1A and Tables S1 to S4). In the 2017–2018 season, the slow growth period was more prolonged in the mid-season variety, in which growth resumption occurred from 39 DAFB (Fig. 1A). On the other hand, pink color initiation started at 40 DAFB in the early-season variety (Fig. 1B), whereas in the mid-season variety, it started at 50 DAFB (Fig. 1A and Table S2). GA$_4$ and GA$_1$ were detected in the fruits of both varieties from 34 to 44 DAFB (Fig. S1B).

Several GAs are present at the time of fruit color initiation in sweet cherry and other non-climacteric fruits$^{5,10}$; therefore, we aimed to investigate the role of GA in ripening triggering by altering GA homeostasis. For this, GA$_3$ treatment was performed when both varieties transitioned from light green to straw yellow fruit color (35 DAFB the 2017–2018 season and 34 DAFB the 2018–2019 season). At the time of GA$_3$ application, both varieties were phenologically similar (Fig. 1; Tables S1 to S4).

GA$_3$ affected ripening parameters at fully ripeness in both varieties, in the 2017–2018 seasons (Table 1), including firmness in both varieties, early-season variety fruit weight, and mid-season variety SSC. Similar results were observed in the 2018–2019 season (Table S5).
Table 1
Ripening related parameters in control- and GA$_3$-treated fruit samples at harvest in early- and mid-season varieties, during 2017–2018 season. d.u., durometer units; SSC, soluble solids content; M.A., malic acid.

| Season     | Variety                  | Treatment | Weight (g) | Firmness (d.u.) | SSC (ºBrix) | Acidity (M.A. %) |
|------------|--------------------------|-----------|------------|-----------------|-------------|------------------|
| 2017–2018  | Early-season var., Celeste | Control   | 9.03$^{a**}$ | 55.63$^a$       | 15.19$^a$  | 2.12$^a$         |
|            |                          | GA$_3^*$  | 9.71$^b$   | 58.33$^b$       | 15.26$^a$  | 1.92$^a$         |
|            | Mid-season var., Bing    | Control   | 8.70$^a$   | 54.06$^a$       | 19.85$^a$  | 3.24$^a$         |
|            |                          | GA$_3$    | 8.61$^a$   | 63.21$^b$       | 18.91$^b$  | 3.19$^a$         |

* GA$_3$ was applied as the commercial product ProGibb ® 40% SG to individual branches a rate of 30 ppm. GA$_3$ treatment was at the light green-to-straw yellow transition of fruits at 35 DAFB in Celeste and Bing.

**For each ripening related parameter, the significance of variation between control- and GA$_3$-treated fruits was tested by one-way ANOVA analysis with Tukey's post hoc test, whereby different letters are significantly different means ($p < 0.05$).

GA$_3$ treatment produced less color in both varieties at harvest, decreased the percentage of dark fruits, and increased the percentage of light fruits in the 2017–2018 season (Figs. 2A-2D), which also occurred the following season (Figs. S2A-S2D). Estimated anthocyanins were significantly reduced in both varieties at harvest (Figs. 2E and 2F).

To further characterize the effect of GA$_3$, IAD was measured from the date of the GA$_3$ treatment in both varieties (Fig. 3).

IAD is a non-destructive ripening index utilized in sweet cherry, correlated with the anthocyanin content$.^{25}$ It reveals the presence of phenolic compounds that act as a screen for chlorophyll absorbance by measuring absorbance differences between 560 nm, 640 nm, and the 750 nm reference$.^{26}$ After the GA$_3$ treatment, the early-season variety had a lower IAD value only at harvest in both seasons (Fig. 3A and Fig. S3A), whereas the mid-season variety had a delayed increase in the IAD values (Fig. 3B and Fig. S3B). This delay observed in the mid-season variety was accompanied by color differences between control- and GA$_3$-treated fruits as soon as 15 days after the treatment (50 DAFB), whereas control- and GA$_3$-treated fruits of the early-season variety had no differences in color at the same date (Fig. 3C).

As both varieties were differentially affected by GA$_3$, we explored the transcriptomic effect produced by GA$_3$ treatment through RNA-seq analysis. For this, we analyzed fruit samples collected minutes before the GA$_3$ treatment in the 2017–2018 season (T0, 35 DAFB), and control- and GA$_3$-treated fruit samples collected four days after the GA$_3$ treatment (CT4 and GT4, respectively, 39 DAFB). In total, 18 samples were sequenced, nine per variety (three points: T0, CT4 and GT4, and three replicates) that yielded a total of 859 million filtered reads (84 Gb of data), with an average length of 98 bp, and a similar quantity of mapped reads to the $P$. avium transcriptome reference that was ca. 68% of the total filtered read count (Table S6).

Clustering of the 100 genes having the highest expression levels revealed that samples behaved as expected, i.e., replicates grouped, except for one replicate of T0 of early-season variety (Sample_ID: CT0_3) that was excluded from subsequent analysis (Fig. S4).
DEGs analysis was performed on the genes that changed between T0 and CT4 (CT4-T0) and between T0 and GT4 (GT4-T0) with a cutoff value of two-fold change and FDR < 0.05. Up and downregulated genes were identified in both varieties (Fig. 4). In the early-season variety, 381 and 808 genes changed only in GT4-T0 and CT4-T0, respectively (Fig. 4A). In mid-season variety, 249 and 495 genes changed only in GT4-T0 and CT4-T0, respectively (Fig. 4B). There were 482 and 729 genes in the early-season variety, and 563 and 314 genes in the mid-season variety, that changed independently of the GA$_3$ treatment (they are in the intercepts of the diagrams). In contrast, the 381 GT4-T0 genes plus the 808 CT4-T0 genes in the early-season variety, and the 249 GT4-T0 genes plus the 495 CT4-T0 genes in the mid-season variety, were GA$_3$-modulated. Therefore, they were used in the subsequent GO enrichment analysis focused on the Biological Process categories.

In the early-season variety, 'response to stimulus', 'response to hormone' and 'response to auxin' were some of the overrepresented GO terms (FDR < 0.05) in the GT4-T0 comparison. On the contrary, 'protein phosphorylation' and 'photosynthesis' were some of the overrepresented GO terms (FDR < 0.05) in the CT4-T0 comparison (Table 2).

Table 2
Gene ontology (GO) enrichment analysis performed on the DEGs that are modulated by GA$_3$ in early- and mid-season varieties. The biological process function ontologies with the lowest $p$-value and FDR less than 0.05 are shown.

| Variety                        | Comparisons       | GO ID       | GO description                | $p$-value  |
|-------------------------------|-------------------|-------------|--------------------------------|------------|
| Early-season var., Celeste    | GT4-T0 total (381)| GO:0050896  | response to stimulus          | 2.284E-06  |
|                               |                   | GO:0042221  | response to chemical          | 6.317E-06  |
|                               |                   | GO:0010033  | response to organic substance | 1.107E-05  |
|                               |                   | GO:0009725  | response to hormone           | 1.519E-05  |
|                               |                   | GO:0009719  | response to endogenous stimulus| 1.647E-05 |
|                               |                   | GO:0009733  | response to auxin             | 2.071E-05  |
|                               | CT4-T0 total (808)| GO:0055114  | oxidation-reduction process   | 9.47E-08   |
|                               |                   | GO:006468   | protein phosphorylation       | 1.62E-07   |
|                               |                   | GO:0008152  | metabolic process             | 2.65E-07   |
|                               |                   | GO:0016310  | phosphorylation               | 2.09E-05   |
|                               |                   | GO:0019419  | sulfate reduction             | 2.57E-05   |
|                               |                   | GO:0015979  | photosynthesis                | 4.86E-05   |
| Mid-season var., Bing         | GT4-T0 up* (168)  | GO:0005975  | carbohydrate metabolic process| 5.293E-04 |
|                               |                   | GO:0055114  | oxidation-reduction process   | 3.986E-08  |
|                               |                   | GO:009607   | response to biotic stimulus   | 3.269E-05  |
|                               |                   | GO:0005975  | carbohydrate metabolic process| 4.311E-05 |
|                               |                   | GO:0006950  | response to stress            | 4.565E-05  |
|                               | CT4-T0 down* (306)|             |                                |            |

*GT4-T0 up and CT4-T0 down comparisons of mid-season variety had more overrepresented categories than GT4-T0 total and CT4-T0 total, respectively; therefore, they were included as these comparisons are more informative.
Regarding the mid-season variety, the GO term 'carbohydrate metabolic process' was overrepresented (FDR < 0.05) in the GT4-T0 comparison with upregulated genes, whereas 'oxidation-reduction process' and 'response to stress', among others, were overrepresented GO terms (FDR < 0.05) in the CT4-T0 comparison that includes downregulated genes (Table 2).

