Targeted Systems Biology Profiling of Tomato Fruit Reveals Coordination of the Yang Cycle and a Distinct Regulation of Ethylene Biosynthesis during Postclimacteric Ripening$^{1[C][W][OA]}$

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The concept of system 1 and system 2 ethylene biosynthesis during climacteric fruit ripening was initially described four decades ago. Although much is known about fruit development and climacteric ripening, little information is available about how ethylene biosynthesis is regulated during the postclimacteric phase. A targeted systems biology approach revealed a novel regulatory mechanism of ethylene biosynthesis of tomato (Solanum lycopersicum) when fruit have reached their maximal ethylene production level and which is characterized by a decline in ethylene biosynthesis. Ethylene production is shut down at the level of 1-aminocyclopropane-1-carboxylic acid oxidase. At the same time, 1-aminocyclopropane-1-carboxylic acid synthase activity increases. Analysis of the Yang cycle showed that the Yang cycle genes are regulated in a coordinated way and are highly expressed during postclimacteric ripening. Postclimacteric red tomatoes on the plant showed only a moderate regulation of 1-aminocyclopropane-1-carboxylic acid synthase and Yang cycle genes compared with the regulation in detached fruit. Treatment of red fruit with 1-methylcyclopropane and ethephon revealed that the shut-down mechanism in ethylene biosynthesis is developmentally programmed and only moderately ethylene sensitive. We propose that the termination of autocatalytic ethylene biosynthesis of system 2 in ripe fruit delays senescence and preserves the fruit until seed dispersal.

In 1972, McMurchie et al. introduced system 1 and system 2 ethylene biosynthesis during fruit development and ripening based upon differences in ethylene production and respiration between climacteric and nonclimacteric fruits (McMurchie et al., 1972). Since its inception, the concept of two systems of ethylene biosynthesis has been broadly adopted and refined. System 1 is characterized by a basal level of ethylene production, is autoinhibitory, and is applicable to all vegetative tissues including developing fruits (Alexander and Grierson, 2002). At a certain moment, coordinated by a yet unknown developmental switch, climacteric fruit start to ripen and ethylene production rises autocatalytically. Ethylene production during system 2 was later characterized to be partially autocatalytically and partially developmentally regulated (Yokotani et al., 2009). The drop in ethylene production after the autocatalytic rise is less investigated. Nonclimacteric fruit ripen without an autocatalytic burst in ethylene production.

The ethylene biosynthesis pathway was unraveled mainly during the late 1970s and early 1980s (Fig. 1). It was shown that 1-aminocyclopropane-1-carboxylic-acid (ACC) is the precursor of ethylene and originates from the amino acid Met (Lieberman and Kunishi, 1966; Adams and Yang, 1979). ACC is formed by ACC synthase (ACS) using S-adenosyl-l-Met (SAM) as a substrate (Murr and Yang, 1975; Boller et al., 1979). SAM itself is produced from the amino acid Met by SAM
synthetase requiring ATP. For each ACC molecule synthesized by ACS, one 5'-methylthioadenosine (MTA) residue is released (Adams and Yang, 1977). MTA is recycled back to Met by the Met or Yang cycle for another round of ethylene biosynthesis. During the first step of Met recycling, MTA is converted to 5'-methylthioribose by 5'-methylthioadenosine nucleosidase (MTN; Adams and Yang, 1977; Wang et al., 1982;
Rzewuski et al., 2007). 5′-methylthioribose is phosphor-ylated to 5′-methylthioribose-1-phosphate (MTR-P) by 5′-methylthioribose kinase (MTK; Kushad et al., 1982; Sauter et al., 2004). Just recently, Pommernenig et al. (2011) characterized two more enzymes of the Yang cycle in plants. The first is MTR-P isomerase (MTI), which catalyzes the isomerization of MTR-P into 5′-methylthioribulose-1-P. In a second step, 5′-methylthioribulose-1-P is converted to 1,2-dihydroxy-3-keto-5-methylthiopentene by a dehydratase-enolase-phosphatase (DEP). 1,2-dihydroxy-3-keto-5-methylthiopentene (also termed acireductone) is converted into 2-keto-4-methylthiobutyrate by acireductone dioxygenase (ARD) in the presence of oxygen (Sauter et al., 2005; Bürstenbinder et al., 2007). Finally, 2-keto-4-methylthiobutyrate is further metabolized to Met by a yet unknown transaminase (Kushad et al., 1983; Pommernig et al., 2011).

The ethylene precursor ACC is converted into ethylene by ACC oxidase (ACO, previously called the ethylene-forming enzyme) in the presence of oxygen (Hamilton et al., 1991; Dong et al., 1992). ACC can also be conjugated to the biologically inactive metabolite 1-(malonylamino)cyclopropane-1-carboxylic acid (MACC) by ACC-N-malonyltransferase (Hoffman et al., 1982; Liu et al., 1983). MACC was thought to be an end metabolite, but Jiao et al. (1986) and Hanley et al. (1989) showed that MACC can be reconverted into ACC. Later it was found that a minor amount of ACC can also be converted into the low abundant conjugate 1-(y-glutamylamino)cyclopropane-1-carboxylic acid (Martin et al., 1995). More recently, Staswick and Tiryaki (2004) showed the presence of a low abundant ACC derivative of jasmonic acid in Arabidopsis (Arabidopsis thaliana).

The enzymes solely involved in ethylene biosynthesis are encoded by multigene families. So far, five different isoforms have been identified for ACO in tomato (Solanum lycopersicum), and they are differentially expressed during fruit development and ripening. During system 1, ACO1 and ACO4 are expressed moderately, whereas ACO3 is transiently expressed during the transition phase. During system 2, ACO1 and ACO4 expression increases severalfold, peaking at the orange stage (Barry et al., 1996; Nakatsuka et al., 1998). A fifth member of the ACO family (ACO5) shows an anaerobically induced expression in both fruit and leaves, but has not yet been profiled during fruit development and ripening (Sell and Hehl, 2005).

