Growth, Yield, and Metabolic Responses of Temperature-stressed Tomato to Grafting onto Rootstocks Differing in Cold Tolerance

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ABSTRACT. Tomato (Solanum lycopersicum cv. Kommeet) plants were either self-grafted, grafted onto the cold-sensitive cultivar Moneymaker or onto LA 1777, a cold-tolerant accession of Solanum habrochaites. Plants were grown at three different temperatures (T): optimal (19.4 °C), intermediate (17 °C), or low (14.6 °C). Grafting tomato ‘Kommeet’ onto LA 1777 increased shoot growth at intermediate and optimal T and root growth at low or intermediate T in comparison with self-grafting or grafting onto ‘Moneymaker’. Reducing T significantly suppressed fruit yield as a result of reduced fruit number per plant without any interaction with grafting treatments. Grafting ‘Kommeet’ onto LA 1777 remarkably restricted fresh and dry fruit mass in comparison with self-grafting of ‘Kommeet’ as a result of a reduction in the fruit number per plant while not influencing flower number per plant and mean fruit mass, regardless of T regime. Negative effect of LA 1777 on fruit setting points to impairment of pollen fertility indicating signals originating from the root. At low and intermediate T, plants grafted onto LA 1777 were capable of increasing soluble carbohydrates, total amino acids, and guaiacol peroxidase activity in roots to higher levels than those grafted onto S. lycopersicum rootstocks while maintaining a significantly lower malondialdehyde content. These differences point to a much weaker oxidative stress in roots of plants grafted onto S. habrochaites than those grafted onto S. lycopersicum rootstocks when exposed to low and intermediate T. Furthermore, plants grafted onto LA 1777 exhibited significantly higher levels of soluble carbohydrates, total amino acids, and guaiacol peroxidase activity in leaves and fruit and superoxide dismutase in fruit at low and intermediate T, which indicates that LA 1777 improved the level of antioxidant compounds in tomato shoots, thereby enhancing its adaptation to lower T than optimal.

Tomato originates from subtropical regions. Its current economic optimum temperature (T) for greenhouse cultivation and fruit set is at 19 to 20 °C (Van der Ploeg and Heuvelink, 2005). Suboptimal T down to 10 °C affects tomato growth and development by shortening internodes that reduce plant length, retard leaf expansion, reduce leaf number and total leaf fresh weight while increasing dry matter content and thickness of leaves arising from increased starch storage (Venema et al., 1999). When T falls below 12 °C, growth is inhibited and will cause irreversible alteration and dysfunctions at the cellular level, depending on species, intensity, and duration of exposure (Kodama et al., 1995).

The most vulnerable stage in the growth cycle of higher plants is the reproductive phase (Nishiyama, 1995) and specifically during the formation of reproductive organs such as flowers, fruit, and seeds. The reproductive phase is affected by the interactions among plant morphology, physiology, and growth conditions (Van der Ploeg and Heuvelink, 2005). According to Van der Ploeg and Heuvelink (2005), optimal T for fruit set is between 18 and 20 °C. T below this level may lead to suppression of yield as a result of decreased fruit set owing to reduction in both quality and quantity of pollen (Domínguez et al., 2005) and retardation of truss appearance rates (Van der Ploeg and Heuvelink, 2005), which restricts fruit number per plant. Indeed, T below the optimal range will impair cell membrane fluidity and permeability, thereby resulting in ion leakage (Abbas, 2012), whereas intra- and extracellular water and nutrient movement are inhibited (Mahajan and Tuteja, 2005; Salinas, 2002), reactive oxygen species (ROS) are generated (Gill and Tuteja, 2010), photosynthesis may be restricted (Theocharis et al., 2012), and finally yield is reduced.