Figure 5. Heat map with the number of DEGs that are modulated by GA3 in each Gene Ontology (GO) category. A, Number of genes in GT4-T0 and CT4-T0 comparisons that are up or downregulated in the early-season variety. B, Number of genes in GT4-T0 and CT4-T0 comparisons that are up or downregulated in the mid-season variety. For A and B, biological processes with $p < 0.001$.

In general, GO terms were specific to a particular comparison, except for the GO category 'oxidation-reduction process' in the early-season variety (Fig. 5A) and 'carbohydrate metabolic process' in the mid-season variety (Fig. 5B), with downregulated genes in CT4-T0 and upregulated genes in the GT4-T0. The GO term 'metabolic process' had the highest number of DEGs modulated by GA3 in the mid-season variety (Fig. 5B). In the early-season variety, this GO term and 'cellular process' were the most abundant (Fig. 5A).

CT4-T0 up and down comparisons contain GA-modulated genes that up or downregulated from T0 to T4, but they no longer changed after the GA3 treatment. In the early-season variety, the CT4-T0 down comparison contained DEGs in 'photosynthesis', 'protein phosphorylation', and 'transmembrane transport' GO categories, among others (Fig. 5A). In contrast, in the mid-season variety, it contained DEGs in the 'response to stress' and 'plant-type secondary cell wall biogenesis' GO categories, among others (Fig. 5B). On the other hand, the 'carotenoid metabolism' GO category contained GA-modulated genes that upregulated from T0 to T4 in the early-season variety.

GT4-T0 up and down comparisons contain GA-modulated genes that did not change from T0 to T4 but are up or downregulated by GA3. DEGs in the 'auxin response' and 'putrescine biosynthetic process' GO categories were present in the GT4-T0 up comparison, in the early-season variety. In contrast, several GO categories related to negative regulation of gene expression were found in the GT4-T0 down comparison in this variety (Fig. 5A). In the mid-season variety, DEGs in the 'cell wall organization or biogenesis' and 'reactive oxygen species metabolic process' GO categories were present in the comparison containing upregulated genes (GT4-T0 up), whereas 'xylan biosynthetic process' had DEGs in the GT4-T0 down comparison in this variety (Fig. 5A).

After that, we investigated the variety-specific genes that respond to GA3 treatment (Fig. 6). In the GT4-T0 comparison of the early-season variety, 436 and 652 genes were up and downregulated, respectively. On the other hand, 477 and 145 genes were up and downregulated in the mid-season variety.

As shown in Table 3, the GO terms 'response to auxin', 'response to stress', and 'DNA packaging', among others, were overrepresented (FDR < 0.01), with genes that exclusively changed in the early-season variety. On the other hand, in the mid-season variety, 'regulation of transcription, DNA-templated' and 'regulation of nucleic acid-templated transcription' were some of the overrepresented GO terms (FDR < 0.01) in the upregulated DEGs that were exclusive of this variety.
Table 3
GO enrichment analysis performed on the DEGs that are modulated by GA$_3$ and unique to each variety. The biological process function ontologies with the lowest $p$-value and FDR less than 0.05 are shown.

| Variety                      | Comparisons       | GO ID             | GO description                              | $p$-value |
|------------------------------|-------------------|-------------------|---------------------------------------------|-----------|
| Early-season var., Celeste   | GT4-T0 total      | GO:0050896        | response to stimulus                        | 1.34E-13  |
| (1088)                       |                   | GO:0055114        | oxidation-reduction process                 | 1.31E-11  |
|                              |                   | GO:0042221        | response to chemical                        | 2.90E-10  |
|                              |                   | GO:0009733        | response to auxin                           | 7.08E-10  |
|                              |                   | GO:0006950        | response to stress                          | 6.87E-09  |
|                              |                   | GO:0006323        | DNA packaging                               | 9.67E-09  |
| Mid-season var., Bing        | GT4-T0 total      | GO:0006355        | regulation of transcription, DNA-templated  | 2.90E-06  |
| (622)                        |                   | GO:1903506        | regulation of nucleic acid-templated         | 3.48E-06  |
|                              |                   | GO:2001141        | regulation of RNA biosynthetic process       | 3.48E-06  |
|                              |                   | GO:0009409        | response to cold                            | 4.13E-06  |
|                              |                   | GO:2000112        | regulation of cellular macromolecule         | 5.81E-06  |
|                              |                   | GO:0051252        | regulation of RNA metabolic process          | 5.94E-06  |

In Fig. 7, the enriched GO terms that had more DEGs unique to early-season variety were 'response to stimulus', 'oxidation-reduction process', 'cellular component organization or biogenesis', 'chromosome organization', 'cell cycle', 'response to hormone', 'response to auxin', among others. In contrast, the GO terms that had more DEGs unique to mid-season variety were 'biological regulation', 'aromatic compound biosynthetic process', 'regulation of transcription, DNA-templated', and 'regulation of nucleic acid-templated transcription', among others (Fig. 7).

As 'response to hormone' was an overrepresented GO, a directed search of hormone-related genes was performed in the CT4-T0 and GT4-T0 comparisons of each variety (Table 4).
Table 4
Differential gene expression as fold change of genes related to hormone pathways in pairwise comparisons CT4-T0 and GT4-T0 of each variety. In gray, comparisons having genes whose absolute fold change value was at least two.