For ACS, nine genes have been identified in tomato so far (ACS1A, ACS1B, ACS2–ACS8) of which five are expressed during fruit ripening. ACS1A and ACS3 are constitutively expressed at a low level during system 1, whereas ACS6 shows a higher expression (Nakatsuka et al., 1998; Barry et al., 2000). During system 2, ACS2 and ACS4 are highly expressed (Yip et al., 1992; Nakatsuka et al., 1998; Barry et al., 2000).

The expression of both ACS and ACO genes is regulated by recently identified transcription factors in tomato. HB-1 is a homeobox Leu zipper protein that binds to the promoter of ACO1, altering its expression (Lin et al., 2008), whereas ETHYLENE RESPONSE FACTOR2 (ERF2) was shown to interact with the GCC box of the ACO3 promoter, stimulating ACO3 expression (Zhang et al., 2009). The RIPENING INHIBITOR (RIN) transcription factor was also shown to interact with the promoter region of ACS-encoding genes ACS2 and ACS4 (Ito et al., 2008; Fujisawa et al., 2011). Martel et al. (2011) showed a clear correlation between RIN expression, RIN abundance, and ACS2 expression.

Early biochemical studies led to the structural and functional characterization of the two enzymes specific for ethylene biosynthesis. The ACO protein belongs to the superfamily of oxygenases and oxidases with a requirement for iron as cofactor and bicarbonate as activator (Dong et al., 1992; Zhang et al., 2004). Ascorbic acid is an essential additive during in vitro activity measurements and most probably also for in vivo functionality, although the exact role of this effector remains uncertain (Rocklin et al., 2004). ACO was localized in the cytosol by immunocytolocalization (Reinhardt et al., 1994; Chung et al., 2002; Hudgins et al., 2006), but some authors reported ACO localization at the plasma membrane (Rombaldi et al., 1994; Ramassamy et al., 1998). Its three-dimensional structure was determined by x-ray crystallography (Zhang et al., 2004). In vitro analysis of ectopically expressed ACO proteins revealed that each isoform has a specific activity with ACO1 being the most active member (Bidone et al., 1998).

The ACS protein belongs to the pyridoxal-5′-P (vitamin B6)-dependent enzymes, requiring pyridoxal-5′-P as an essential cofactor and is located in the cytosol (Boller et al., 1979). ACS can be posttranslationally stabilized by phosphorylation to prevent degradation by the ubiquitin/proteasome pathway (Tatsuki and Mori, 2001; Joo et al., 2008; Kamiyoshihara et al., 2010). The three-dimensional structure of apple (Malus domestica) ACS homodimer was obtained by Capitani et al. (1999). It was shown that ACS proteins can form heterodimers that increase both the structural and functional complexity (Tsuchisaka and Theologis, 2004).

Climacteric fruit ripening was most extensively studied in tomato (Alexander and Grierson, 2002). ACC content has been profiled in the past during system 1 and system 2 and shows a close correlation with the level of fruit ethylene production (Hoffman and Yang, 1980; Su et al., 1984). ACC levels decline after the climacteric peak, but rise again at later stages (Hoffman and Yang, 1980). MACC follows a similar pattern, although this metabolite has not yet been profiled during postharvest stages to our knowledge. Met, on the other hand, was rarely investigated in relation to ethylene biosynthesis. Metabolome analysis showed that Met levels increase during tomato fruit ripening (Carrari and Fernie, 2006; Oms-Oliu et al., 2011). To our knowledge, no information is available on SAM and MTA profiles in ripening fruit.

In this study, we aimed to investigate fruit development, climacteric ripening, and postclimacteric development during postharvest storage of tomato by means of a targeted systems biology approach. Our intent was to integrate metabolic, enzymatic, and transcriptomic
profiles to characterize the regulation of the ethylene biosynthesis pathway including the Yang cycle. The current understanding of preclimacteric and climacteric fruit ripening (mainly confirming literature) is expanded and linked to new findings on the poorly investigated postclimacteric phase of ethylene biosynthesis. This approach revealed a novel regulatory mechanism for ethylene biosynthesis during the postclimacteric ripening of tomato fruit.

RESULTS
Fruit Physiology and Quality

Various physiological parameters were measured to obtain a representative data set describing fruit development, climacteric ripening, and postharvest storage. Straight after sampling, fruit ethylene production and respiration rates were measured together with color and firmness, two important fruit quality parameters. Overall, fruit ethylene production showed a typical climacteric behavior with low ethylene production during development (system 1) and a climacteric rise during ripening (system 2; Fig. 2A). During postclimacteric development ethylene production decreased close to, without quite reaching, the basal level. A typical high respiration rate was observed for very small developing fruit that declined to a basal level at the immature green stage. Fruit respiration rate subsequently showed a distinct increase at the onset of fruit ripening and declined again during further ripening and storage (Fig. 2B). Fruit color showed a logistic shift from green to red during ripening (Supplemental Fig. S1A). Fruit firmness measurements showed that softening began at the breaker stage and continued until the red stage was reached (Supplemental Fig. S1B).

Metabolic Profiling Reveals that ACC Is Not Limiting for Postclimacteric Ethylene Production

The relation between ethylene production and intermediate metabolites was further investigated by metabolic profiling. Free Met (Fig. 3A) continuously accumulated throughout fruit development and postharvest storage. Met is a general metabolite that appeared not to be limiting because it was present at a micromolar concentration compared with the nanomolar concentration of other metabolites. SAM and MTA profiles have not yet been previously published for tomato. The SAM profile (Fig. 3B) showed a clear correlation with ethylene production: low during fruit development, followed by a drastic increase during ripening and a steady decline during the postclimacteric stages. SAM levels began to increase in parallel with ethylene production but reached maximal levels at the breaker stage prior to the ethylene maximum, which was reached at the orange stage. MTA followed a similar profile like SAM, albeit less pronounced and around 20 times lower in concentration (Fig. 3C).

