Metabolic engineering of tomato fruit organic acid content guided by biochemical analysis of an introgression line

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Organic acid content is regarded as one of the most important quality traits of fresh tomatoes. However, the complexity of carboxylic acid metabolism and storage means that it is difficult to predict the best way to engineer altered carboxylic acid levels. Here we have used a biochemical analysis of a tomato introgression line with increased levels of fruit citrate and malate at breaker stage to identify a metabolic engineering target that was subsequently tested in transgenic plants. Increased carboxylic acid levels in introgression line 2-5 were not accompanied by changes in the pattern of carbohydrate oxidation by pericarp discs or the catalytic capacity of TCA cycle enzymes measured in isolated mitochondria. However, there was a significant decrease in the maximum catalytic activity of aconitase in total tissue extracts suggesting that a cytosolic isoform of aconitase was affected. To test the role of cytosolic aconitase in controlling fruit citrate levels, we analysed fruit of transgenic lines expressing an antisense construct against SlAco3b, one of the two tomato genes encoding aconitase. A GFP-fusion of SlAco3b was dual targeted to cytosol and mitochondria, while the other aconitase, SlAco3a was exclusively mitochondrial when transiently expressed in tobacco leaves. Both aconitase transcripts were decreased in fruit from transgenic lines and aconitase activity was reduced by about 30% in the transgenic lines. Other measured enzymes of carboxylic acid metabolism were not significantly altered. Both citrate and malate levels were increased in ripe fruit of the transgenic plants and, as a consequence, total carboxylic acid content was increased by 50% at maturity.
**Introduction**

Tomato is an important food crop of high economic value and represents a model species for fleshy fruit physiology and ripening (Giovannoni, 2004; Mueller, 2009). The breeding history of tomato has been dominated by a focus on traits that benefit the grower such as yield, storage characteristics and field performance (Schuch, 1994; Giovannoni, 2006; Cong et al., 2008). As a result there has been an unintentional loss of consumer quality traits such as flavour and nutritional value and this has focussed recent interest on the molecular genetics of such traits (Giovannoni, 2001; Causse et al., 2002; Causse et al., 2004; Fraser et al., 2009; Mounet et al., 2009; Enfissi et al., 2010; Centeno et al., 2011). The accumulation of a range of soluble metabolites is critically important for both flavour and nutrition. Tomato fruit undergo substantial changes in their metabolite content and composition during ripening (Carrari et al., 2006). Fruit flavour is influenced both by volatile and non-volatile metabolites (Buttery et al., 1987; Goff and Klee, 2006; Carli et al., 2009). Of the non-volatile metabolites, the balance between sugars and acidic compounds is of major importance for flavour (Tieman et al., 2012). Perceived flavour of tomato fruit is a complex issue and simple associations of metabolites with flavour traits do not always hold true. Nevertheless, a network analysis of several tomato genotypes demonstrated a strong correlation between tomato fruit flavour and the main acidic metabolites - carboxylic acids, glutamate and aspartate (Carli et al., 2009).

The breeding or engineering of improved flavour by increasing the amount of specific metabolites in ripe fruit requires an understanding of the biochemical and molecular factors that regulate their accumulation during the ripening process. In this study, we focused on the accumulation of citrate and malate, the most abundant of the acidic metabolites in tomato fruit. Carboxylic acids are a major component of the osmotic potential that drives cell expansion through water uptake in the expansion phase of fruit growth (Liu et al., 2007). The concentrations of citrate and other carboxylic acids fall during this expansion phase as the cell contents are diluted (Baxter et al., 2005; Carrari et al., 2006). However, during the final stages of ripening the level of citrate (and to a lesser extent other carboxylic acids) increases again such that it is present at high abundance in the ripe fruit. It is not known how these changes in organic acid levels are brought about. The maximal catalytic activities of enzymes of the tricarboxylic acid cycle generally decline during fruit development and there are no pronounced changes in activities during the latter stages of ripening that correlate with the rise in organic acid levels (Steinhauser et al., 2010).

Regulation of metabolite levels is a complex issue. At the most basic level, the amount of a metabolite will change because of a difference between influx into that metabolite pool and efflux from it (Kruger and Ratcliffe, 2009). In the case of citrate, for example, one could envisage that a change in the balance of flux through the citrate synthase and aconitase reactions could be responsible for a change in citrate levels. However, the situation is complicated because citrate is synthesised in the mitochondrion but accumulates in the vacuole (Martinoia et al., 2007). Thus, transport of citrate between these subcellular compartments will also have a bearing on its rate of accumulation (Shiratake and Martinoia, 2007). Export from the mitochondrion is by counter-exchange with other carboxylic acids which are imported. Therefore the capacity of other parts of the TCA cycle will affect the rate of export of citrate. Active uptake into the vacuole is ultimately dependent on the tonoplast membrane potential generated by the proton-pumping V-type ATPase, although citrate transport appears to occur by facilitated diffusion rather than active uptake in tomato fruit (Oleski et al., 1987). Given that TCA cycle flux is critical for synthesis of ATP, one can see the extent to which carboxylic acid metabolism as a whole and citrate accumulation are intertwined. Accumulation in the vacuole is also a function of both influx and efflux of citrate from the vacuole (Shimada et al., 2006) and subsequent metabolism by cytosolic isoforms of aconitase.
and isocitrate dehydrogenase. Given this complexity and the variety of flux modes within carboxylic acid metabolism (Sweetlove et al., 2010), it is not obvious what the best strategy for engineering increased accumulation of carboxylic acids in fleshy fruits would be. This is reflected in the range of enzymes that have been proposed to control fruit citrate accumulation including PEP carboxylase (Guillet et al., 2002), PEP carboxykinase (Famiani et al., 2005), citrate synthase (Sadka et al., 2000) and aconitase (Sadka et al., 2000; Degu et al., 2011).