| Name                                      | ID                              | GO Biological Process | CT4-T0 | GT4-T0 | CT4-T0 | GT4-T0 |
|-------------------------------------------|---------------------------------|-----------------------|--------|--------|--------|--------|
| gibberellin-regulated protein 11-like     | Pav_sc0001364.1_g150.1.mk       | Not identified        | -19.5  | -8.4   | -9.4   |        |
| gibberellin-regulated protein 1-like      | Pav_sc0000044.1_g560.1.mk       | Not identified        |        |        |        | 2.2    |
| gibberellin-regulated protein 4-like      | Pav_sc0000766.1_g050.1.mk       | Not identified        |        |        |        | 2.9    |
| gibberellin-regulated protein 9           | Pav_sc0000350.1_g1060.1.mk      | Not identified        | -5.8   | -3.8   |        |        |
| gibberellin 2-beta-dioxygenase-like       | Pav_sc0003135.1_g610.1.mk       | Not identified        | -5.1   |        |        |        |
| gibberellin 2-beta-dioxygenase 1          | Pav_sc0000143.1_g470.1.mk       | Not identified        | -4.2   | -2.0   |        |        |
| gibberellin 2-beta-dioxygenase 1-like     | Pav_sc0000095.1_g1110.1.mk      | Not identified        | -2.6   |        |        |        |
| DELLA protein GAI-like                    | Pav_sc0000704.1_g110.1.mk       | Not identified        |        |        |        | 2.2    |
| gibberellin receptor GID1B-like           | Pav_sc0000848.1_g080.1.mk       | Response to endogenous stimulus |        |        |        | -3.1   |
| phytoene synthase 2 chloroplastic         | Pav_sc0001124.1_g480.1.mk       | Metabolic proces      | 2.3    |        |        |        |
| 9-cis epoxycarotenoid dioxygenase NCED1 chloroplastic-like | Pav_sc000095.1_g1080.1.mk | Metabolic proces | 4.0    |        |        |        |
| probable carotenoid cleavage dioxygenase 4 chloroplastic | Pav_sc0000354.1_g230.1.mk | Metabolic proces | -4.2   | -2.1   | -2.9   | -3.9   |
| abscisic acid 8'-hydroxylase 2 isoform X1 | Pav_sc0000563.1_g270.1.mk       | Metabolic proces      | -4.9   |        |        |        |
| abscisic acid 8'-hydroxylase 4            | Pav_sc0002234.1_g030.1.mk       | Metabolic proces      | -5.1   | -2.5   |        |        |
| abscisic stress ripening                  | Pav_sc0005261.1_g020.1.br       | Not identified        | 2.8    | 2.3    |        |        |
| abscisic stress-ripening protein 2        | Pav_sc0002659.1_g070.1.br       | Not identified        | 2.9    |        |        |        |
| Name                                           | ID                                      | GO Biological Process                  | Early-season variety | Mid-season variety |
|------------------------------------------------|-----------------------------------------|----------------------------------------|----------------------|--------------------|
| abscisic acid stress ripening protein homolog  | Pav_sc0000863.1_g110.1.m                | Not identified GO                      | 3.1                  | 2.8                |
| abscisic stress-ripening protein 2            | Pav_sc0000863.1_g060.1.br               | Not identified GO                      | 3.3                  | 2.5                |
| abscisic acid receptor PYL4                    | Pav_sc0001341.1_g250.1.mk               | Not identified GO                      | 2.1                  | 2.6 2.6            |
| protein C2-DOMAIN ABA-RELATED 4-like          | Pav_sc0000464.1_g320.1.mk               | Not identified GO                      | 2.2                  | 2.0                |
| protein C2-DOMAIN ABA-RELATED 7               | Pav_sc0000221.1_g240.1.mk               | Not identified GO                      | 2.1                  | 2.2                |
| probable protein phosphatase 2C 12            | Pav_sc0000852.1_g1120.1.mk              | Cellular protein modification process   | 2.1                  | 2.9 3.3            |
| probable protein phosphatase 2C 39            | Pav_sc0000975.1_g200.1.mk               | Cellular protein modification process   | .                    | 2.2                |
| probable phosphatase 2C-like protein 44       | Pav_sc0000119.1_g610.1.mk               | Cellular protein modification process   | .                    | -3.0 -4.4          |
| auxin-responsive protein SAUR50               | Pav_sc0000480.1_g730.1.mk               | Response to endogenous stimulus        | 10.8                 | 13.8 6.8 4.7       |
| auxin-responsive protein IAA29                | Pav_co4052989.1_g010.1.mk               | Response to endogenous stimulus        | 2.8                  | 3.5                |
| auxin-responsive protein IAA29                | Pav_sc0000311.1_g1160.1.mk              | Response to endogenous stimulus        | 5.9                  | 5.3                |
| auxin-responsive protein IAA13                | Pav_sc0002717.1_g060.1.br              | Response to endogenous stimulus        | 2.6                  |                    |
| auxin-responsive protein IAA11 isoform X1     | Pav_sc0002181.1_g160.1.mk               | Response to endogenous stimulus        | 2.6                  |                    |
| auxin-responsive protein IAA8-like isoform X2 | Pav_sc0002327.1_g560.1.mk               | Response to endogenous stimulus        | 4.3                  | 3.4 2.9            |
| auxin-induced protein IAA6                    | Pav_sc0000716.1_g200.1.mk               | Response to endogenous stimulus        | 2.3                  | 2.0 2.3 2.5        |
| protein kinase PINOID                         | Pav_sc0000481.1_g260.1.mk               | Response to endogenous stimulus        | 3.4                  | 4.3 4.4 6.5        |
| Name                                                                 | ID                                          | GO Biological Process                                      | CT4_T0 | GT4_T0 | CT4_T0 | GT4_T0 |
|----------------------------------------------------------------------|---------------------------------------------|------------------------------------------------------------|--------|--------|--------|--------|
| probable indole-3-pyruvate monooxygenase YUCCA10                      | Pav_sc0000382.1_g010.1.mk                   | Response to endogenous stimulus                           | 6.2    | 3.0    | 4.2    |        |
| auxin-induced protein 15A                                            | Pav_sc0006480.1_g020.1.br                  | Response to endogenous stimulus                           | 7.3    | 10.6   |        |        |
| auxin-induced protein 15A                                            | Pav_sc0006480.1_g030.1.br                  | Response to endogenous stimulus                           | 3.9    | 5.4    |        |        |
| auxin-induced protein 15A-like                                       | Pav_sc0007626.1_g030.1.br                  | Response to endogenous stimulus                           | 10.8   | 15.0   |        |        |
| auxin-induced protein 15A-like                                       | Pav_sc0008108.1_g010.1.mk                   | Response to endogenous stimulus                           | 5.1    | 8.4    |        |        |
| auxin-induced protein 15A-like                                       | Pav_sc0004469.1_g010.1.br                  | Response to endogenous stimulus                           |        | -2.7   | -3.0   |        |
| auxin-induced protein 15A-like                                       | Pav_sc0004655.1_g010.1.br                  | Response to endogenous stimulus                           |        | -3.7   | -3.0   |        |
| auxin-induced protein 15A-like                                       | Pav_sc0004655.1_g020.1.br                  | Response to endogenous stimulus                           |        | -3.4   |        |        |

Name | ID                                              | GO Biological Process                                      | CT4_T0 | GT4_T0 | CT4_T0 | GT4_T0 |

1-aminocyclopropane-1-carboxylate oxidase | Pav_sc0001084.1_g100.1.mk | Metabolic process | 3.0 | 2.8 |
1-aminocyclopropane-1-carboxylate oxidase | Pav_sc0002206.1_g340.1.mk | Metabolic process | 6.6 | 5.3 |
1-aminocyclopropane-1-carboxylate oxidase homolog 1-like | Pav_sc0000583.1_g270.1.mk | Metabolic process | -12.5 |
ethylene-responsive transcription factor ERF003-like | Pav_sc0000890.1_g720.1.mk | Nucleobase-containing compound metabolic process | 7.0 | 5.2 | 22.3 | 25.3 |
ethylene-responsive transcription factor 1B-like | Pav_sc0000583.1_g510.1.mk | Nucleobase-containing compound metabolic process | -7.8 | -6.0 | 12.6 |
ethylene-responsive transcription factor 12 | Pav_sc0001305.1_g990.1.mk | Nucleobase-containing compound metabolic process | 3.0 | 3.0 | 5.9 | 6.3 |
Several gibberellin 2-beta-dioxygenase related genes were downregulated in the CT4-T0 and GT4-T0 comparisons only in the early-season variety. Regarding the ABA biosynthetic pathway, two abscisic acid 8'-hydroxylase related genes were downregulated in the early-season variety. Concerning the ABA response, several abscisic stress ripening-related genes were upregulated in the CT4-T0 comparison only in the early-season variety, which was less marked in the GT4-T0 comparison. Regarding the auxin response, several genes that encode putative auxin AuxIAA response repressors were induced in the CT4-T0 comparison of the early-season variety, and they were less upregulated in the GT4-T0 comparison. Some genes coding for auxin-induced protein 15A and 15A-like were upregulated in CT4-T0 and were upregulated even more in GT4-T0, whereas other groups of auxin-induced protein 15A-like were downregulated in the mid-season variety. In contrast, several ethylene-responsive transcription factor genes were more upregulated in the mid-season variety than in the early-season variety.

Given that 'negative regulation of gene expression' and 'DNA packaging' and 'nucleosome assembly' GO terms were overrepresented, a directed search of genes related to these categories was performed (Table 5).
Table 5
Differential gene expression as fold change of genes related to transcriptional and epigenetic regulation in pairwise comparisons CT4-T0 and GT4-T0 of each variety. In gray, comparison having genes whose absolute fold change value was at least two.

| Name                                           | ID                                      | GO Biological Process                                      | Early-season variety | Mid-season variety |
|------------------------------------------------|-----------------------------------------|------------------------------------------------------------|----------------------|--------------------|
| dof zinc finger protein DOF1.2                  | Pav_sc0001556.1_g080.1.mk               | Nucleobase-containing compound metabolic process           | 3.1                  | 3.3                |
| transcription factor MYB44                     | Pav_sc0000625.1_g100.1.mk               | Not identified GO                                           | 3.8                  | 3.9                |
| transcription factor MYB44-like                | Pav_sc0000800.1_g060.1.mk               | Not identified GO                                           | .                    | .                  |
| transcription factor MYB44-like                | Pav_sc0000852.1_g360.1.mk               | Not identified GO                                           | 6.4                  | 5.5                |
| transcription factor MYB44-like                | Pav_sc0001807.1_g120.1.mk               | Not identified GO                                           | 4.3                  | 3.8                |
| transcription repressor MYB6-like              | Pav_sc0000464.1_g910.1.mk               | Not identified GO                                           | -3.0                 | -2.8               |
| tannin-related R2R3 MYB transcription factor   | Pav_sc0000129.1_g770.1.br               | Not identified GO                                           | -10.0                | -9.0               |
| transcription factor bHLH94-like               | Pav_sc0006212.1_g040.1.mk               | Not identified GO                                           | -6.8                 | -10.0              |
| transcription factor bHLH92                    | Pav_sc0000998.1_g640.1.mk               | Not identified GO                                           | -6.7                 | -8.6               |
| probable WRKY transcription factor 13          | Pav_sc0002442.1_g080.1.mk               | Nucleobase-containing compound metabolic process           | -8.5                 | .                  |
| probable WRKY transcription factor 70          | Pav_sc0001582.1_g320.1.mk               | Nucleobase-containing compound metabolic process           | -3.4                 | 3.9                |
| Name | ID | Early-season variety | Mid-season variety |
|------|----|----------------------|-------------------|
| probable WRKY transcription factor 46 | Pav_sc0000254.1_g030.1.mk | Nucleobase-containing compound metabolic process | -3.0 | 4.5 4.0 |
| probable WRKY transcription factor 26 | Pav_sc0000375.1_g500.1.mk | Nucleobase-containing compound metabolic process | . | 2.4 3.2 |
| probable WRKY transcription factor 11 | Pav_sc0000624.1_g520.1.mk | Nucleobase-containing compound metabolic process | . | 2.6 2.6 |
| probable WRKY transcription factor 7 | Pav_sc0000852.1_g460.1.mk | Nucleobase-containing compound metabolic process | . | 2.8 3.2 |
| probable WRKY transcription factor 40 isoform X1 | Pav_sc0000890.1_g500.1.mk | Nucleobase-containing compound metabolic process | . | 5.4 6.8 |