As a result of interaction of stress- and sugar-signaling pathways under abiotic stress conditions (Gupta and Knaur, 2005), plant metabolism is modulated. Indeed, during cold stress, sucrose accumulation in leaves leads to feedback inhibition of photosynthesis (Chiou and Bush, 1998; Ruelland et al., 2009). Thus, cellular components and structures are modified by suboptimal T exposure resulting in, for example, swelling of mitochondria and disruption in protein lipid accumulation and protein denaturation and stability (Kratsch and Wise, 2000; Siddiqui and Cavicchioli, 2006). Moreover, cellular dehydration caused by suboptimal T stress is attributed to reduced water...
uptake by roots (Yadav, 2010) and inhibition of stomatal closure (Wilkinson et al., 2001). Suboptimal T reduces not only water, but also nutrient uptake, thereby causing osmotic stress (Chinnasamy et al., 2007). In addition, suboptimal T causes peroxidation of unsaturated membrane lipids (Theocharis et al., 2012) thereby increasing membrane permeability to nutrients and ions that result in ion leakage (Gao et al., 2008). Such modifications are capable of disorganizing the entire plastid leading to growth reduction with an indirect yield loss. Furthermore, at suboptimal T, reactions of the Calvin cycle are down-regulated, and this may lead to overreduction of the photosynthetic light reactions and generation of ROS (Ensminger et al., 2006).

Reactive oxygen species are generated in plant cells even during a normal metabolic process such as photosynthesis and respiration (Apel and Hirt, 2004). Under optimal growth conditions, the damaging effect of ROS is prevented through the scavenging activity of antioxidants such as superoxide dismutase (SOD, EC 1.12.1.1) and guaiacol peroxidase (GPOD, EC 1.11.1.7) (Sundar et al., 2004). ROS scavenging capacity can be enhanced by hormones such as cytokinins (Zhou et al., 2007). In addition, suboptimal T causes peroxidation of unsaturated membrane lipids (Theocharis et al., 2012) thereby nutrient uptake, thereby enhancing the growth rate and yield performance of S. lycopersicum as rootstocks, characterized by a vigorous root system and tolerance to suboptimal T, may minimize yield losses in elite cultivars of grafted tomato when exposed to suboptimal T. However, when the balance between scavenging activity of antioxidants and ROS formation is disturbed as a result of suboptimal T stress, oxidative damage is caused.

Suboptimal T affects pollen quality and quantity (Dominguez et al., 2005), thereby reducing fruit set and concomitantly fruit number per plant. Long-term exposure to suboptimal T results in larger fruit (Van der Ploeg and Heuvelink, 2005).

To avoid yield losses resulting from suboptimal T, indoor production of this economically important crop has become crucial. However, high-energy costs required for greenhouse heating, increasing energy prices, and environmental concerns about the emission of CO₂ during the combustion of fossil fuel are forcing the greenhouse industry to improve its energy efficiency and to seek alternative approaches for enhancing plant tolerance (Ntatsi et al., 2013). An increase in tolerance might considerably reduce fuel cost (Venema et al., 2008). Already a reduction of 1 °C from the optimum T leads to a saving of 8% in energy costs (Elings et al., 2005). A possible strategy would be the introduction of new tolerant cultivars. Unfortunately, the very low genetic diversity of the existing cultivars (Venema et al., 2008) in combination with limitations in breeding methodology has led this approach to fail up to date.

Therefore, a promising tool to achieve this goal is grafting onto rootstocks that are known to affect canopy development, disease resistance, or cold-hardiness (Schwarz et al., 2010). The Solanaceae family includes many wild relatives of the cultivated tomato species S. lycopersicum with different degrees of tolerances to abiotic and biotic stresses. However, S. lycopersicum is characterized by low genotype diversity with respect to cold stress responses (Schwarz et al., 2010) and, therefore, the use of other Solanaceae species or of interspecific breeds as rootstocks might be an alternative. Accessions of S. habrochaites [synonym Lycopersicon hirsutum (Spooner et al., 2005)] are of particular interest as potential germplasm sources to widen the genetic variation in low T tolerance of the cultivated tomato (Venema et al., 2005, 2008). This wild tomato species originates from an altitude of 3200 m (Rick et al., 1994) where an adaptation to suboptimal T stress can be expected (Patterson et al., 1978). Superiority of S. habrochaites in terms of cold adaptability is attributed to adaptive mechanisms alleviating cell damage and retaining its reproduction ability under suboptimal T conditions (Venema et al., 2008).