To identify a suitable metabolic engineering strategy for increasing carboxylic acid content of tomato, we undertook a detailed biochemical study of an introgression line of tomato (Eshed and Zamir, 1995). A specific line with an introgressed segment from *Solanum pennellii* on chromosome 2 (IL2-5) was identified that showed reproducible increases in citrate and malate during the later stages of fruit development but had minimal changes in gross developmental characteristics such as fruit size and number. Although the introgressed segment contained many genes, and metabolites other than carboxylic acids were altered, we reasoned that a focused metabolic analysis would identify which of the proposed mechanisms for controlling fruit citrate accumulation was responsible for the increased citrate and malate content in the introgression line. The effectiveness of this mechanism as a viable target for metabolic engineering of carboxylic acid content could then be tested in transgenic plants.
Results

Fruit of introgression line 2-5 have increased malate and citrate content during development.

To identify a suitable line from the *Solanum lycopersicum* x *Solanum pennellii* introgression population (Eshed and Zamir, 1995), we grew a subset of lines that were free from major phenotypic changes (Gur et al., 2004) and measured fruit carboxylic acid content in pericarp tissue using $^1$H NMR. We wished to identify lines in which carboxylic acids were increased at several stages of fruit development prior to the final ripe stage. This was because the biochemical changes that influence accumulation of carboxylic acids during ripening are likely to be set prior to the final stage of maturation.

Based on our analysis, we focused on one introgression line, IL2-5, in which fruit citrate and malate content was increased (Fig. 1). Fruits were analysed at three developmental ages (30, 40 and 55 days after anthesis (DAA)). Tomatoes at 40 DAA were at breaker stage and by 55 DAA they were fully ripe. In comparison to the *Solanum lycopersicum* M82 parent line, citrate was significantly increased at each of the developmental ages, with the difference between IL2-5 and M82 increasing during development such that ripe fruit of IL2-5 contained 60% more citrate than those of M82. Malate levels in IL2-5 were also significantly increased by a similar degree at 30 and 40 DAA but in ripe fruit there was no difference compared to M82. Fumarate levels showed reciprocal changes to malate, with decreases in IL2-5 at 30 and 40 DAA, although the absolute amount of fumarate was a factor of one thousand less than malate. Other carboxylic acids were not quantifiable by $^1$H NMR of fruit extracts. These metabolite changes were not due to an altered rate of development of IL2-5 fruit because the main changes in carotenoid content, a key indicator of fruit developmental stage, were coincident in M82 and IL2-5 fruit (Supplemental Fig. 1).

To assess whether other parts of the central metabolic network were also altered in IL2-5 fruits we determined the content of amino acids. Because amino acid synthesis draws on precursors from a number of different sectors of central metabolism, they are a useful indicator of the state of the central metabolic network. As one might expect given the number of potential genetic changes due to chromosomal introgression, changes in a range of amino acids were observed (Fig. 2). Of the 19 measured amino acids, 10 were significantly increased in IL2-5 fruit. Several of these (glutamate, threonine and methionine) are either directly connected to carboxylic acid metabolism or draw on carboxylic acid metabolism as a source of carbon skeletons for their biosynthesis. However, in addition there were also increases in amino acids that are synthesised from precursors from glycolysis (alanine, serine, glycine, valine, leucine, isoleucine) and the oxidative pentose phosphate pathway (phenylalanine).

Carbohydrate oxidation fluxes in IL2-5 fruit

To assess whether the increases in citrate and malate were accompanied by major changes in flux through the central network of carbon metabolism, we incubated discs cut from the pericarp of IL2-5 and M82 fruit at 40 DAA (at which stage the amounts of both citrate and malate were significantly greater in IL2-5) with positionally labelled [14C]glucose. Evolved $^{14}$CO$_2$ was quantified at intervals over a 24h period. The ratios of $^{14}$CO$_2$ released from differently labelled glucose molecules reveals information about the relative rates of different routes of carbohydrate oxidation (Ap Rees, 1980). The total amount of [14C]glucose metabolised by M82 and IL2-5 discs was not significantly different suggesting that capacity for uptake of glucose and overall metabolic rate were similar between the two lines. The rate of oxidation of [14C]glucose to $^{14}$CO$_2$ was linear between 2 and 12h for each positionally labelled glucose (except for [2,14C]glucose which was only linear for the first 5h) for
both M82 and IL2-5 pericarp discs suggesting that a metabolic steady state was achieved (Supplemental Fig. 2). There were no significant differences in the relative rates of oxidation of any of the different labelled positions of glucose between IL2-5 and M82 (Fig. 3) suggesting that there were no substantial changes in flux through the pathways of central metabolism. Of particular relevance is the release of label from the C3,4 and C2 positions of glucose which occurs predominantly via the TCA acid cycle. The release of CO\textsubscript{2} from C3,4 glucose relative to other carbon positions was the same in both tomato lines (Fig. 3 A-C). Similarly, the ratios of CO\textsubscript{2} release from C2 : C1 and C1 : C6 were not significantly different between lines (Fig. 3D, E). These results indicate that the flux through the oxidative steps of the TCA cycle relative to other major carbon oxidation pathways such as the oxidative pentose phosphate pathway is unchanged in pericarp discs of the IL2-5 line.