| Name | ID | GO Biological Process | CT4-T0 | GT4-T0 | CT4-T0 | GT4-T0 |
|------|----|-----------------------|--------|--------|--------|--------|
| DNA (cytosine-5)-methyltransferase 1-like | Pav_sc0000113.1_g130.1.mk | DNA metabolic process | -5.2 | -4.4 | -2.1 | -2.4 |
| homocysteine S-methyltransferase 1 | Pav_sc0001051.1_g010.1.mk | Biosynthetic process | -2.2 | -2.6 |     |     |
| histone-lysine N-methyltransferase ASHR3 | Pav_sc0000065.1_g730.1.mk | Cellular protein modification process | -5.8 | -6.6 | -2.6 | -3.6 |
| histone acetyltransferase KAT6B | Pav_sc0001313.1_g160.1.mk | Not identified GO | -6.3 | -5.5 | -5.0 | -4.7 |
| histone-lysine N-methyltransferase ATXR6 | Pav_sc0000138.1_g1060.1.mk | Cellular protein modification process | -2.4 | -2.9 |     |     |
| histone-lysine N-methyltransferase SETD1B-like | Pav_sc0000363.1_g430.1.mk | Not identified GO | -4.1 | -3.3 |     |     |
Many genes coding for MYB44-like transcription factor were upregulated in CT4-T0, but they were less induced in the GT4-T0 comparison in the mid-season variety. In contrast, other transcription factor genes were downregulated in the early-season variety, including a gene coding for a putative tannin-related R2R3 MYB. Finally, both varieties had downregulated genes encoding putative histone methyl transferases and a DNA (cytosine-5)-methyltransferase 1-like in the CT4-T0, which were slightly less expressed in the GT4-T0 comparison.

Discussion

Physiological differences between early- and mid-season varieties

Sweet cherry varieties are considered early-, mid-, or late-season depending on the harvest time, which depends on the flowering time and the fruit developmental period. In our experimental conditions, early- and mid-season varieties flowered the same day in the 2017–2018 season and within a two-day time frame in the 2018–2019 season. Therefore, the varieties utilized in this work differed mainly in their fruit developmental length and, more specifically, in the duration of the slow-growth period that occurs during the light green-to-straw yellow transition, which was longer in the mid-season variety (Fig. 1A), similar to a previous report. These results imply that the ripening processes, including color initiation, begin earlier in the early-season varieties. In fact, in the early-season variety, pink color initiates at 40 DAFB (Fig. 1B), whereas in the mid-season variety, it occurred from 50 DAFB onwards, several days after growth resumption (Fig. 1A).

GA1 and GA4 have been reported to be present at the onset of ripening in sweet cherry fruits, and GA4 negatively correlates with ripening parameters. We found that the color initiation differences between both varieties were accompanied by differences in the GA4 and the GA1 content. For instance, the early-season variety, which colors first, had lower GA4 levels than the mid-season variety at 38 and 44 DAFB (Fig. S1B); and GA1 levels were higher in the mid-season variety at 38 DAFB (Fig. S1C); therefore, possibly low levels of GAs are needed for the ripening triggering. In strawberry, another non-climacteric fruit, color change coincides with the GA4 decrease between the white and the red stage. In the non-climacteric sweet pepper (Capsicum annuum L.), silencing of DNA methylase gene CaMET1-like1 caused repressed expression of GA biosynthesis genes and led to premature ripening. Taking this evidence together, GA could be a negative regulator of ripening, explaining the differences in the timing of the color development initiation between contrasting sweet cherry varieties.

Differential effect of GA3 on fruit quality parameters between early- and mid-season varieties

To further investigate the GA’s role in ripening triggering, we altered GA homeostasis by treating fruits with GA3, a GA commonly used in agronomic practices. The treatment increased firmness in both varieties. The firmness effect of GA3 may be associated with changes in cell wall composition. Kondo and Danjo found that during sweet cherry fruit ripening, cell wall-bound neutral sugars – galactose, arabinose, among others – were reduced, which is associated with fruiting softening, and that GA3 treatment prevented these decrease, supporting the role of GA in fruit firmness maintenance.

It is commonly accepted that early-season varieties are unresponsive to GA3; this is based on some reports showing no effect of GA3 on some ripening related parameters. For instance, Choi et al. found that GA3 did not affect fruit size and firmness in early-ripening genotypes. In contrast, we found that GA3 impaired color, reduced IAD and increased fruit weight at harvest in the two seasons evaluated (Fig. 3A and Fig. S3A; Table 1 and Table S5), similar to
our recent report in other early-season variety, where IAD was lower in GA$_3$-treated fruits compared with control fruits$^{17}$. In the early-season variety Satohnishiki, GA$_3$ treatment decreased anthocyanin fruit content and modified the sugar accumulation pattern$^{19}$.

On the other hand, it is usually affirmed that GA$_3$ affects ripening-related parameters in mid- or late-season varieties; this is because of a delay in softening, fruit size increase, and soluble solids accumulation$^{14,15,16}$. However, since the fruits ripen normally and only need more time to acquire the desired features, it would be correct to refer to a delay. The idea of a ripening impairment arises from studies that analyzed only the harvest point and found less color in the GA$_3$-treated fruits$^{16,28}$. Therefore, the monitoring of fruit-ripening parameters is crucial for a better understanding of the physiological effect of GA$_3$ on late- or mid-season varieties. For this, we used the IAD ripening index$^{26}$, already utilized in the context of sweet cherry fruit ripening$^{25}$. We observed a color difference as soon as 15 days after the GA$_3$ treatment (Fig. 3C), accompanied by an IAD delay in the GA$_3$-treated fruits of the mid-season variety (Fig. 3B). This delay was also found in other mid-season variety$^{17}$, which supports the idea that mid-season varieties respond differently to GA$_3$ treatment than early-season varieties. GA$_3$-treated fruits of the mid-season variety had less color, anthocyanin content, and IAD value at harvest (Fig. 2 and S2). However, the IAD value of GA$_3$-treated fruits reached similar control values four or five days later (Fig. 3 and S3), suggesting that there is no ripening impairment in the mid-season variety; rather, a normal ripening process occurs but is delayed. A previous report shown that late- and mid-season varieties had delayed color development after the GA$_3$ treatment, and maturity was two to three days delayed in the GA$_3$-treated fruits$^{16}$.

**GA ripening control differs between early- and mid-season varieties at the transcriptomic level**

The differential physiological response to exogenous GA$_3$ between early- and mid-season varieties, and the advanced color initiation in the early-season variety, which is accompanied by less GAs levels, led us to hypothesize that the ripening process of these contrasting varieties is very different at the molecular level, and possibly it may be differentially modulated by GA. Therefore, we studied the effect of GA at the transcriptomic level by altering the GA homeostasis with exogenous application of GA$_3$. This treatment was applied on the same date in both varieties in the 2017–2018 season, when they were at a similar physiological stage, as revealed by the IAD parameter (Fig. 3A and B); allowing the transcriptomic data to be comparable between varieties. The date selected for the treatment was 35 DAFB in the 2017–2018 season, five and 15 days before early- and mid-season variety color initiation, respectively, to characterize the molecular events modulated by GA at the onset of ripening and the early differences between both varieties.

To characterize the effect of GA$_3$, we considered CT4-T0 and GT4-T0 comparisons, which contain GA-modulated genes since they exclude the genes that changed independently of the GA$_3$ treatment (the intercepts of the diagrams; Fig. 4). As mentioned, CT4-T0 up and down comparisons contain genes that changed from T0 to T4 but did not change in the GT4-T0 comparison, i.e., GA-modulated genes since they no longer changed after the GA$_3$ treatment. Therefore, it can be deduced that GA$_3$ avoids their up or downregulation.

GT4-T0 up and down comparisons also contain GA-modulated, since their gene expression did not change from T0 to T4 in the control fruits, but the GA$_3$ treatment up or downregulated them.

The GO terms ‘protein phosphorylation’ and ‘photosynthesis’ were overrepresented in the CT4-T0 comparison of the early-season variety (Table 2), with genes downregulated in this comparison (Fig. 5A). This finding means that these
genes downregulated from T0 to T4 in the control fruits, but they no longer changed after the GA$_3$ application. Therefore, it is deduced that GA$_3$ impedes this downregulation, which was expected since the photosynthetic rate and capacity are usually positively controlled by GAs$^{29,30}$. 