Recent studies indicated that moderately suboptimal T has no adverse impact on root growth of high-altitude accessions of S. habrochaites (Venema et al., 2008). Therefore, a reduction in upward transport capacity of water and nutrients is not expected when this S. habrochaites accession line is exposed to suboptimal T (Venema et al., 2008). Similarly, imbalances in supply rate of root-deprived phytohormones to the scion caused by exposure to suboptimal T are also not expected. This hypothesis is further strengthened by findings indicating that use of rootstocks with a vigorous root system results in increased nutrient and water uptake, thereby enhancing the growth rate and yield performance of the whole plant (Lee, 1994).

In this article, the hypothesis is tested that using wild relatives of S. lycopersicum as rootstocks, characterized by a vigorous root system and tolerance to suboptimal T, may minimize yield losses in elite cultivars of grafted tomato when exposed to suboptimal T. The following genotypes were used as rootstocks: 1) the cold-tolerant S. habrochaites accession LA 1777 (Venema et al., 2008); and 2) the elite cultivar Moneymaker, which appears to be cold-sensitive (Dominguez et al., 2005; Venema et al., 2008). The two rootstocks were grafted to the same commercial tomato cultivar Kommeet. Moreover, a control treatment with 3) self-grafted ‘Kommeet’ plants was applied to avoid confounding of effects imposed by grafting incision with those arising from the rootstock genotype. Impact of rootstock genotype on plant growth and yield under conditions of suboptimal (i.e., intermediate or low) T was assessed by measuring characteristics of shoot and root growth, fruit, yield, and fruit quality. To elucidate the physiological mechanisms underlying the observed effects on growth and yield, carbohydrate accumulation, carbon-to-nitrogen (C/N) ratio, oxidation characteristics, and antioxidant enzyme activities were also measured.

Materials and Methods

Plant material, growth conditions, and measurements. The experiment was carried out at the Leibniz Institute of Vegetable and Ornamental Crops, Großbeeren, Germany (lat. 52°20' N, long. 13°18' E, alt. 40 m). A commercial tomato cultivar, Kommeet, was self-grafted or grafted onto the tomato cultivar Moneymaker or onto the line accession LA 1777 (S. habrochaites). Seeds of LA 1777 were sown in coarse sand on 20 June 2011; the two other cultivars were sown 1 week later. Grafting was performed 1 month later, when seedlings had developed three to four true leaves.

Grafted plants were transplanted on 11 Aug. 2011 into six 60-m² greenhouse cabins and cultivated at a heating set point of 18/16 °C (day/night). Roof ventilation was used once air T reached 27 °C for both day and night. At a relative humidity above 80%, the vents were opened for exchange with outside air to reduce relative humidity. Pure CO₂ was supplied to keep CO₂ concentration in all cabins at ambient levels matching the outside environment.

Grafted plants were transferred into two troughs (8 × 0.2 × 0.07 m) located in the middle of the cabins where they were grown in a nutrient film technique. To manage the climate inside the cabin and to maintain appropriate growth, the two troughs were bordered from both sides by three further troughs. Each trough contained 12 plants. A standard nutrient solution (EC 3.7 dS·m⁻¹, pH 5.6) for tomato was prepared by mixing stock solutions and rain water (De Kreij et al., 1997) and was supplied to the troughs every 10 min for 20 s. The composition of the...
nutrient solution was measured periodically in the laboratory and corrected when necessary.

Once the second truss was flowering, on 9 Sept., set points for heating by day/night were shifted within 3 d to 20/18, 16/14, and 11/9 °C in each of the two greenhouse cabins. Different set points for heating in combination with global radiation and outside T resulted in mean cabin T during the treatments of 19.4 (optimal), 17 (intermediate), and 14.6 °C (low).

Differences in the mean T resulted mainly from the differences during the night phase, whereas during the daytime and more specifically the middle of the day, T differences between treatments were slight as a result of solar radiation and equal set points for ventilation. However, toward the end of the experiment, daytime solar radiation could not compensate for lower heating set points in the low T treatments resulting in mean average T in the low T treatment of only 11 °C. Treatments had minimum/maximum T as follows: optimal (14.1/30.0 °C), intermediate (12.7/30.7 °C), and low (9.2/30.1 °C).