**Maximum catalytic activities of enzymes of carboxylic acid metabolism**

The labelling experiments give a broad overview of the fluxes in central carbohydrate metabolism, but the complexity of the carbohydrate oxidation network means that it is impossible to ascribe oxidation of a specifically labelled carbon atom exclusively to a single metabolic pathway. Moreover the approach cannot account for complementary changes in flux through parallel pathways in different subcellular compartments. To address this latter issue, we measured the maximum catalytic activity of enzymes of carboxylic acid metabolism, both in isolated mitochondria and in crude tissue extracts to assess the partitioning of metabolic capacity between mitochondria and other subcellular compartments. Mitochondria were isolated from 40 DAA fruit from IL2-5 and M82 plants and the mitochondrial activities of seven of the eight enzymes of the TCA cycle measured. There were no significant differences in the activities of these enzymes in IL2-5 mitochondria compared to M82 (Fig. 4A). We also assessed the total cellular activity of detectable enzymes of carboxylic acid metabolism in cases where there are known to be extra-mitochondrial isoforms (Fig. 4B). Total aconitase activity was significantly lower in IL2-5 fruit (about one third of the activity in M82 fruit). The other four enzymes measured were not significantly changed. Given that aconitase activity in isolated mitochondria was the same in the two lines, this result indicates that the cytosolic isoform of aconitase is present at substantially lower levels in IL2-5 relative to M82.

**Metabolic engineering of carboxylic acid content of tomato fruit**

Aconitase catalyses the conversion of citrate to isocitrate and it follows that a reduction in the activity of aconitase could lead to an accumulation of citrate. To examine if the reduction of total aconitase activity is brought about at the transcriptional level, and would therefore be a good target for genetic engineering, we assessed aconitase transcript levels using semi-quantitative RT-PCR. The tomato genome contains two genes encoding aconitase (Kamenetzky et al., 2010), Solyc07g052340 and Solyc12g005860. The two genes show a high degree of similarity (the predicted cDNA sequences are 88% identical) and both are most similar to ACO3 in Arabidopsis (Bernard et al., 2009). Accordingly, we refer to them as SlAco3a and SlAco3b, respectively. The RT-PCR suggests that the SlAco3a transcript was slightly reduced in IL2-5 fruit (at 40 DAA) in comparison to M82 (Fig. 5). SlAco3b showed a less consistent pattern with an apparent decrease in abundance in some IL2-5 samples but not others (Fig. 5).

To aid in the choice of which Aco gene to target for metabolic engineering, it would be helpful to know the subcellular localisation of the respective gene products. In Arabidopsis, there are three aconitase genes, ACO1 encoding a cytosolic isoform while the products of ACO2 and ACO3 are located in mitochondria (Bernard et al., 2009). The aco-1 mutant allele in *Solanum pennellii* (which corresponds to SlAco3b) is deficient in both cytosolic and mitochondrial aconitase protein.
suggesting that in tomato this gene-product is dual-targeted (Carrari et al., 2003). To investigate the subcellular targeting of both tomato Aco proteins, the SlAco genes were transiently expressed as C-terminal GFP fusions in tobacco leaves (Fig. 6). The two gene products showed clear differences in subcellular localisation. SlAco3a co-localised closely with mito-mCherry indicating an exclusively mitochondrial localisation. In contrast, SlAco3b has a more complex distribution, appearing throughout the cytosol, but was also present in punctate bodies co-localising with mito-mCherry. This suggests that SlAco3b is dual-targeted to both cytosol and mitochondria and is consistent with the decrease in both cytosolic and mitochondrial aconitase in the aco-1 mutant.

The changes in Aco transcripts suggests that Aco genes would be good targets for the engineering of altered carboxylic acid levels in tomato fruit and that to replicate the decrease in cytosolic aconitase activity in IL2-5, SlAco3b should be repressed. Therefore we characterised fruit enzyme and metabolite levels in transgenic plants expressing an 800 bp antisense fragment of SlAco3b (van der Merwe et al., 2010). These transgenic plants were originally generated as part of a study of the role of the TCA cycle in root metabolism (van der Merwe et al., 2010) and their fruit have not been previously characterised.

Two of the transgenic lines (ACO-19 & ACO-38) were grown and allowed to set fruit. Fruit growth, size and number in each transgenic line were indistinguishable from WT. Tomatoes were harvested at breaker (40 DAA) and ripe (55 DAA) stages and extracted for transcript, enzyme and metabolite determinations. Unsurprisingly, given the high degree of sequence similarity between the two SlAco genes, both SlAco3a and SlAco3b transcripts were significantly decreased in both transgenic lines in fruit at both 40 and 55 days after anthesis (Fig. 7). The exception was line ACO19 in which SlAco3b was significantly altered in 40-DAA fruit but not fruit from the later developmental time point. However, aconitase activity was decreased by about 30% compared to WT in both lines at both stages of fruit development (Table 1). Although the activity of aconitase increased developmentally in WT and transgenic lines between the 40- and 55-DAA stages, the relative decrease in activity in the transgenic lines was maintained at around 30%. None of the other measured enzymes of carboxylic acid metabolism were significantly different in the transgenic fruit compared to WT at either stage of development (Table 1).