On the other hand, we observed that 'auxin response' was an overrepresented GO category in the GT4-T0 comparison, which means that the transcript levels of the genes in this category only changed after the GA$_3$-treatment. The genes in the overrepresented GO category 'auxin response' upregulated in the GT4-T0 comparison (Fig. 5A), and some of them were exclusive of the early-season variety (Fig. 7). Several putative genes of the Aux/IAA gene family coding for auxin-responsive proteins IAA29, IAA13, IAA6, among others, were upregulated from T0 to T4. These are mainly repressors of auxin responses in fruits$^{31}$, therefore in the early-season variety, the auxin responses should turn off as ripening is initiating, since auxin usually antagonizes the ABA and ethylene promoting effect$^{6,7}$. In this regard, auxin response should be activated by GA, since several genes encoding putative Aux/IAA repressors were downregulated in the GT4-T0 comparison. Reduced expression of these repressors should activate the auxin response; therefore, GA may be a positive regulator of the auxin pathway.

Interestingly, several genes encoding auxin-induced protein 15A and 15A-like were more upregulated in GT4-T0 comparison of the early-season variety than in the CT4-T0 comparison, whereas other different genes encoding a protein 15A-like were downregulated in the mid-season variety. These results demonstrate that there is an antagonistic IAA response between both varieties. It also suggests a gene specialization, where only early-season auxin-induced protein 15A and 15A-like genes might be GA-responsive.

'Carotene metabolic process' GO category had two upregulated genes in the CT4-T0 comparison (Fig. 5A). Carotenoids are precursors of ABA, a key regulatory hormone that triggers ripening in non-climacteric fruits$^{2,32,33}$. Several genes encoding putative ASR (abscisic acid stress ripening proteins) were upregulated only in the early-season variety. The ASR genes encode transcription factors induced by ABA and abiotic stress in several plant tissues, including fruits$^{34}$. Additionally, two genes encoding an ABA degrading enzyme, abscisic acid 8'-hydroxylase 1 (CYP707A1), were more downregulated in the CT4-T0 than in the GT4-T0 comparison; hence, it is deduced that GA is a positive regulator of these genes. The silencing of PavCYP707A2 in sweet cherry led to a delay in the transcript accumulation of several genes of the ABA and anthocyanin synthetic pathways$^{35}$, supporting its role as a negative regulator of ripening. GA$_3$ enhances its expression according to our results (Table 5), in agree to our recent report showing that GA$_3$ increases the transcript abundance of PavCYP707A2 five days after the treatment, but only in the early-season variety$^{17}$.

In the mid-season variety, the GO term 'carbohydrate metabolic process' was overrepresented in both the GT4-T0 and the CT4-T0 comparisons (Table 2). Interestingly, this category's genes were downregulated in the CT4-T0 comparison, whereas they were upregulated in the GT4-T0 comparison (Fig. 5B), implying that GA strongly activates these genes. Several genes encoding putative xyloglucan endotransglucosylase/hydrolases were strongly upregulated by GA$_3$ in the mid-season variety (data not shown). These enzymes are cell wall-modifying proteins and loosen cell walls hence conferring cell wall extensibility. Interestingly, transient size increase was observed only in the mid-season variety four days after the GA$_3$ treatment (Fig. S5).

Another overrepresented GO term in the mid-season variety was 'response to stress' (Table 2). Genes in this category were downregulated in CT4-T0 but not in GT4-T0. Therefore, it is deduced that GA prevents their downregulation. A gene coding for a putative ethylene biosynthetic gene, 1-aminocyclopropane-1-carboxylate oxidase homolog 1-like, was also downregulated in CT4-T0 but not in GT4-T0. Possibly GA has a positive effect on ethylene synthesis, and
this hormone, in turn, maintains activated some stress responses. Ethylene seems to play a promoting role in non-climacteric fruit ripening\textsuperscript{36}; therefore, it is intriguing that GA, a hormone that antagonizes ripening, increases ethylene levels. This supports the idea that GA, in late- or mid-season varieties, does not negatively affect ripening. Additionally, there is a more increased ethylene response in the mid-season variety; hence, ethylene pathway might be significant in this variety.

The GO terms 'regulation of transcription, DNA-templated' and 'regulation of nucleic acid-templated transcription' were overrepresented in the mid-season variety, and they included genes that exclusively changed in this variety (Table 3; Fig. 7). Several genes coding for MYB44-like transcription factors were upregulated only in the mid-season variety (Table 5). Additionally, they were GA-downregulated as they were less upregulated in the GT4-T0 comparison. Also, a gene coding for an MYB44 transcription factor was present in both varieties, but it was more strongly expressed in the mid-season variety. MYB44 proteins suppress the expression of \textit{PP2C} genes encoding type 2C protein phosphatases\textsuperscript{37}. PP2Cs are key proteins of the ABA regulatory network and are negative regulators of subfamily 2 of SNF1-related kinases (SnRK2s), therefore negative regulators of ABA responses\textsuperscript{38}. \textit{PavSnRK2s} and \textit{PavPP2Cs} are expressed during sweet cherry fruit development\textsuperscript{8}. Since \textit{MYB44} related genes were upregulated in the mid-season variety, they could positively regulate the ABA response.

Interestingly, in the GT4-T0 comparison, these genes were less upregulated than in the CT4-T0 comparison, suggesting that GA downregulates the expression of \textit{MYB44} genes. This could be a mechanism for GA to antagonize the ABA pathway. MYB44 transduce environmental signals and mediate stress responses\textsuperscript{39}. It also mediates the interaction of hormones; for instance, it regulates the antagonistic interaction between salicylic acid and jasmonic acid pathways\textsuperscript{39}, besides the modulation of the ABA signaling. Therefore, it seems to be a crucial regulatory element for converging pathways.

\textbf{Ripening differences between maturity time contrasting varieties: what is in common and what is different?}

We observed that some ripening-related molecular features were more activated in the early-season variety, including genes related to photosynthesis, carotenoid, and ABA pathway. Photosynthesis is a negative biomarker of ripening\textsuperscript{40}, and in sweet cherry, chlorophyll decrease is one of the first events at the ripening initiation, even before sugar and anthocyanin content increase\textsuperscript{33}. 'Photosynthesis' was an overrepresented category in the early-season variety (Table 2), and several photosynthesis-related genes were downregulated in the CT4-T0 comparison (Fig. 5A). This might be exclusive of the the early-season phenotype, as this GO term was not overrepresented in the mid-season variety.

Auxin counteracts the ripening processes\textsuperscript{2,6}. Here we found that a probable indole-3-pyruvate monooxygenase \textit{YUCCA10} was upregulated in both varieties from T0 to CT4. This agrees with the increase in the IAA content detected in both varieties from 34 to 44 DAFB (Fig. S6A). Therefore, possibly this hormone prevents that the ABA ripening triggering occur too early. On the other hand, GA seems to control the IAA response, as discussed above, rather than the IAA biosynthesis, with differences between varieties in the expression of auxin induced genes.

Carotenoid and ABA biosynthesis is a positive biomarker of ripening\textsuperscript{40}. In the early-season variety, the ‘carotenoid metabolism’ category contained genes upregulated in the CT4-T0 comparison (Fig. 5A). Phytoene synthase and \textit{NCED1} like genes were upregulated in this comparison in both varieties (Table 4), possibly involved in carotenoid production and ABA biosynthesis. Additionally, in the CT4-T0 comparison, a probable carotenoid cleavage dioxygenase 4 chloroplastic gene was upregulated in the early-season variety, which probably degrades carotenoids,
since in chrysanthemum (*Chrysanthemum morifolium*) a possible orthologue of this gene contributes to white petal color\(^4\). Regarding the ABA biosynthesis, in sweet cherry, NCED1 transcript variations correlate with ABA increase during ripening\(^4\). The NCED1 upregulation in CT4-T0 implies that ABA increases in the early-season variety at the onset of ripening. We found that ABA content increases from 34 to 44 DAFB in the early-season variety, which occurred later in the mid-season variety (Fig. S6B).

Finally, phenylpropanoid early branches are negative biomarkers of non-climacteric ripening\(^40\). In this regard, we found that a tannin-related R2R3 MYB transcription factor was strongly repressed in the early-season variety, compared with the mid-season variety.