ACC and MACC metabolic profiles were similar to previously published results (Fig. 3, D and E), with low levels during preclimacteric development and a strong increase during climacteric ripening (Hoffman and Yang, 1980; Su et al., 1984). During postharvest storage, ACC content leveled off, whereas MACC content slowly continued to increase. MACC content was about 60 times higher than ACC content and steadily increased, indicating that MACC is most probably an end product. In conclusion, the high ACC content during the postclimacteric stages suggests that ACC content was not limiting for ethylene biosynthesis during these stages.

Enzymatic Profiling Reveals that ACO Catalyzes the Rate-Limiting Step during Postclimacteric Ethylene Biosynthesis

To relate metabolic changes to key enzymes of ethylene biosynthesis, we determined the in vitro enzyme
activities of ACO, ACS, and MTN during fruit development, ripening, and postharvest storage (Fig. 4). Large differences in absolute activity values were observed between ACS and the other enzymes analyzed. ACS is known to have a low abundance and stability. Together with the tedious extraction procedure, this might have caused the low in vitro activity. During preclimacteric development, ACO activity increased whereas ACS activity decreased, supporting the general observation that ACS is the rate-limiting enzyme for ethylene biosynthesis. Once the fruit started to ripen, ACO and ACS activities increased 2- to 3-fold. At the light orange to orange stage, when ethylene production had reached its maximum, ACO and ACS activities both declined. Surprisingly, ACS activity rose again at the end of ripening and during subsequent postharvest storage, whereas ACO activity dropped during storage, almost to the low activity observed in preclimacteric fruit. Reduced ACO activity and a sustained activity of ACS resulted in the accumulation of ACC (Fig. 3D). This strongly suggests that ACO is rate limiting for ethylene biosynthesis during the postclimacteric stages.

MTN activity was high during early fruit development but rapidly dropped to an intermediate level during preclimacteric ethylene production and remained low during climacteric ripening (Fig. 4C). The activity rose again at the pink to red ripe stage, just prior to the decrease in MTA levels (Fig. 3C). This delay may indicate a feed-forward activation of MTN by MTA. During postharvest storage, MTN activity reached a maximum similar to ACC and MACC. All in all, MTN activity shows no direct correlation with climacteric ethylene production but correlated well with ACC/MACC synthesis during the postclimacteric stages.

**Western-Blot Analysis Confirms Both ACO and ACS Activity**

To study the regulation of ethylene biosynthesis at the protein level, ACO and ACS protein content was visualized by western blotting (Fig. 5). The antibodies used were designed to recognize a consensus peptide epitope of four tomato ACO isoforms (ACO1 to -4) and of four tomato ACS isoforms (ACS1, ACS2, ACS4, and ACS6). The ACO western blot showed a clear band just above the 37-kD marker corresponding to the predicted masses of all tomato ACO isoforms. During early fruit development, hardly any ACO protein was detected, although a very faint band was visible prior to ripening. At the onset of ripening, ACO content increased toward a maximum around the breaker to pink stage. From the red ripe stage onward,
ACO protein levels decreased again. Abundance of the 37-kD ACO protein thus correlated well with high ACO activity observed during fruit ripening (Fig. 4A).

Western-blot analysis of ACS showed a faint band just above 50 kD, corresponding to the mass of ACS (Fig. 5B). The weak signal is probably caused by the low ACS abundance in tomato pericarp and the weak affinity of the anti-ACS antibodies. An unspecific band, just slightly higher than the ACS band, blurs the signal. Nevertheless, ACS appeared to be present during early fruit development with the highest signal intensity observed at the breaker stage corresponding to the high ACO activity measured at this stage (Fig. 4B). During further ripening, the ACS protein level remained constant and was still clearly detectable during the postclimacteric stages when high ACS activity was observed. Overall changes in ACS protein abundance were similar to the changes observed for ACO activity.

Transcriptional Profiling Reveals Similarities between ACS and Yang Cycle Gene Expression

The burst of ethylene synthesis is a hallmark of climacteric tomato fruit ripening. Surprisingly, postclimacteric development with low ethylene production was accompanied by high ACS activity and abundance and an accumulation of ACC. To better understand the regulation of ethylene biosynthesis and Met recycling at the transcriptional level, expression profiles of ACO, ACS, SAM5, and Yang cycle genes were studied throughout tomato fruit development, ripening, and postharvest storage by real-time quantitative PCR (qPCR). These expression profiles were compared with the recently obtained RNA sequencing data of the Tomato Genome Consortium for tomato fruit of the same maturity stages for both the greenhouse cultivar Heinz 1706 and the old species Solanum pimpinellifolium (Tomato Genome Consortium, 2012). Figure 6 shows the expression profiles of all known ACO isoforms in tomato. The results confirm previous reports on ACO expression patterns visualized by northern blotting but also revealed some differences (Barry et al., 1996; Nakatsuka et al., 1998). ACO2 and ACO4 transcript levels are highest during system 1, whereas in a previous report ACO2 transcripts were not detected (Barry et al., 1996). ACO1 was moderately expressed during preclimacteric development. During the transition stage ACO1 and ACO3 transcript levels increased, whereas the expression levels of ACO2 and ACO4
dropped. ACO1 was the only ACO gene still expressed at elevated levels in the postclimacteric stage, likely contributing to the low rate of ethylene synthesis observed in these fruit. In parallel with ACO1 expression, ACO abundance (Fig. 5A), ACO activity (Fig. 4A), and ethylene production (Fig. 2A) continuously declined during fruit storage. ACO3 expression dropped from the light orange stage on to a very low level that was maintained during postharvest storage. The expression level of ACO4 was very low during ripening, contrary to the report by Nakatsuka et al. (1998). The yet uncharacterized expression profile of ACO5 showed a similar expression pattern as ACO3 with elevated levels during ripening. In general, the RNA sequencing data matched the expressions profiles for all ACO isoforms.