To establish whether the transgenic manipulation had the predicted effect on citrate levels, fruit carboxylic acids were quantified by GC-MS (Table 2). In ripe fruit, citrate was significantly increased in both transgenic lines by about 40 % confirming successful metabolic engineering. In contrast to the introgression line, however, this increase in citrate was not apparent at the earlier developmental stage (40 DAA) in the transgenic fruit. The increases in citrate in ripe fruit were roughly proportional to the decrease in aconitase levels in both the introgression and transgenic lines. In IL2-5, total aconitase activity decreased by 66% and citrate increased by 60%. In the transgenic lines, aconitase decreased by approximately 30% and citrate increased by 40%.

In the transgenic fruit other carboxylic acids were also significantly altered compared to wild type in ripe fruit but, as with citrate, not at the early 40 DAA stage. Malate increased and fumarate and succinate decreased (Table 2). These changes were similar to those seen in the introgression line at 40 DAA. However, the changes in malate and fumarate in the introgression lines were not apparent in ripe fruit, indicating that the developmental timing of the perturbation of this sector of carboxylic acid metabolism is different in the transgenic lines. The fact that the transgenic lines were not an exact biochemical phenocopy of the introgression lines is to be expected given the presence of many other background genetic changes in the introgression lines and differences in the timing and extent of change of aconitase activity in the transgenic lines. Nevertheless, in quantitative terms, the changes in fumarate and malate were proportional to the decrease in aconitase activity in both the introgression and transgenic lines. Fumarate decreased
proportionately to aconitase: fumarate decreased by 54% in the introgression line and an average
of 37% in the transgenic lines (approximately matching the 66% and 30% decreases of aconitase,
respectively). Malate increased by 120% in the introgression fruit and an average of 76% in the
transgenic fruit, meaning an increase of roughly twice the decrease in aconitase activity. Total
carboxylic acid (malate + citrate + succinate + fumarate) increased from an average of 99 μg g⁻¹
FW in WT fruit, to 140 and 157 μg g⁻¹ FW in ACO-19 and ACO-22 fruit, respectively.
Discussion

Decreased activity of cytosolic aconitase correlates with altered carboxylic acid content in fruit of an introgression line.

Fruit of the selected introgression line, IL2-5, contained elevated levels of citrate and malate and a decrease in fumarate at breaker stage relative to the M82 parent line. The increase in citrate persisted until ripeness under our growth conditions, although this trait may not be stable under all conditions because no increase in citrate was recorded in the ripe fruit of this introgression line when grown in the field (Schauer et al., 2006; Schauer et al., 2008). We found no evidence of major changes in fluxes through relevant metabolic pathways in the IL2-5 fruit reflecting the non-intuitive relationship between metabolite levels and flux (Williams et al., 2008; Kruger and Ratcliffe, 2009).

There was, however, a significant decrease in the maximum catalytic activity of one of the enzymes of carboxylic acid metabolism – aconitase. This change in activity was not apparent in isolated mitochondria, suggesting that it was a cytosolic isoform of aconitase that was altered. Analysis of transiently expressed GFP fusions of the two tomato aconitases in tobacco leaves showed that while SlAco3a is localised exclusively in mitochondria, SlAco3b appears to be dual targeted to both mitochondria and cytosol. There is a precedent for this in yeast in which aconitase is thought to be dual-targeted by inefficient import into mitochondria or by release of the mitochondrially-imported enzyme back into the cytosol (Regev-Rudzki et al., 2005).

The relationship between aconitase activity and genotype in the introgression line is not direct because the two aconitase genes lie on chromosomes 7 and 12 and are therefore not within the introgressed segment on chromosome 2. RT-PCR suggests a slight decrease in expression of the two SlAco genes so perhaps the introgressed region in IL2-5 contains a transcriptional regulator of the aconitase gene. This possibility is supported by the identification of several enzyme QTLs in the same introgression line population in which the introgressed region does not contain the structural gene of the enzyme concerned (Steinhauser et al., 2011).

Aconitase activity is a determinant of fruit citrate levels

A decrease in activity of aconitase provides an obvious link to accumulation of citrate, its substrate. Perturbation of the TCA cycle could also provide an explanation for changes in other carboxylic acids. Further evidence that the change in citrate is directly linked to aconitase activity and not some other background effect in the introgression line was provided by direct manipulation of aconitase activity in transgenic plants. In fact, the increase in citrate in ripe fruit was quantitatively proportional to the decrease in aconitase in both the introgression and transgenic lines, suggesting that aconitase activity is a major determinant of ripe fruit citrate levels. In support of this, pharmacological inhibition of aconitase in citrus fruit also led to an increase in citrate levels (Degu et al., 2011). Aconitase activity is also linked to citrate levels in other tissues in tomato: leaf citrate levels were also increased in the aco-1 mutant (Carrari et al., 2003). However, it is worth noting that, in the transgenic lines, the decreased aconitase activity at the earlier fruit developmental stage (40 DAA) did not result in a significant increase in citrate level. This suggests that other factors, in addition to aconitase activity, influence fruit citrate level. The difference in the relationship between aconitase and citrate at this developmental stage in transgenic versus introgression-line fruit is most likely due to the difference in background genotype (Moneymaker vs M82, respectively).