Chromosome reorganization, changes in histone methylation profile and DNA hypomethylation are characteristic features of ripening initiation\(^42\). For instance, anthocyanin-deficient apple mutant had higher methylation levels in the promoter of a *MdMYB* gene\(^43\). Silencing of a methylase gene in the non-climacteric sweet pepper led to premature ripening\(^27\). We observed several downregulated genes in the CT4-T0 comparison encoding putative histone modification enzymes and DNA (cytosine-5)-methyltransferase 1-like (Table 5). These genes were downregulated in both varieties, suggesting that chromosome remodeling and DNA hypomethylation could be relevant at this stage of development. However, some were slightly more downregulated in the early-season varieties, suggesting that these processes could be advanced in this variety, which could lead to earlier ripening initiation in early-season varieties. Additionally, there are some early-season variety-exclusive genes, including histone-lysine N-methyltransferase *ATXR6*, which is involved in the transcriptional repression in Arabidopsis\(^44\). On the other hand, some genes in the 'DNA packaging' GO categories were exclusive of the early-season variety (Table 3), suggesting that though DNA related changes may occur in both varieties, they have variety-dependent particularities.

The GA pathway seemed to be downregulated earlier in the early-season variety. Supporting this idea, several gibberelin 2-beta-dioxygenase related genes were downregulated from T0 to T4 in the early-season variety (Table 4). These genes encode putative enzymes involved in GA degradation, previously characterized in grapevine fruits\(^9\). With low GA levels, these genes downregulate as they act in a negative feedback for controlling GA levels. Therefore, possibly a low GA content may be required for the onset of ripening to occur in the early-season variety, as it was observed (Fig. S1B and Fig. S1C).

GA\(_3\) treatment upregulates the DNA (cytosine-5)-methyltransferase in the early-season variety, so possibly higher methylation levels occur when GA content is elevated, thus impairing the ripening processes. The effect of GA\(_3\) on IAA and ABA pathways in the early-season variety, as discussed above, also supports that GA exerts a negative effect on the ripening processes.

Higher GA levels could be retarding the onset of ripening in the mid-season variety, possibly to reach full embryo development. The regulatory module controlling the timing of ripening initiation could involve GA repression of *MYB44-like* gene expression, a putative negative regulator of ABA signaling genes, *PP2Cs*. Less expression of *MYB44-like* genes could lead to accumulation of negative *PP2Cs* transcripts. Therefore, more *PP2Cs* expression could imply that more ABA is needed to surpass this effect, and this could explain the ripening delay upon the GA\(_3\) treatment. In the mid-maturing sweet cherry variety Lapins, GA\(_3\) increases *PavPP2C3* and *PavPP2C4* transcripts and delays ripening\(^17\), which supports our findings.

The summary of possible molecular interactions controlling ripening in early- and mid-season varieties is depicted in Fig. 8, where a possible model is proposed. This work shows evidence supporting GA's divergent role in the ripening process of two contrasting sweet cherry varieties and provides a better understanding of non-climacteric fruit ripening.
Materials And Methods

Plant material

Sweet cherry (*Prunus avium*) fruits of four years old trees grafted on Cab-6P rootstocks growing in the *arboretum* of INIA Los Tilos Experimental Station, located in Buin, Región Metropolitana, Chile (33°42' S, 70°42' W), were selected for the experiments during the 2017–2018 and 2018–2019 seasons. The plants were in a 4 x 2 array, had regular nutritional, irrigation, and phytosanitary management, and did not receive plant growth regulator treatments. The varieties selected for the analyses were red varieties 'Bing' and 'Celeste', which have differences in the harvest time since 'Bing' is a mid-season variety, whereas 'Celeste' ('Celeste ®, denomination Sumpaca, 13S.24.28) is an early-season variety. These varieties' phenology was determined every 2–4 days during both seasons based on fruit color variations from October in 2017 and 2018 (Tables S1 to S4). Three trees were used per variety for all the analyses and measurements, where each tree is a biological replicate. 0 DAFB (days after full bloom) was assigned when 50% of the flowers were open and was September 28 for both varieties in the 2017–2018 season, whereas in the 2018–2019 season, it was October 1 and 3 for early- and mid-season varieties, respectively.

**GA₃ treatment and sampling**

Three trees for each variety were used for GA₃ treatments performed in 2017–2018 and 2018–2019. Six branches were selected from each tree, three control branches and three GA₃-treated branches, following Usenik’s methodology. GA₃ (ProGibb ® 40% SG) was dissolved in water and applied to individual branches with a hand sprayer to run-off at a rate of 30 ppm when the fruits of each variety were in late Stage II of development, and fruit color transitioned from green to straw yellow. Control branches were treated with water and protected from spraying with GA₃, according to Usenik et al. The treatment was performed at the same hour of the day in both varieties and both seasons (12:00 GMT). Ten fruits per repeat (branch), i.e. 30 fruits per treatment, were sampled, with sample size according to Luo et al. Fruit samples were homogenous in color, size and form, and without any visible defect. The fruits were immediately frozen in liquid nitrogen after removal from the tree and stored at -80°C until used for RNA-seq analysis, anthocyanins estimations, and hormone quantification.

**Fruit parameters**

For fruit width, a caliper was used to measure the equatorial diameter at the fruits’ widest point. VIS/NIF device Cherry Meter (T.R. ® Turoni, Italy) was used to measure the ripening index, IAD (Index of Absorbance Difference), according to Nagpala et al. Non-destructive fruit width and IAD assessments were performed during the growing season until ripeness. Fruit width and IAD were measured individually in 20 fruits randomly selected from each tree. Fruit weight, firmness, soluble solids content, and acidity (malic acid) were measured at the ripeness of each variety. Fruit weight was quantified using a portable mini scale, and firmness was determined on two opposite cheeks using a durometer device (Durofel T.R. ® Turoni, Italy), according to San Martino et al. Pocket Brix-Acidity Meter (PAL-BX|ACID3, ATAGO USA, Inc.) was used to quantify soluble solids content (SSC) and acidity as a percentage of malic acid, as reported by Sediqi et al. Color distribution was determined at harvest using a CTIFL color chart with 1 to 4 values (1 = light red, 2 = red, 3 = dark red, 4 = light mahogany). Fruit parameters were measured individually in 25 fruits randomly selected from each tree. For acidity and SSC, a subset of five fruits of the 25 fruits was measured, according to Chavoshi et al.

**RNA extraction and RNA seq-analysis**
For RNA-seq analysis, samples were collected minutes before the GA₃ treatment (T0 samples) and four days after the GA₃ treatment (GT4 samples) or control treatment (CT4 samples) in the 2017–2018 season. RNA was extracted from 18 samples consisting of three replicates of T0, CT4, and GT4 samples (nine samples per variety). T0 samples of early- and mid-season varieties had a similar color and IAD value, which was around 0.4 in T0 samples in the 2017–2018 season. For the RNA-seq analysis the 2017 samples were used (GA₃ application on November 2, 2017; 35 DAFB, 12:00 GMT in both varieties).

Total RNA was isolated from 0.5 g of ground flesh- and peel-enriched tissue using the CTAB-method with minor modifications, according to Meisel et al. GeneJET RNA Cleanup and Concentration Micro Kit (Thermo Scientific™, San Diego, CA, USA) was used for purifying the RNA samples. Purity values, A₂₆₀/₂₃₀ and A₂₆₀/A₂₈₀, were around 2.0 in all the samples.

For RNA-seq analysis of the 18 fruit samples, 1µg of RNA with RIN (RNA integrity number) > 7.0 was used to generate cDNA libraries of the fruit samples using the Illumina TruSeq stranded mRNA Library Preparation Kit, according to manufacturer's instructions. The libraries were sequenced in an Illumina platform (Illumina NovaSeq6000 at Macrogen Inc.), and 100 bp paired-end reads were generated. Data was generated using the base-calling software CASAVA v1.8.2 for forward and reverse segments.

**Analysis of differentially expressed genes**

Adaptors, low-quality bases, and short sequences trimming were performed using CLC Genomics Workbench version 11.0.1 following parameters: Q ≥ 20; no more than two ambiguities; final read length ≥ 50 bp. Sequence mapping to the reference genome was performed using CLC Genomics Workbench version 11.0.1 with the following parameters: similarity = 0.9; length fraction = 0.6; insertion/deletion cost = 3; mismatch cost = 3, and unspecific match limit = 10. Expression levels were normalized by calculating RPKM (Reads Per Kilobase Million) value. Transcript abundances were fitted using a general linear model (GLM) and differential expression of treatments tested with the Wald test against control groups. Differentially expressed genes (DEGs) were defined as having an absolute fold-change value of at least two between T4 and T0, with an adjusted p-value using a false discovery rate (FDR) with at least a 95% significance.