Figure 7 shows the expression profiles of the ACS gene family. The expression profile of ACS1 (A and B together) fluctuated throughout fruit development. During preclimacteric development, ripening, and again during postharvest storage, ACS1 was transiently upregulated. Transcripts of ACS2 and ACS4 were not detected during initial development, but expression strongly increased during the autocatalytic ethylene production phase of system 2. Expression of both genes peaked during the breaker stage along with the peak in ACS abundance (Fig. 5B) and activity at this stage (Fig. 4B). ACS2 and ACS4 transcript levels declined during further ripening and peaked again during the postclimacteric stages matching ACS activity. ACS6 was expressed during early fruit development, but mRNA levels quickly decreased. During ripening, ACS6 was not expressed, but transcript levels gradually increased during postharvest storage.

Transcripts of the other four ACS members (ACS3, ACS5, ACS7, and ACS8) were not detected at any stage. A phylogenetic relationship analysis of all ACS genes in tomato revealed that the fruit ripening-related ACS genes group together, whereas the remaining four nonfruit-ripening specific members that were not expressed also grouped (Supplemental Fig. S2). The RNA sequencing data matched the qPCR expression profiles for all ACS genes and confirmed high expression of ACS2 and ACS4 at the red stage. No RNA reads were found for ACS3, ACS5, ACS7, and ACS8.

It has been proposed that high rates of ethylene biosynthesis in climacteric fruit are supported by recycling of the ethylene precursor Met via the Yang cycle (Baur and Yang, 1972). To test this hypothesis, we analyzed transcript abundance of the known Yang cycle genes MTN, MTK, ARD1, ARD2, MTI, and DEP in relation to ethylene biosynthesis in tomato fruit (Fig. 8). MTN, MTK, ARD1, and ARD2 showed a coordinated expression pattern. During fruit development, expression levels were low and increased just prior to ripening at the mature green stage. Around the orange stage, transcript levels dropped slightly before increasing.

Figure 6. Gene expression (relative copies) of ACO1 (A), ACO2 (B), ACO3 (C), ACO4 (D), and ACO5 (E) during fruit development, ripening, and postharvest storage. Error bars represent the so of the mean of six biological replicates. See Figure 2 for maturity stage annotations. For comparison, normalized RNA sequencing data (secondary y axis) for defined maturity stages from cv Heinz 1706 (light gray circles) and S. pimpinellifolium (open circles) were included.
to even higher levels during postharvest storage. These expression profiles showed a striking similarity to the expression profiles of ACS1, ACS2, and ACS4 and to ACS activity. In contrast, the recently identified phloem-specific Yang cycle genes DEP and MTI (Pommerrenig et al., 2011) did not show a comparable pattern. Both genes were mainly expressed in young developing fruit and showed a moderate constitutive expression during ripening and postharvest storage. Again the RNA sequencing data confirmed the expression profiles for all Yang cycle genes except for MTN, for which no RNA reads were found, although the qPCR results provided in this study show a strong up-regulation.

Although SAMS is not exclusively used for ethylene synthesis, the production of SAM from Met is an important requirement in the ethylene pathway. Four SAMS genes (SAMS1–SAMS4) were identified in tomato, for which the expression profiles during fruit development are shown in Supplemental Figure S3. SAMS1 and SAMS4 showed a similar profile with only a high expression during fruit development. SAMS2 and SAMS3 were also highly expressed during fruit development, but SAMS2 was also up-regulated during postharvest storage. It is worth mentioning that none of the SAMS genes were expressed at high levels during climacteric ripening, although SAM levels peaked during these stages. The RNA sequencing data also matched the SAMS expression profiles in general. Despite technical and cultivar differences, the RNA sequencing data of the Tomato Genome Consortium (2012) largely confirmed all measured expression profiles. This indicates that the observed profiles are representative for modern greenhouse cultivars (cv Bonaparte and Heinz 1706) as well as for the old species S. pimpinellifolium, and likely for tomato fruit in general.

Effect of Harvest on Ethylene Biosynthesis

When fruit have reached their red ripe stage and ethylene biosynthesis declines, ACS activity and expression, together with the expression of Yang cycle genes, increased. As these changes coincided with harvest, we decided to investigate whether they occurred as a result of fruit detachment. Figure 9 shows the effect of harvesting red tomatoes on fruit ethylene production, related enzyme activities, and metabolites. Detached red fruit showed a reduced level of free ATP (Fig. 9A), indicating that energy supply was more limiting in detached fruit. Ethylene profiles for both attached and detached fruit showed the characteristic decline in ethylene production for postclimacteric fruit (Fig. 9B). However, fruit left on the plant had a significantly higher ethylene production rate than fruit removed from the vine. This was also reflected by a higher ACO activity (Fig. 9C). It is interesting to note that red fruit kept on the plant did not show an increase in ACS activity, as was observed in detached fruit (Fig. 9D). The increase in ACS activity, together with a lower consumption by ACO, resulted in an ACC content that was significantly higher in detached fruit as compared with attached fruit (Fig. 9E). Differences in MACC levels were also observed, albeit less pronounced (Fig. 9F). As a consequence of the increased ACS activity in detached fruit, SAM levels...
were slightly lower (Fig. 9G). MTA levels, on the other hand, did not differ significantly (Fig. 9H). Although one would expect a higher MTN activity to cope with the additional MTA coming from the high ACS activity, MTN activity was significantly lower in detached fruit (Fig. 9I). Perhaps other pathways (e.g., polyamine biosynthesis) might have reduced their MTA input because of the detachment. Nonetheless, the findings indicated that MTA levels are tightly regulated, independent of harvest. Taken together, these results showed that harvesting red tomatoes triggers ACS activity and, at the same time, inhibits ACO activity with a concomitant effect on ethylene biosynthesis metabolites.