Citrate synthase might also be expected to influence citrate levels and this was demonstrated in the leaves of transgenic tomato plants with reduced citrate synthase (Sienkiewicz-Porzucek et al.,...
287 However, there was no direct relationship between citrate synthase activity and citrate levels in arsenite–treated citrus (Sadka et al., 2000). Transgenic manipulation to decrease the activity of NAD-isocitrate dehydrogenase, the enzyme immediately downstream of aconitase, had no effect on tomato leaf citrate levels (Sienkiewicz-Porzupek et al., 2010). It appears that there is not a simple relationship between carboxylic acid levels and the activity of the enzymes that catalyse their interconversion (Nunes-Nesi et al., 2008). This disconnect between metabolite levels and maximum catalytic activities of enzymes extends throughout the network of central carbon metabolism (Sulpice et al., 2010) and the effect of altered aconitase activity on citrate levels may therefore be considered somewhat exceptional.

296 **Aconitase activity also affects the accumulation of other carboxylic acids**

297 In addition to increasing citrate content in ripe fruit, transgenic suppression of aconitase also led to a substantial increase in malate as well as decreases in succinate and fumarate. Perturbation of several carboxylic acids by manipulation of a single TCA cycle enzyme is not uncommon (Araujo et al., 2012) and is a reflection of the interconnected nature of carboxylic acid metabolism. Similar changes in carboxylic acid levels were seen in the introgression line where the change in aconitase activity was restricted to the cytosol.

307 Interpretation of these changes is complicated by the compartmentation of tricarboxylic acid metabolism and may not be directly related to mitochondrial carboxylic acid metabolism. For instance, the levels of predominantly vacuolar carboxylic acids such as citrate and malate are influenced by the rate of export of these metabolites from mitochondria. In heterotrophic Arabidopsis cells, these export fluxes are an order of magnitude lower than the mitochondrial TCA cycle fluxes (Williams et al., 2008). Thus, changes in the accumulation rate of citrate and malate can be caused by proportionally small changes in mitochondrial TCA cycle flux that may be within the error range of the flux estimate (and therefore undetectable). This most likely explains the lack of detectable change of overall carboxylic acid oxidation in the introgression line fruit and further suggests that flux through the cytosolic pathway of citrate metabolism is low in relation to the mitochondrial pathway. Alternatively, there could be a compensatory increase in flux through the mitochondrial pathway (Morgan et al., 2008).

315 The opposing changes in fumarate and malate may also be explained by the compartmentation of these metabolites. In mitochondria, the interconversion of fumarate and malate is close to equilibrium and so one would expect the levels of these two metabolites to follow one another. The observed decreased fumarate and increased malate levels in both the transgenic and introgression line fruit reflects the fact that the measured malate is mainly extra-mitochondrial (vacuolar). Most likely, an increased efflux of malate from the mitochondrion leads to increased accumulation of malate in the vacuole. Malate concentration in the mitochondrion is probably decreased, in line with the decreased fumarate content.

323 The function of cytosolic aconitase in ripening fruit is unclear. One suggestion is that it is involved in metabolism of citrate released from the vacuole, to provide an entry point into amino acid metabolism or the GABA shunt in citrus (Degu et al., 2011). However, in tomato fruit, citrate is accumulating at the phase of ripening under consideration, and so extensive efflux of citrate from the vacuole seems unlikely. An alternative possibility is that metabolism of citrate in the cytosol contributes to cytosolic NADPH provision by providing a substrate for the NADP-dependent isocitrate dehydrogenase, or is important for generation of carbon skeletons for the synthesis of glutamate and aspartate, both of which accumulate substantially in the later phases of tomato ripening (Baxter et al., 2005).
Conclusion

This study demonstrates that individual lines of genetic mapping populations can provide useful information to guide metabolic engineering strategies. Although such lines contain relatively large regions of introgressed DNA from a genetically distinct parental line, detailed biochemical analysis can pinpoint the main point of metabolic disturbance and highlight potential candidate proteins that can be tested in a targeted manner in transgenic plants. Here, the introgression line allowed us to focus specifically on aconitase amongst a myriad of possible targets for manipulation of accumulation of carboxylic acids in tomato fruit. One could envisage further refinement of the transgenic manipulation by using fruit specific promoters that are more finely tuned to the appropriate developmental stage and take into account the variations in metabolism within different fruit tissues (Moco et al., 2007; Matas et al., 2011).
Materials and methods

Plant material: Seeds of *Solanum lycopersicum*, cv M82 and introgression line 2-5 were kindly supplied by the Tomato Genetics Resource Centre (TGRC, Davis, USA). Transgenic *Solanum lycopersicum* cv ‘Moneymaker’ expressing an antisense aconitase construct were previously described (van der Merwe et al., 2010). Plants were grown in a 16 h photoperiod at 22-23°C day temperature, 20-22°C night temperature and with supplementary lighting to maintain an irradiance of 250-400 μmol m⁻²s⁻¹. Plants were grown in a standard potting compost supplemented with slow-release fertiliser. Tomato feed was applied during flowering and fruit set and a 0.5% (w/v) calcium chloride solution was sprayed directly onto all developing fruit weekly to help control blossom end rot.