Functional annotation of *P. avium* reference transcripts was performed by BLAST2GO software version 5.2.5. First, a BLASTx search was performed against the NCBI NR database with an e-value cutoff of 1e-6 and HSP length cutoff of 33. Then, INTERPROSCAN analysis was performed with BLAST2GO default parameters. Finally, the data from BLAST searches, INTERPROSCAN terms, enzyme classification codes (EC), and metabolic pathways (KEGG, Kyoto Encyclopedia of Genes and Genomes) were merged in gene ontology (GO) terms for a comprehensive functional range cover in the functional annotation. The BLAST2GO program defaults were used in all mapping and annotation steps, and the false discovery rate (FDR) cutoff was set to a 0.05% probability level. GO over-representation analysis was performed with the Fisher’s Exact Test Enrichment Analysis using BLAST2GO tools and integrated FatiGO package with default parameters.

Venn diagrams were constructed using the online tool VENNY, whereas Heatmaps were constructed with the online tool Morpheus (https://software.broadinstitute.org/morpheus/)

**Hormone quantification**

ABA, IAA, GA₁ and GA₄ were measured in both varieties at 34, 38, and 44 DAFB in the 2018–2019 season. For the extraction, 10 mg of flesh- and peel-enriched tissue was freeze-dried, ground, and suspended in 80% methanol − 1%
acetic acid solution containing internal standards (deuterium-labeled hormones; OlChemim Ltd., Olomouc, Czech Republic). The mix was shaken for one hour at 4°C, and the extracted fraction was maintained at -20°C overnight. The samples were centrifuged, and the supernatant was vacuum dried and then dissolved in 1% acetic acid. A reverse-phase column (OasisHLB) was used, and the eluate was dried and dissolved in 5% acetonitrile – 1% acetic acid. An autosampler and reverse-phase UHPLC chromatography column, 2.6 µm Accucore RP-MS, 100 mm x 2.1 mm (ThermoFisher Scientific, San Diego, CA, USA) were used. Then the hormones were separated using a gradient of acetonitrile (2%-55%) containing 0.05% acetic acid, at a rate of 400 µL/min over 22 min. ABA, IAA, GA$_1$ and GA$_4$ were detected in a Q-Exactive mass spectrometer (Orbitrap detector; ThermoFisher Scientific; San Diego, CA, USA). Targeted Selected Ion Monitoring and Electrospray Ionization in the negative mode were used to detect the hormones. The quantifications were performed using external calibration curves with the Xcalibur 4.0 and TraceFinder 4.1 SP1.

**Analysis of anthocyanins**

soluble solids were determined on a small aliquot of the juice with a hand-held refractometer

soluble solids were determined on a small aliquot of the juice with a hand-held refractometer

soluble solids were determined on a small aliquot of the juice with a hand-held refractometer

Anthocyanins were measured in control- and GA$_3$-treated fruits at the fully ripeness of both varieties using three replicates per date in the 2017–2018 season. For anthocyanins extraction, 0.1 g of flesh- and peel-enriched tissue was freeze-dried and ground. The tissue was mixed with 80% methanol solution, sonicated, shaken for 20 min, and left overnight in the dark at 4°C. The samples were centrifuged for 10 min at 4°C and 4,000 rpm and the supernatant filtered using a 0.22 µm PFTE membrane. Anthocyanins were separated using a Waters Acquity HSS T3 column, 1.8 µm, 100 mm x 2.1 mm, in a UPLC-MS/MS Waters Acquity system (Milford, MA, USA), as reported by Arapitsas et al. The anthocyanins were detected in a mass spectrometry Waters Xevo TQMS instrument with an ESI source. For data processing, Mass Lynx Target Lynx Application Manager was used. Cyanidin-3-O-rutinoside was used to estimate anthocyanin since it represented more than 98% of the total anthocyanins in both varieties.

**Statistical analysis**

One-way ANOVA analysis with Tukey's *post hoc* test was used for assessing differences between control- and GA$_3$-treated fruits in estimated anthocyanins and fruit parameters, where three replicates per date were used to establish the mean differences, using the InfoStat software.

**Abbreviations**
Abscisic acid (ABA); days after full bloom (DAFB); IAD (Index of Absorbance Difference); gibberellin (GA); gibberellin A1 (GA$_1$); gibberellin A3 (GA$_3$); gibberellin A4 (GA$_4$); gibberellin A7 (GA$_7$).

**Declarations**

**Data Availability Statement**

The raw reads sequences from this work were submitted to NCBI's Sequence Read Archive through the BioProject ID: PRJNA683645.

**Acknowledgments**

We like to thank INIA Los Tilos, especially Gustavo Azócar. Also, we like to thank Simón Miranda, Natalia Molina, Carla Trigo and Elisabeth Sarabia for their technical support.

**Conflict of Interest**

The authors declare that they have no conflict of interest.

**Funding**

FONDECYT Fondecyt de Postdoctorado 3180138

FONDECYT Fondecyt Regular 1171016

FONDECYT Scholarship Grant 21190238

**Author contribution statement**

N.K., J.M. and L.M. conceived, designed, and conducted the research. N.K., C.P. and M.A. performed RNA extractions. J.M. analyzed RNA-seq results. N.K., C.P., M.A. and A.T. conducted physiological measurements in the field. E.C. performed hormone quantification. C.P., S.M. and S.M. measured fruit anthocyanin content. N.K., J.M. and L.M. wrote the manuscript. S.M., S.M., E.C., J.M.D., B.S. and L.M. performed critical review on the manuscript.

**References**

1. Fortes, A. M., Teixeira, R. T., & Agudelo-Romero, P. Complex interplay of hormonal signals during grape berry ripening. *Molecules, 20*, 9326-9343 (2015).
2. Jia, H. et al. Abscisic acid, sucrose, and auxin coordinately regulate berry ripening process of the Fujiminori grape. *Integr. Genomics* 17, 441-457 (2017).
3. Wheeler, S., Loveys, B., Ford, C., & Davies, C. The relationship between the expression of abscisic acid biosynthesis genes accumulation of abscisic acid and the promotion of *Vitis vinifera* berry ripening by abscisic acid. *Aust. J. Grape Wine Res.* 15, 195-204 (2009).
4. Ren, J. et al. Role of abscisic acid and ethylene in sweet cherry fruit maturation: molecular aspects. *New Zeal. J. Crop Hort. Sci.*, 39, 161-174 (2011).
5. Teribia, N., Tijero, V., & Munné-Bosch, S. Linking hormonal profiles with variations in sugar and anthocyanin contents during the natural development and ripening of sweet cherries. *New Biotechnol.*, 33, 824-833 (2016).
6. Gouthu, S. & Deluc, L. G. Timing of ripening initiation in grape berries and its relationship to seed content and pericarp auxin levels. *BMC Plant Biol.* **15**, 46 (2015).

7. Davies, C., Boss, P. K. & Robinson, S. P. Treatment of grape berries, a non-climacteric fruit with a synthetic auxin, retards ripening and alters the expression of developmentally regulated genes. *Plant Physiol.* **115**, 1155-1161 (1997).

8. Wang, Y. et al. Transcriptional regulation of *PaPYLs*, *PaPP2Cs* and *PaSnRK2s* during sweet cherry fruit development and in response to abscisic acid and auxin at onset of fruit ripening. *Plant Growth Regul.* **75**, 455-464 (2015).

9. Giacomelli, L. et al. Gibberellin metabolism in *Vitis vinifera* during bloom and fruit-set: functional characterization and evolution of grapevine gibberellin oxidases. *J. Exp. Bot.* **64**, 4403-4419 (2013).

10. Csukasi, F. et al. Gibberellin biosynthesis and signalling during development of the strawberry receptacle. *New Phytol.* **191**, 376-390 (2011).

11. Fortes, A. M. et al. Transcript and metabolite analysis in Trincadeira cultivar reveals novel information regarding the dynamics of grape ripening. *BMC Plant Biol.* **11**, 149 (2011).

12. Gambetta, G. A., Matthews, M. A., Shaghasi, T. H., McElrone, A. J. & Castellarin, S. D. Sugar and abscisic acid signaling orthologs are activated at the onset of ripening in grape. *Planta* **232**, 219-234 (2010).

13. Kondo, S. & Danjo, C. Cell wall polysaccharide metabolism during fruit development in sweet cherry 'Satohnishiki' as affected by gibberellic acid. *Jpn. Soc. Hortic. Sci.* **70**, 178-184 (2001).

14. Choi, C., Toivonen, P., Wiersma, P. A. & Kappel, F. Effect of gibberellic acid during development of sweet cherry fruit: physiological and molecular changes. *Acta Hort.* **636**, 429-495 (2004).

15. Kappel, F. & MacDonald, R. A. Gibberellic acid increases fruit firmness, fruit size, and delays maturity of 'Sweetheart' sweet cherry. *Am. Pom. Soc.* **56**, 21 (2002).

16. Usenik, V., Kastelec, D. & Štampar, F. Physicochemical changes of sweet cherry fruits related to application of gibberellic acid. *Food Chem.* **90**, 663-671 (2005).