Figure 10 shows expression profiles of some ACO, ACS, and Yang cycle genes. ACO1 expression was slightly higher in attached fruit concomitant with elevated ethylene production of fruit on the vine (Fig. 10A). Both attached and detached fruit showed upregulation of ACS2 within 5 d after harvest, but this was more pronounced in detached fruit (Fig. 10B). Expression of MTN, MTK, ARD1, and ARD2 was also upregulated in postclimacteric fruit (Fig. 10, C–F). Again, tomatoes that were stored off the plant showed a more pronounced increase of transcripts. Note that the higher MTN expression in detached fruit did not result in an increased MTN activity (Fig. 9D). Overall, these profiles suggest that harvesting red fruit promotes ACS activity through an elevated expression of ACS2 and inhibits ACO activity and ethylene production. At the same time, the expression of Yang cycle genes is elevated, activating the recycling part of the ethylene pathway likely to metabolize the ACS side-product, MTA.

Ethylene Sensitivity Is Lost during Postclimacteric Fruit Development

System 1 and system 2 have previously been characterized as autoinhibitory and autocatalytic mechanisms
to regulate ethylene biosynthesis. To test whether postclimacteric tomatoes with a declining ethylene production are susceptible to ethylene-dependent feedback regulation, experiments were carried out with exogenously applied ethylene and with 1-methylcyclopropane (1-MCP), an inhibitor of ethylene perception. Application of the ethylene-releasing compound ethephon to red fruit transiently increased ethylene production within 4 d whereas 1-MCP transiently reduced ethylene biosynthesis compared with control fruit (Fig. 11). At 8 d and later, control fruit had an equal ethylene production compared with treated fruit, suggesting only a temporal effect. The overall decline in ethylene production observed in the postclimacteric stage could not be overcome by any of the treatments, indicating that down-regulation of ethylene biosynthesis after fruit maturation was developmentally programmed and only moderately ethylene sensitive.

**PCA Reveals Profile Similarities**

To further analyze this large and complex data set, principal component analysis (PCA) was performed on all parameters measured. The correlation loadings plot of the first two PCs is shown in Supplemental Figure S4. PC1 explained 33%, and PC2 explained 25% of the total variance. Some variables were closely clustered, indicating that those metabolites/proteins/gens showed a similar profile. ACC and MACC grouped and were associated with postharvest storage. Fruit firmness and respiration (CO₂) were associated more with small developing fruit. The PCA also
confirmed that the Yang cycle genes (MTN, MTK, ARD1, and ARD2) correlated well with ACS activity. ACO activity correlated with the expression of ACO1, ACO3, and ACO5 whereas ethylene production correlated strongly with SAM and MTA content and with ACS2 and ACS4 expression, and was associated mainly with breaker to orange fruit.

**DISCUSSION**

**Regulation of Postclimacteric Ethylene Biosynthesis Is Distinct from System 1 and System 2 Ethylene Production**

A targeted systems biology approach allowed us to characterize the ethylene biosynthesis pathway and the Yang cycle of developing tomato fruit in depth. Based upon the results described above and previous findings from literature, the regulation of postclimacteric ethylene biosynthesis was found to be different from the autocatalytic phase (system 2) and different from the autoinhibitory phase (system 1). The postclimacteric ethylene production is characterized by a gradual decline in ethylene biosynthesis when pink-red fruit have reached their maximum in autocatalytic ethylene production. It is essential that tomato fruit stop the autocatalytic rise when the ripening process is nearly accomplished. Additional ethylene from this point on will enhance senescence and fruit decay. It may be advantageous that red fruit delay their senescence phase to maximize the time available for seed dispersal. The exact mechanism by which autocatalytic ethylene production is shut down is only poorly understood. We found that the regulatory mechanisms by which ethylene biosynthesis is shut down operate at the level of ACO1 transcript and ACO protein abundance. On the other hand, ACS gene expression and activity increase again during postharvest storage of tomato. Enhanced ACS activity, together with reduced ACO activity, results in an
accumulation of ACC indicating that ACO is the rate-limiting step of postclimacteric ethylene biosynthesis.

How ACO and ACS expression is controlled during the postclimacteric phase remains unknown. Elucidation of the initiation of fruit ripening and the associated climacteric ethylene production can provide some clues. A few key transcriptional regulators were identified as essential activators of the ripening process. RIN-MADS is a MADS-box transcription factor that binds to the promoter of ACS2, ACS4, and other ripening-associated genes. Therefore, RIN is a crucial and early regulator of fruit ripening (Vrebalov et al., 2002; Fujisawa et al., 2011). Perhaps ACS2 expression during postclimacteric ripening could be induced by a sustained activity of RIN-MADS. HB-1 encodes a homeodomain-zip homeobox transcription factor that directly interacts with the promoter of ACO1 and is responsible for the activation of ACO1 expression during ripening (Lin et al., 2008). Loss of HB-1 activity might terminate autocatalytic ethylene production by reducing ACO1 expression. If and how these regulators shut down ethylene biosynthesis remains to be investigated. It is also plausible that ethylene response factors (ERFs) are responsible for the direct (interaction with ACO and/or ACS promoter regions, like ERF2) or indirect (through deactivating the expression of the regulators mentioned above) shut down of ethylene biosynthesis. The ERF gene AP2a was identified as a negative regulator of ethylene biosynthesis and was also found to be ethylene induced (Karlova et al., 2011). Repression of AP2a expression resulted in an overproduction of ethylene (Chung et al., 2010; Karlova et al., 2011). Down-regulation of ethylene biosynthesis by ethylene-induced ERFs would mean that ethylene turns down its own production during postclimacteric ripening, similar to system 1 and system 2 ethylene biosynthesis. An overview summarizing the main ethylene-related features of system 1 and system 2 and of the regulatory mode of postclimacteric ripening described here is given in Figure 12 and Supplemental Table S1.