Analysis of carboxylic acids by ¹H NMR: freeze-dried pericarp tissue was extracted in 70% methanol-d₄ / 30% D₂O exactly as described by (Le Gall et al., 2003). ¹H NMR spectra were recorded at 20°C on a Varian Unity Inova 600 spectrometer (Varian Inc., Palo Alto, USA) using a 5-mm HCN triple resonance z-gradient probe and the standard Varian pulse sequence with a relaxation delay of 2 s, including a presaturation pulse to suppress the residual water signal, a 90° pulse angle, a spectral width of 10 ppm, and a 4 s acquisition time. Methanol-d₄ was used for the internal lock signal and 320 transients were collected for each spectrum. Spectra were processed and analyzed using NUTS (Acorn NMR Inc., Livermore, USA). A 1 Hz line broadening was applied before Fourier transformation and peaks were integrated manually within NUTS.

Analysis of carboxylic acids by GC-MS: frozen pericarp tissue powder was extracted in chloroform-methanol and carboxylic acids quantified by GC-MS as described previously (Roessner et al., 2001), following a procedure optimised for tomato tissue (Roessner-Tunali et al., 2003).

Analysis of amino acid content: frozen pericarp tissue powder was extracted in 0.1 M HCl and proteinogenic amino acids quantified by HPLC (Bruckner et al., 1995).

Carbohydrate oxidation fluxes: freshly harvested pericarp discs (10 mm diameter, 3 mm thick) were washed in 10 mM MES-KOH (pH 6.5). Eight discs were placed into 100 ml flasks containing 5.0 ml 10 mM MES-KOH (pH 6.5) with 1 mM glucose supplemented with [1-¹⁴C]-, [2-¹⁴C]-, [3,4-¹⁴C]-, or [6-¹⁴C]-glucose (3.7 kBq/flask). Evolved ¹⁴CO₂ was trapped in 0.5 ml of 10% (w/v) KOH and radioactivity quantified by liquid scintillation counting (Harrison and Kruger, 2008).

Isolation of mitochondria: fresh tomato pericarp tissue was roughly chopped and 50-100 g was placed in a square-section polycarbonate container with 200-300 ml of extraction medium (0.3 M sucrose, 25 mM tetrasodium pyrophosphate, 25 mM TES-KOH (pH 7.5), 2 mM EDTA, 10 mM KH₂PO₄, 1% (w/v) polyvinylpyrrolidone (PVP-40), 1% (w/v) BSA, 20 mM ascorbic acid). The sample was homogenised using multiple short bursts (less than 1 s) of a Status Polytron blender (Kinematica Inc., Bohemia, USA) on a low setting (4). The sample was filtered through one layer of miracloth and two layers of muslin. The filtrate was centrifuged at 1,085 g for 5 min. The pellet was discarded and the supernatant centrifuged for 15 min at 23,500 g. The pellet was resuspended in wash buffer (0.3 M sucrose, 10 mM TES-KOH (pH 7.2), 0.1% (w/v) BSA) to an approximate final volume of 2 ml before being layered onto a 35 ml gradient of 0-4.4% (w/v) PVP-40 in 18% (v/v) Percoll. The gradient was centrifuged at 40,000 g for 40 min and the mitochondria removed from the band near the bottom of the gradient using a 5 ml pipette. The mitochondria were diluted with wash buffer and centrifuged twice at 27,000 g for 15 min. The final pellet was resuspended in a minimal volume of wash buffer.
Enzyme assays: maximum catalytic activities of the enzymes of carboxylic acid metabolism were measured spectrophotometrically in desalted extracts of pericarp tissue or in isolated mitochondria according to (Morgan et al., 2008).

Semi-quantitative RT-PCR: the following primers were used to amplify Solyc07g052340 (fwd, CATGAAACAACCTTGCCAGTG; rev, GCAGCTTCTGCTCCTAACACGTC) and Solyc12g005860 (fwd, TCCACAAAGATAGCCCTGT; rev, TCCCATCCTACCAAAGTTGC) from fruit pericarp cDNA. Cycle numbers and annealing temperatures were optimised for each template to ensure that the amplification reaction was tested in the exponential phase. Products were visualised using ethidium bromide after separation by agarose gel electrophoresis.

Quantitative real-time PCR: the following primers were used to amplify Solyc07g052340 (fwd GCCGCTTGCTCAACTCTAC, rev GACTCCACCTCAGGACAGA) and Solyc12g005860 (fwd TGTGGTTATTGCTCAGGTGA, rev CAACACCGTATCTCCACCTCA). cDNA amplification was quantified using the fluorescent intercalating dye SYBR Green in an iCycler detection system (Bio-Rad, Hemel Hempstead, UK). Transcript abundance was normalised against ubiquitin3, amplified using the following primers (fwd, AGGTTGATGACACTGGAAAGGTT, rev ATCGCTCCAGCTTGGTCTGTA).

Subcellular localisation of SlAco-GFP fusion proteins transiently expressed in tobacco leaves. SlAco3a and SlAco3b were PCR amplified from fruit cDNA using the following primer pairs: SlAco3a_for (caccATGTATAGTAATACAGCTCGCAAGTAC) and SlAco3a_rev (TTGCTTTGTCAATTGACGAATGACATATTG); SlAco3b_for (caccATGTACGTTTCTTCTTCTGTTCACATCAAAC) and SlAco3b_rev (TTGCTGACTCAGCTGCAATGACGTATG). Purified PCR products were sub-cloned into pUB10-cGFP (Grefen et al., 2010) and transformed into Agrobacterium tumefaciens strain GV3101. Transformed agrobacteria were infiltrated into mature Nicotiana tabacum leaves together with agrobacterium containing expression constructs for mitochondrial-targeted mCherry (Nelson et al., 2007). Four days after infiltration, excised leaf tissue was imaged on a Leica DM6000 CS confocal microscope (Leica Microsystems, CMS GmbH Mannheim, Germany). GFP was excited with an Argon 488 laser at 35 % power and GFP signal was collected between 505 and 525 nm. The mCherry fluorophore was excited with the HeNe 543 laser set to 25 % power and emitted signal was collected between 595 to 620 nm. Chlorophyll fluorescence was collected with HeNe 543 excitation at 20 % power between 650 to 720 nm. Images were processed using Leica LAS AF Lite software.