Mean daily photosynthetically active radiation (PAR) at the canopy top decreased from 13.7 mol·m⁻² during the first 4 weeks after planting to 2.5 mol·m⁻² during the last 4 weeks of the experiment. Mean daily integral of PAR over the entire treatment period was 7.4 mol·m⁻²·d⁻¹. The experiment was ended on 15 Dec. 2011.

**Growth characteristics.** Total number of flowers and fruit was counted at each truss beginning with Truss 3 until Truss 8 at four dates, 20, 34, 59, and 95 d after treatment start, to calculate a mean for the cultivation time. Leaf area was calculated as a mean of five fully expanded leaves. We measured the width of Leaves 11 to 15 counted from the shoot tip at the same dates as mentioned before (Schwarz and Klärning, 2001). At the end of the experiment, two plants of each grafted treatment were harvested per experimental unit, divided into stem, leaves, fruit, and roots. Subsequently, fresh weight of each of these plant parts was determined. Thereafter, all plant parts were dried separately at 70 °C to determine dry plant biomass and its allocation into different plant parts. Subsamples were used for later chemical analyses. Fresh root samples of ≈1 g were collected from each experimental unit to measure root length (meters per gram) and mean root diameter (millimeters) using an image analyses system (WinRhizo; Regent Instruments, Sainte-Foy, Quebec, Canada). With these results and root fresh weight, total root surface area and total root length were estimated as described in Ntatsi et al. (2013).

**Yield.** Harvesting started on 26 Oct. and finished 15 Dec. Number of harvested fruit and total fruit yield per plant were recorded up to the sixth truss. Marketable yield was classified as fruit without cracking, blossom-end rot (BER), blotchy ripening, deformations, and small fruit (diameter less than 30 mm). Marketable fruit as well as fruit with BER were counted, fresh, and dry-weighed.

**Tissue analyses.** Powder from oven-dried root, leaf, and fruit samples were used to determine the total carbon and nitrogen content of different plant parts. The Dumas combustion technique was performed with the samples using a C/N analyzer (Elementar, Hanau, Germany).

Soluble sugars were measured according to Geigenberger and Stitt (1993), starch according to Sonnewald (1992), and total amino acids according to Moore and Stein (1948) using a 96-position microplate spectrophotometer (Synergy HT; BioTek Instruments, Winooski, VT). Lipid peroxidation was estimated in leaves and roots by determining the concentration of thiobarbituric acid-reactive substances and quantified on its product, malondialdehyde (MDA), according to the method described by He et al. (2009) and Hodges et al. (1999). MDA was not detected in fruit and therefore data are presented only for roots.

For enzyme analysis, 0.1 g of each pulverized frozen sample (leaf, root, or fruit) was homogenized with ice-cold 25 mm HEPES buffer (pH 7.8) including 0.2 mm EDTA, 2 mm ascorbate, and 2% (w/v) polyvinylpyrrolidone. The homogenate was centrifuged at 4 °C and 17,562 g, for 5 min. Supernatants were used for enzyme analysis. All steps in preparation of enzyme extract were carried out at 4 °C. Protein concentration was determined as described by Bradford (1976) with bovine serum albumin as the standard. GPOD activity was determined according to Cakmak and Marschner (1992) as modified by Egley et al. (1983). Increase in absorbance at 510 nm caused by guaiacol oxidation (E = 26.6 mM·cm⁻¹) was measured over 50 min. Glutathione reductase (GR) activity was assayed according to Rao et al. (1996) with some modifications. Briefly, the reaction mixture in a total volume of 200 μL consisted of 25 mm potassium buffer (pH 7.0), 0.1 mm EDTA, 0.5 mm oxidized glutathione, 0.12 mm NADPH, and 10 μL plant sample. GR activity was measured by following the decrease in absorbance of oxidized glutathione at 340 nm (E = 6.2 mM⁻¹·cm⁻¹). SOD activity was determined according to the method of Rao and Stresty (2000). One unit of enzyme activity was defined as the amount of enzyme required to impose a 50% inhibition in the rate of nitro blue tetrazolium reduction measured at 560 nm.