The Termination of Climacteric Ethylene Production Is Developmentally Programmed

Mature green tomatoes treated with 1-MCP show a reduced climacteric ethylene production and do not ripen like untreated fruit (Hoeberichts et al., 2002; Tassoni et al., 2006). When treated with ethylene, fruit show an increased ethylene production and ripen faster (Sims, 1969). With either treatment, climacteric ethylene biosynthesis was severely altered. By contrast, postclimacteric red fruit that were treated with ethephon or 1-MCP did not show such a pronounced alteration in ethylene production. The decline in ethylene production was not overcome by ethephon nor inhibited by 1-MCP, except for a temporal and minor effect 4 d after treatment. This temporal effect of 1-MCP-treated red ripe tomatoes was also observed by Hoeberichts et al. (2002). How the change in ethylene sensitivity is achieved in postclimacteric tomatoes remains to be elucidated. Ethylene sensitivity is inversely proportional to the amount of ethylene receptors (Hua and Meyerowitz, 1998; Tieman et al., 2000). This means that a high receptor abundance lowers ethylene sensitivity and vice versa. During climacteric ripening of tomato, the expression of ethylene receptors NR (ETR3), ETR4, and ETR6 is induced (Lashbrook et al., 1998, Kevany et al., 2007). Receptor abundance may also be regulated at the level of protein stability (Kevany et al., 2007). However, it remains to be investigated if receptor abundance is key to the reduced ethylene sensitivity observed during postclimacteric development. Regulation at other ethylene signaling components is also conceivable. In any case, the decline in ethylene biosynthesis observed during postclimacteric ripening appears to be developmentally programmed and only moderately influenced by ethylene signaling.
The Yang Cycle Genes Are Coordinately Expressed and Show a Distinct Up-Regulation during Postclimacteric Ripening

The Yang cycle genes MTN, MTK, ARDI, and ARD2 showed very similar expression profiles throughout tomato fruit development, ripening, and postharvest storage. By contrast, MTK and ARDI were mainly expressed in a constitutive manner. Arabidopsis AMTI and AtDEP were described as being specifically expressed in the phloem (Pommerrenig et al., 2011). It was shown that the vascular bundle tissue of tomato fruit pericarp showed a more complex transcriptome profile compared with overall pericarp tissue, as used in this study (Matas et al., 2011). Possibly tomato fruit pericarp possesses other enzymes that catalyze these steps of the Yang cycle than the ones analyzed, although a database search did not reveal additional homologous genes. Perhaps certain Yang cycle metabolites are being translocated between the pericarp tissue and vascular cells for further processing. Recently, it was shown in the Arabidopsis double mtn/mtk mutant that efficient recycling of MTA by the Yang cycle is essential for vascular tissue development (Waduwara-Jayabahu et al., 2012).

The gene expression profiles of MTN, MTK, and ARD show a basal expression during preclimacteric fruit development and an up-regulation with the onset of ethylene production at the mature green stage. This is in accord with the long-proposed function of the Yang cycle to preserve the reduced sulfur of MTA, the by-product of ACS during high rates of ethylene biosynthesis (Baur and Yang, 1972; Bürstenbinder et al., 2007). Nevertheless, these Yang cycle genes show their highest expression during the postclimacteric phase when ethylene production continuously declined. High Yang cycle activity coincides with the up-regulation of ACS genes and ACS enzyme activity. As only little ethylene is produced postclimacterically, the signal that controls the Yang cycle may come from the ACS products ACC or MTA. It was previously described that MTN1 protein abundance and MTN enzyme activity in Arabidopsis were induced by MTA (Bürstenbinder et al., 2010). In rice (Oryza sativa), it was shown that OsMTN expression and MTN activity increased in the youngest internode upon submergence, also linking it to ethylene and ACS (Rzewsuki et al., 2007). However, early experiments showed that breeder tomato pericarp discs infiltrated with the ACS activity inhibitor aminoethoxyvinyl-Gly did not alter MTN activity (Wang et al., 1982). Independent of ethylene synthesis or ACS activity, it appears to be crucial to maintain MTA levels within a narrow concentration range. It was shown that loss of MTA metabolism in Arabidopsis mth knockout plants causes morphological disorders in Arabidopsis (Waduwara-Jayabahu et al., 2012). This evidence points toward a crucial role for MTA in regulating the initial step of the Yang cycle. However, it still remains to be elucidated how this is achieved and how MTN and ARD genes are coordinately regulated.

One possibility would be that their respective substrates accumulate with higher MTN activity resulting in an independent, yet coordinated, regulation of the Yang cycle. Alternatively, a common transcription factor could be activated that coordinately regulate MTN, MTK, ARDI, and ARD2 genes in tomato fruit. The data presented here clearly indicate that ethylene is not the crucial signal to regulate the Yang cycle as originally stated by Baur and Yang (1972). Rather, the regulation of the Yang cycle in tomato pericarp is linked to ACS activity and/or MTA levels.

Fruit Detachment Induces ACS and Inhibits ACO

Detachment of fruit from the mother plant is a serious intervention. Fruit are cut off from nutrition and water, causing stress. Resources and energy need to be preserved as the fruit suddenly has to switch from a heterotrophic metabolism to an autotrophic metabolism. This will ultimately result in a lower content of free ATP, as observed in detached tomato fruit. Previous studies demonstrated that detached preclimacteric tomato fruit have a normal climacteric ethylene production rate (Nakatsuka et al., 1997). Preclimacteric harvested apples show a higher ethylene production rate compared with attached fruit (Sfakiota and Dilley, 1973; Lin and Walsh, 2008) whereas harvested melon (Cucumis melo) fruit did the opposite (Bower et al., 2002). Little information is available about ethylene production in fruit detached post ripening. Our results show that the decline in ethylene production during postclimacteric ripening is enhanced when the fruit is harvested. This reduction is caused by an inhibition of both ACO expression and activity. Despite of a reduced ACO activity, ACS expression and activity increase following detachment. It is known that local wounding of the calyx and fruit water stress by dehydration can induce ACS expression in detached persimmon fruit (Nakano et al., 2003). Harvested tomato fruit are also confronted with water loss during storage (Hertog et al., 2004), which might in turn induce ACS expression and activity. Or perhaps another “tree factor” is involved (Sfakiota and Dilley, 1973). Along with ACS activity, expression of Yang cycle genes also increased in detached fruit. This is in accord with the view that regulation of the Yang cycle is closely linked to ACC/MTA production.