17. Kuhn, N. et al. Gibberellic acid modifies the transcript abundance of ABA pathway orthologs and modulates sweet cherry (*Prunus avium*) fruit ripening in early- and mid-season varieties. *Plants* **9**, 1796 (2020).

18. Alkio, M., Jonas, U., Declercq, M., Van Nocker, S. & Knoche, M. Transcriptional dynamics of the developing sweet cherry (*Prunus avium*) fruit: sequencing, annotation and expression profiling of exocarp-associated genes. *Hort research* **1**, 1-15 (2014).

19. Wei, H. et al. Comparative transcriptome analysis of genes involved in anthocyanin biosynthesis in the red and yellow fruits of sweet cherry (*Prunus avium*). *PLoS One* **10**, e0121164 (2015).

20. Guo, X. et al. Transcriptomic analysis of light-dependent anthocyanin accumulation in bicolored cherry fruits. *Plant Physiol. Biochem* **130**, 663-677 (2018).

21. Chai, L. et al. RNA sequencing reveals high resolution expression change of major plant hormone pathway genes after young seedless grape berries treated with gibberellin. *Plant Sci.* **229**, 215-224 (2014).

22. Jiang, S., Luo, J., Xu, F. & Zhang, X. Transcriptome analysis reveals candidate genes involved in gibberellin-induced fruit setting in triploid loquat (*Eriobotrya japonica*). *Plant Sci.* **174**, 924 (2016).

23. Wen, B. et al. Identification and characterization of cherry (*Cerasus pseudocerasus* Don) genes responding to parthenocarpy induced by GA$_3$ through transcriptome analysis. *BMC genetics* **20**, 65 (2019).

24. Rodrigo, F. J. & Guerra, M. *La fruticultura del S. XXI. Ed. Cajamar*, 107-123. (2014).
25. Nagpala, E. G. L. et al. Cherry-Meter: an innovative non-destructive (vis/NIR) device for cherry fruit ripening and quality assessment. *Acta Hortic*. **1161**, 491-496 (2013).

26. Costa, G., Vidoni, S. & Rocchi, L. Use of non-destructive devices to support pre-and postharvest fruit management. *Acta Hort.* **1119**, 329-335 (2014).

27. Xiao, K. et al. DNA methylation is involved in pepper (*Capsicum annuum*) fruit ripening regulation and interacts with phytohormones. *J. Exp. Bot.* **71**, 1928–1942 (2020).

28. Einhorn, T. C., Wang, Y. & Turner, J. Sweet cherry fruit firmness and postharvest quality of late-maturing cultivars are improved with low-rate, single applications of gibberellic acid. *HortScience*. **48**, 1010-1017 (2013).

29. Wen, Y., Su, S. C., Ma, L. Y., & Wang, X. N. Effects of gibberellic acid on photosynthesis and endogenous hormones of *Camellia oleifera* in 1st and 6th leaves. *J. Forest Res.* **23**, 309-317 (2018).

30. Zhou, B. et al. Heterologous expression of a *Gibberellin 2-Oxidase* gene from *Arabidopsis thaliana* enhanced the photosynthesis capacity in *Brassica napus*. *J. Plant Biol.* **54**, 23-32 (2011).

31. Liu, D. J., Chen, J. Y. & Lu, W. J. Expression and regulation of the early auxin-responsive *Aux/IAA* genes during strawberry fruit development. *Biol. Rep.* **38**, 1187-1193 (2011).

32. Luo, H. et al. The role of ABA in the maturation and postharvest life of a non-climacteric sweet cherry fruit. *Plant Growth Regul.* **33**, 373-383 (2014).

33. Shen, X. et al. A role for PacMYBA in ABA-regulated anthocyanin biosynthesis in red-colored sweet cherry cv. Hong Deng (*Prunus avium*). *Plant Cell Physiol.* **55**, 862-880 (2014).

34. Carrari, F., Fernie, A. R. & Iusem, N. D. Heard it through the grapevine? ABA and sugar cross-talk: the ASR story. *Trends Plant Sci.* **9**, 57-59 (2004).

35. Li, Q. et al. PacCYP707A2 negatively regulates cherry fruit ripening while PacCYP707A1 mediates drought tolerance. *Exp. Bot.* **66**, 3765-3774 (2015).

36. Sun, L. et al. Reciprocity between abscisic acid and ethylene at the onset of berry ripening and after harvest. *BMC Plant Biol.* **10**, 257 (2010).

37. Nguyen, N. H. & Cheong, J. J. AtMYB44 interacts with TOPLESS-RELATED corepressors to suppress protein phosphatase 2C gene transcription. *Biophys. Res. Comm.*, **507**, 437-442 (2018).

38. Umezawa, T. et al. Type 2C protein phosphatases directly regulate abscisic acid-activated protein kinases in *Arabidopsis*. *Natl Acad. Sci.*, **106**, 17588-17593 (2009).

39. Seo, J. S. et al. Expression of the *Arabidopsis AtMYB44* gene confers drought/salt-stress tolerance in transgenic soybean. *Breed.* **29**, 601-608 (2012).

40. Fasoli, M. et al. Timing and order of the molecular events marking the onset of berry ripening in grapevine. *Plant Physiol.* **178**, 1187-1206 (2018).

41. Ohmiya, A., Kishimoto, S., Aida, R., Yoshioka, S. & Sumitomo, K. Carotenoid cleavage dioxygenase (*CmCCD4a*) contributes to white color formation in chrysanthemum petals. *Plant Physiol.* **142**, 1193-1201 (2006).

42. Gallusci, P., Hodgman, C., Teyssier, E. & Seymour, G. B. DNA methylation and chromatin regulation during fleshy fruit development and ripening. *Plant Sci.* **7**, 807 (2016).

43. El-Sharkawy, I., Liang, D. & Xu, K. Transcriptome analysis of an apple (*Malus × domestica*) yellow fruit somatic mutation identifies a gene network module highly associated with anthocyanin and epigenetic regulation. *Exp. Bot.* **66**, 7359-7376 (2015).

44. Jacob, Y. et al. ATXR5 and ATXR6 are H3K27 monomethyltransferases required for chromatin structure and gene silencing. *Struct. Biol.* **16**, 763 (2009).
45. San Martino, L., Manavella, F. A., García, D. A. & Salato, G. Phenology and fruit quality of nine sweet cherry cultivars in South Patagonia. *Acta Hort* **795**, 841-848 (2005).

46. Sediqi, A. G., Kramchote, S., Itamura, H. & Esumi, T. Physiological changes in sweet cherry fruit in response to physical damage. *Acta Hort* **1235**, 495-502 (2017).

47. Chavoshi, M. et al. Phenotyping protocol for sweet cherry (*Prunus avium*) to facilitate an understanding of trait inheritance. *J. Am. Pomol. Soc.* **68**, 125-34 (2014).

48. Meisel, L. et al. A rapid and efficient method for high quality total RNA from peaches (*Prunus persica*) for functional genomics analyses. *Res.* **38**, 83–88. (2005).

49. Shirasawa, K. et al. The genome sequence of sweet cherry (*Prunus avium*) for use in genomics-assisted breeding. *DNA Res.* **24**, 499-508 (2017).

50. Draper, N. R. & Smith, H. *Applied regression analysis*. John Wiley & Sons. **326** (1998).

51. Trapnell, C. et al. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and Cufflinks. *Protoc.* **7**, 562-578 (2012).

52. Conesa, A. & Götz, S. Blast2GO: A comprehensive suite for functional analysis in plant genomics. *J. Plant Genomics* **2008**, 619832 (2008).

53. Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. Basic local alignment search tool. *Mol. Biol.* **215**, 403-410 (1990).

54. Hunter, S. et al. InterPro: the integrative protein signature database. *Ac. Res.* **37** (suppl_1), D211-D215 (2009).

55. Al-Shahrour, F., Díaz-Uriarte, R. & Dopazo, J. FatiGO: a web tool for finding significant associations of Gene Ontology terms with groups of genes. *Bioinformatics* **20**, 578-580 (2004).

56. Oliveros, J. C. An interactive tool for comparing lists with Venn Diagrams. http://bioinfogp. cnb. csic. es/tools/venny/index. html (2007).

57. Seo, M., Jikumaru, Y. & Kamiya, Y. Profiling of hormones and related metabolites in seed dormancy and germination studies. Seed Dormancy, *Humana Press*, 99-111 (2011).

58. Arapitsas, P., Perenzoni, D., Nicolini, G., & Mattivi, F. Study of sangiovese wines pigment profile by UHPLC-MS/MS. *Agr. Food Chem.* **60**, 10461–10471 (2012).

59. Di Rienzo, J. A. et al. (2012). InfoStat versión 2012. Grupo InfoStat, FCA. Argentina: Universidad Nacional de Córdoba.