Statistical analyses: Student’s t test (two-tailed, unequal variance) was used to determine the significance of differences in enzyme activities and metabolite levels between tomato lines. The mean ± SEM of ratios of 14CO2 release from positionally labelled glucose were subjected to log transformation prior to repeated measures analysis of variance (ANOVA) based on Type III sums-of-squares (SPSS/PASW Statistics 18, IBM Corp, Armonk, New York, USA). Homogeneity of variance of the dependent variable in repeated measures ANOVA was confirmed using Levene’s test prior to assessing the significance of differences between plant lines. Only statistical differences for which P < 0.05 are considered significant.

Acknowledgements

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References


Table 1. Maximum catalytic activities of enzymes of carboxylic acid metabolism in fruit of wild type and transgenic tomato plants expressing an antisense aconitase construct. Tomatoes harvested at 40- and 55 days after anthesis (DAA) were extracted and the maximal catalytic activities of six enzymes of carboxylic acid were metabolism measured. Wild type and two independent transgenic lines (ACO 22 and ACO 38) were assessed. Each value is the mean ± SEM of six biological replicates. Values significantly different from wild type are indicated in bold (t-test, P<0.05).

Abbreviation: DH, dehydrogenase

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Enzyme activity (nmol min⁻¹ gFW⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>40 DAA</td>
</tr>
<tr>
<td></td>
<td>WT</td>
</tr>
<tr>
<td>Aconitase</td>
<td></td>
</tr>
<tr>
<td></td>
<td>20.1 ± 5.3</td>
</tr>
<tr>
<td>Succinate DH</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.73 ± 0.12</td>
</tr>
<tr>
<td>Fumarase</td>
<td></td>
</tr>
<tr>
<td></td>
<td>190 ± 32.3</td>
</tr>
<tr>
<td>NAD+-malate DH</td>
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</tr>
<tr>
<td></td>
<td>3521 ± 145</td>
</tr>
<tr>
<td>Citrate synthase</td>
<td></td>
</tr>
<tr>
<td></td>
<td>11.3 ± 1.8</td>
</tr>
<tr>
<td>NADP-isocitrate DH</td>
<td></td>
</tr>
<tr>
<td></td>
<td>22.5 ± 2.5</td>
</tr>
</tbody>
</table>

Table 2. Carboxylic acid content of fruit of transgenic plants expressing an antisense aconitase construct. Fruit extracts were analysed using GC-MS. Tomatoes were harvested at 40 and 55 days after anthesis (DAA) as indicated. Data are the mean ± SEM of six independent fruits. Values in bold are significantly different from WT (t-test, P<0.05)

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Metabolite level in fruit (relative to WT)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>40 DAA</td>
</tr>
<tr>
<td></td>
<td>WT</td>
</tr>
<tr>
<td>Citrate</td>
<td>1.00±0.04</td>
</tr>
<tr>
<td>Succinate</td>
<td>1.00±0.02</td>
</tr>
<tr>
<td>Fumarate</td>
<td>1.00±0.08</td>
</tr>
<tr>
<td>Malate</td>
<td>1.00±0.08</td>
</tr>
</tbody>
</table>
FIGURE LEGENDS

Fig. 1. Carboxylic acid content of fruit of M82 and introgression line 2-5 during fruit development. Organic acid content was determined by \(^1\)H NMR for M82 (triangles) and IL2-5 (squares) in extracts of pericarp tissue at 30, 40, and 55 DAA (days after anthesis). Each value is the mean ± SEM (n=6). * indicates values significantly different from M82, (t-test, P<0.05)

Fig. 2. Amino acid content of tomato fruit (55 days after anthesis) from M82 and introgression line 2-5. Black bars, M82; grey bars introgression line 2-5. Values are the mean ± SEM (n=4). * P≤0.05 (t test).

Fig. 3. Relative rates of oxidation of specific labelled glucose carbons in tomato fruit from M82 and introgression line 2-5. Pericarp discs from fruit harvested 40 days after anthesis were incubated with [1-\(^14\)C], [2-\(^14\)C], [3,4-\(^14\)C ] or [6-\(^14\)C ]-glucose (1 mM). The \(^14\)CO\(_2\) released by metabolism was monitored at intervals throughout a 24h incubation period and quantified by liquid scintillation counting. Ratios of cumulative \(^14\)CO\(_2\) release from different combinations of positionally labelled glucose for IL 2-5 (squares, solid line) and M82 (triangles, dotted line) are presented. Data are the ratio of the mean ± SEM (n = 4). There were no significant differences (repeat measures ANOVA) in the ratios of \(^14\)CO\(_2\) released by M82 and IL 2-5 for any of the combinations of positionally labelled substrates analysed (F < 3.78; d.f. = 1,6; P > 0.10)

Fig. 4. Maximum catalytic activities of enzymes of carboxylic acid metabolism in fruit from M82 and introgression line 2-5. Enzyme activities in (A) isolated mitochondria and (B) crude pericarp extracts from 40 DAA fruit. Data are mean ± SEM (n=3). For clarity the NAD-malate dehydrogenase activity is divided by ten. * Indicates significantly different from M82 (t test, P<0.05.). Abbreviations: ICDH, isocitrate dehydrogenase; 2-OGDH, 2-oxoglutarate dehydrogenase; DH, dehydrogenase. Co-enzyme specificity is indicated where relevant.