CONCLUSION

A targeted systems biology approach revealed a novel mode of regulation of ethylene biosynthesis, which sets in when tomato fruit have reached their climacteric maximum around the pink-red stage. It is characterized by a decline in ethylene production that appears to be developmentally programmed and that is regulated at the level of ACO. Harvest of red tomatoes hastens the decline in ethylene production and at the same time promotes the expression and activity of
ACO. Increased ACS activity causes an influx of MTA into the Yang cycle, which is highly active during postclimacteric ripening, revealing a close link between ACC synthesis and Yang cycle regulation. Finally, we observed that Met recycling from MTA is important in the Yang cycle, which is highly active during postclimacteric ripening, revealing a close link between ACC synthesis and Yang cycle regulation. Furthermore, we investigated the role of Yang cycle genes in regulating Yang cycle activity and found that ACS activity is crucial for Yang cycle regulation.

**MATERIALS AND METHODS**

**Plant Material**

Tomato plants (*Solanum lycopersicum* 'Bonaparte') were grown in a greenhouse at the Research Station for Vegetable Production of Hooigraten (Belgium). Plants were cultivated hydroponically on rock wool substrate under natural daylight with controlled humidity (70%) and temperature (23°C/21°C d/night). Fruit of different developmental and ripening stages were harvested (August 2009), transported to the lab, analyzed for ethylene production and fruit quality attributes (color and firmness), and further processed the same day (Van de Poel et al., 2010). Ten independent fruits of each developmental stage were selected, and pericarp tissue of individual fruit was flash-frozen in liquid nitrogen, crushed with a Grindomixer (Retsch) and stored at −80°C. Subsequently, a postharvest experiment was conducted by harvesting red ripe tomato fruit and storing them for 12 d at shelf life conditions (18°C and 80% relative humidity) with regular sampling (10 fruit each time) as described above.

**The Effect of Harvest on Ethylene Biosynthesis**

In a separate experiment, the effect of harvest on ethylene biosynthesis was investigated by comparing red ripe tomato fruit attached and detached from the plant in time (June 2010). For the detached fruit, red ripe tomatoes were harvested from the greenhouse (as described above) and stored for 14 d at shelf life conditions (18°C and 80% relative humidity). At the same time, red ripe fruit that remained attached to the vine were marked and kept on the plant for the same 14 d. Both attached and detached fruit were sampled simultaneously several times during the same storage period (10 independent fruit for each sampling time/treatment). Ethylene production was assessed the same day of sampling and pericarp tissue of individual fruit was subsequently flash frozen in liquid nitrogen, crushed, and stored at −80°C.

**1-MCP and Ethephon Treatment of Red Harvested Tomatoes**

Red ripe tomatoes were harvested and treated for 24 h at 18°C in airtight containers with 5 μL L⁻¹ 1-MCP (SmartFresh, AgroFresh) or 300 μL of ethephon Classic (a kind gift from Bayer CropScience N.V.). The ethephon solution was adjusted to pH 6.0 to facilitate rapid ethylene release. Control fruit were also kept at 18°C in similar sealed containers during the experimental period. Subsequent sampling was done on 10 fruits per time point per treatment. Ethylene production was measured, and pericarp tissue was flash frozen in liquid nitrogen, crushed, and stored at −80°C.

**Ethylene and Metabolite Quantification**

Fruit ethylene production was quantified after 1 h incubation according to the procedure described by Bulens et al. (2011). Fruit respiration rate was quantified by measuring CO₂ production simultaneously with ethylene production by gas chromatography (CompactGC, Interscience). ACC was extracted and quantified by the Lizaeda and Yang (1979) method recently optimized by Bulens et al. (2011). MACC was obtained after acidic hydrolysis according to Hoffman et al. (1982) and also recently updated by Bulens et al. (2011). SAM was extracted and quantified by capillary electrophoresis (P/ACE MDQ, Beckman Coulter) with UV detection according to Van de Poel et al. (2010). Met content was measured by gas chromatography-mass spectrometry (Agilent) as described by Oms-Oliu et al. (2011). MTA and ATP were quantified by HPLC after chloroacetaldehyde derivatization according to Bürrstenbinder et al. (2007).

**Protein Extraction and in Vitro Activity**

ACO was extracted and in vitro activity determined according to Verderidis and John (1991) updated by Bulens et al. (2011). The incubation buffer consisted of a 100-μM Tris buffer, pH 8.0, instead of the described 400-μM 3-(N-morpholino)propanesulfonic acid buffer, pH 7.2. The incubation time was optimized to 15 min. ACS was extracted and in vitro activity determined exactly as described by Bulens et al. (2011). MTN was extracted with a cold 50 mM of potassium phosphate, pH 7.2, from 200 mg of crushed frozen tissue. The mixture was vortexed vigorously and incubated on ice for 10 min. Subsequently, it was centrifugated at 18,320g for 30 min at 4°C. The supernatant was collected for the in vitro activity assay. The enzymatic MTN activity was measured spectrophotometrically according to Durn et al. (1994), updated by Bürrstenbinder et al. (2010). Briefly, 15 μg of total protein yield was used together with 50 mM of potassium phosphate buffer, pH 7.2, 1 mM MTA, and 2.5 mM of iodonitrotetrazolium chloride (prepared fresh). This mixture was incubated for 2 h with 0.2 units of xanthine oxidase (Grade III), and absorption was measured periodically at 470 nm with a spectrophotometer (SpectraMax M2, Molecular Devices).

**Western Blotting**

Polyclonal antibodies were developed (GenScript, GE Healthcare) against a consensus epitope for four different ACO isoforms (ACO1, ACO2, ACO3, and ACO6) and a consensus epitope for four ACS isoforms (ACS1, ACS2, ACS4, and ACS6: LADPGDAFLVP). The total protein content of ACO and ACS extracts was determined by the Bradford assay (Bradford, 1976). For ACO and ACS, 15 and 25 μg of total protein content, respectively, was mixed with 10X SDS-sample buffer (Laemmli buffer), denatured for 5 min at 95°C, and loaded on a 12-well Criterion XT 4% to 12% Bis-Tris gel (Bio-Rad). SDS-PAGE was performed for 1 h at 180 V in an XT-MES buffer (Bio-Rad). Subsequent electro blotting was carried out for 1 h and 20 min at 100 V on a Protran nitrocellulose membrane (Whatman, Maidstone) or a polyvinylidene difluoride membrane (GE Healthcare) in the presence of transfer buffer (25 mTris, 140 mGly, 20% [v/v] methanol). The membrane was blocked for 1 h in TBS-T (Tris-buffered saline plus Tween 20; 25 mM Tris, 125 mM NaCl, and 0.1% [v/v] Tween 20) containing 5% milk powder for ACS and 5% bovine serum albumin for ACO. After blocking, the membrane was incubated overnight at 4°C with primary antibody solution (1/1000 anti-ACO AB in TBS-T with 5% milk powder or 1/100 anti-ACS AB in TBS-T with 5% bovine serum albumin). Subsequently, the membrane was washed 5 times for 5 min in TBS-T. Secondary antibody (1/2000 anti-rabbit-horseradish peroxidase-linked AB; Cell Signaling Technologies, Inc.) incubation was carried out for 2 h at 4°C in the presence of TBS-T with 5% milk powder or 5% bovine serum albumin. Again, the membrane was washed. Enhanced chemiluminescence was performed with Pierce ECL western-blotting substrate (Thermo Fischer Scientific), and band intensity was detected with the ImageQuant LAS4000 system (GE Healthcare).

**RNA Extraction and Reverse Transcription-qPCR**

All procedures described below are in accordance with the Minimum Information for Publication of Quantitative Real-Time PCR Experiments guidelines where applicable (Bustin et al., 2009). Total RNA was extracted with hot (65°C) RNA extraction buffer (55 mM 0.5 ethyltrimethyl ammonium bromide, 1.34 m NaCl, 25 mM EDTA, 2% [w/v] polyvinylpyrrolidone, 2% [v/v] β-mercaptoethanol, 0.05% [v/v] spermide) from 4 g of frozen crushed tissue. The mixture was heated for 2 min at 65°C and subsequently homogenized for 2 min and further incubated at 65°C for 15 min. An equal volume of chloroform:isoamyl alcohol (24:1, v/v) was added, shaken vigorously, and subsequently centrifuged at 3,090g for 5 min at 4°C. The supernatant was discarded. The RNA pellet was washed with 0.5 μL of 7.5 m LiCl overnight at 4°C. The mixture was centrifugated at 14,000g for 30 min at 4°C, and the supernatant was discarded. The RNA pellet was washed with 300 μL of 70% cold ethanol and centrifuged again at 14,000g for 10 min at 4°C. The ethanol supernatant was neutralized with 3 M sodium acetate and ethanol leftovers were vaporized under vacuum for 15 min at room temperature. The pellet was dissolved in 75 μL of an RLT buffer and subsequent purification, and DNA removal was done with the Qiagen RNeasy Plus Mini Kit according to the manufacturer's protocol (Qiagen GmbH). RNA integrity was checked on a 1% agarose gel
stained with ethidium bromide and RNA content was quantified spectroscopically in a 384-well microtiter plate (UV-star, Greiner Bio-One GmbH) by measuring absorption at 260 nm. One microgram of the purified RNA was reverse transcribed into complementary DNA (cDNA) by the Quant iTect Rev. Transcription Kit (Qiagen) according to the manufacturer’s protocol and subsequently stored at ~80°C.

The reverse transcription-qPCR reaction consisted of forward and reverse primer (3.75 μM), reverse transcription-template, water, and Absolute QPCR SYBR Green mix (ABgene, Ltd.). Primers and their properties are listed in Supplemental Table S2. qPCR was performed with the Rotor-Gene Q (Qiagen) and molecular measurements. B.V.d.P., I.B., M.L.A.T.M.H., S.V., M.S., and G. Pittoors from Pitoma BVBA are acknowledged for supplying plant material. J. Shukhtina (KU Leuven) is acknowledged for her help with certain experiments. R.D., M.W., S.V., Y.O., and R.H. helped with specifying the primers.

J.K., E.W., M.P.D.P., M.S., B.M.N., and A.H.G. designed the research. I.B., B.V.d.P., S.V., and G. Pittoors from Pitoma BVBA are acknowledged for supplying plant material. J. Shukhtina (KU Leuven) is acknowledged for her help with certain experiments. R.D., M.W., S.V., Y.O., and R.H. helped with specifying the primers.

Statistical Analysis, PCA, and Trend Visualization
Statistical differences were analyzed with the one-way ANOVA procedure using the Statistical Analysis Software (SAS Enterprise Guide 4.2; SAS Institute, Inc.). Confidence intervals were set at 95%. PCA was performed with The Unscrambler X 10.1 (CAMO software AS). Graphs were made with Origin 8. Smoothing splines were used and ran for 1 million generations.

Phylogenetic Analysis
Phylogenetic analyses were executed with Mr. Bayes 3.2 freeeware (Ronquist and Huelsenbeck, 2003) using cDNA sequences of all nine ACS isoforms (for annotation see Supplemental Table S1). Default settings of the 4by4 Nucmodel were used and ran for 1 million generations.

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