Fig. 5. Semi-quantitative RT-PCR analysis of aconitase transcript abundance in M82 and IL2-5 fruit at 40 DAA. Replicate PCR products amplified from cDNA template prepared from three independent fruit of M82 and IL2-5 plants. The transcript encoding ubiquitin (UBQ) was used as a control.

Fig. 6. Subcellular localisation of SlAco3a and SlAco3b. Gene constructs consisting of SlAco genes fused C-terminally to GFP and driven by the Arabidopsis ubiquitin 10 promoter were transiently expressed in N. tabacum leaves by agro-infiltration. GFP fluorescence was colocalised with mitochondrial-targeted mCherry (mito-mCherry). GFP fluorescence of SlAco3a-GFP (A) and SlAco3b-GFP (B) with mito-mCherry signal as indicated. Chlorophyll auto-fluorescence is also shown. Scale bars = 20 \(\mu\)m.

Fig. 7. Abundance of SlAco3a and SlAco3b transcripts in fruit from antisense transgenic lines. RNA was extracted from fruit from WT and transgenic lines (ACO 22 and ACO38) at 40 and 55 days after anthesis (DAA). SlAco3a and SlAco3b transcripts were quantified by quantitative real-time PCR. Mean abundance values (relative to WT) ± SEM of 3 biological replicates are shown. * indicates significantly different from WT (p<0.05, t-test).

Supplemental Figure 1. Carotenoid content of M82 and IL2-5 fruit during development. Carotenoids were measured in the pericarp of fruit from M82 (square, solid line) and IL 2-5 (triangle, dotted line) at specific days after anthesis (DAA). The component carotenoids were lutein (A), phytoene (B), \(\beta\)-carotene (C), \(\alpha\)-tocopherol (D) and lycopene (E). Data are the mean ± SEM from individual measurements of fruit from three separate plants. * indicates values significantly different from M82 (t-test, P<0.05).
Supplemental Figure 2. Rate of oxidation of $^{14}$C[glucose] to $^{14}$CO$_2$ by tomato pericarp discs from M82 and IL2-5 fruits is linear over the first 12h. Pericarp discs from fruits harvested 40 days after anthesis were incubated with [1-$^{14}$C], [2-$^{14}$C], [3,4-$^{14}$C] or [6-$^{14}$C]-glucose (1 mM). The $^{14}$CO$_2$ released by metabolism was monitored at intervals throughout a 12 h incubation period and quantified by liquid scintillation counting. Data are expressed as the percentage of supplied $^{14}$C and values are means ± SEM (n = 4).
Fig 1

- **Citrate**: The graph shows an increase in the metabolite amount (nmol mgDW⁻¹) with fruit age (DAA). The data points indicate a significant increase at certain DAA stages, marked with asterisks.

- **Malate**: Similar to citrate, malate also shows an increase with fruit age. The graph highlights a peak and subsequent decline, followed by another increase. Asterisks denote significant changes.

- **Fumarate**: The graph for fumarate is different from citrate and malate. It displays a notable increase with fruit age, followed by a plateau and a slight decrease towards the end. Asterisks indicate significant differences.

The x-axis represents fruit age (DAA), while the y-axis represents metabolite amount (nmol mgDW⁻¹).
Fig 3

Ratio of $^{14}$CO$_2$ release from:

A: $[1-^{14}$C$]$glucose
B: $[2-^{14}$C$]$glucose
C: $[6-^{14}$C$]$glucose
D: $[3,4-^{14}$C$]$glucose
E: $[1-^{14}$C$]$glucose
F: $[6-^{14}$C$]$glucose

Incubation time (h)

0 2 4 6 8 10 12 24 0 2 4 6 8 10 12 24
Fig 4

A

Enzyme activity (nmol min$^{-1}$ mg protein$^{-1}$)

- M82
- IL 2-5

B

Enzyme activity

- M82
- IL 2-5

* indicates a significant difference.
Fig 5

<table>
<thead>
<tr>
<th>Transcript</th>
<th>M82</th>
<th>IL 2-5</th>
</tr>
</thead>
<tbody>
<tr>
<td>SIAco3a (Solyc07g052340)</td>
<td>![Image]</td>
<td>![Image]</td>
</tr>
<tr>
<td>SIAco3b (Solyc12g005860)</td>
<td>![Image]</td>
<td>![Image]</td>
</tr>
<tr>
<td>UBQ</td>
<td>![Image]</td>
<td>![Image]</td>
</tr>
</tbody>
</table>
A) SlAco3a-GFP, mito-mCherry, chlorophyll merge
B) SlAco3b-GFP, mito-mCherry, chlorophyll merge
Transcript abundance (relative to WT)

**Fig 7**

**SI Aco3a**

- WT
- ACO19
- ACO38

40 DAA
55 DAA

**SI Aco3b**

- WT
- ACO19
- ACO38

40 DAA
55 DAA