**Statistical analysis.** Three T treatments were combined with three grafting combinations in a two-factorial design with two replications per T and five per grafting combination in each T replication. Data were subjected to a two-factorial analysis of variance. Factor effects were evaluated using Fisher’s F-test. When T or grafting combination had a significant impact but the interaction between them was not significant, the means between the three T levels tested and the three grafting combinations, respectively, were separated using Duncan’s multiple range test at a significance level of P ≤ 0.05. The same test was used to separate means of all nine treatments when the interaction was found to be significant. Statistical analyses were carried out using the STATISTICA software package Version 9.0 (StatSoft, Tulsa, OK).

**Results**

**Vegetative growth and root morphology.** Root fresh (Fig. 1A) and dry mass (Fig. 1B) of plants grafted onto LA 1777 were significantly larger than self-grafted ‘Kommeet’ plants at both low (14.6 °C) and intermediate T (17 °C). However, at optimum T (19.4 °C) and in self-grafted ‘Kommeet’, they remained the same, and thus no significant difference was found between them. Plants grafted onto ‘Moneymaker’ exhibited consistently lower root fresh and dry mass than those grafted onto the other two rootstocks at all T levels. Total root area (Fig. 1C) and total root length (Fig. 1D) were larger for self-grafted plants and those grafted onto LA 1777 than for plants grafted onto ‘Moneymaker’ at 14.6 and 17 °C. At 19.4 °C, the total root area of plants grafted onto LA 1777 was similar to those found in plants grafted onto ‘Moneymaker’.

Grafting ‘Kommeet’ onto the S. habrochaites accession LA 1777 increased fresh leaf mass (Fig. 1E), dry leaf mass (Fig. 1F), and total leaf area (Fig. 1G) at optimal or intermediate T in
comparison with self-grafted ‘Kommeet’ or grafting onto the cold-sensitive tomato ‘Moneymaker’. However, at low T, fresh and dry leaf mass and total leaf area of plants grafted onto LA 1777 decreased to similar levels with those measured in self-grafted ‘Kommeet’. Nevertheless, fresh and dry leaf mass and total leaf area measured in ‘Kommeet’ grafted onto ‘Moneymaker’ were also reduced at 14.6 °C, and thus they remained at significantly lower levels than in plants grafted onto LA 1777. Depending on differences reported previously and their extent at 14.6 and 17 °C, root/shoot ratio (Fig. 1H) was significantly higher in ‘Kommeet’ grafted onto LA 1777 than in the other two treatments. However, at 19.4 °C, the self-grafted ‘Kommeet’ exhibited a significantly higher root/shoot ratio than those grafted onto LA 1777.

**Yield Characteristics.** Total fresh and dry mass of fruit per plant and marketable fruit fresh weight were strongly reduced when plants were exposed to low T in comparison with intermediate or optimal T regardless of grafting treatment (Table 1). A decrease in T from 19.4 to 17 °C reduced only fresh and marketable fruit mass but had no significant impact on fruit dry mass. In contrast, average fresh weight of a tomato fruit was enhanced by reduction of T from 19.4 to 17 or 14.6 °C regardless of grafting treatment. Remarkably, intermediate T reduced appreciably fresh fruit mass affected by BER, whereas at 14.6 °C, no fruit with BER was recorded. No interaction was observed between T and rootstock genotype.

Total and marketable yield of ‘Kommeet’ grafted onto LA 1777 was significantly lower than that obtained from self-grafted
Table 1. Effects of three different temperatures (T) and tomato scion (S) ‘Kommeet’ (KO) self-grafted or grafted onto the rootstocks (R) LA 1777 (LA) or ‘Moneymaker’ (MM) on fruit fresh, dry, average and marketable mass, and fruit with blossom-end rot (BER).<sup>z</sup>

| Treatment | Fresh (g/plant) | Dry (g/plant) | Avg (g/fruit) | Marketable (g/plant) | BER (g/plant) |
|-----------|----------------|--------------|--------------|----------------------|---------------|
| T         |                |              |              |                      |               |
| Low       | 819 c          | 25.80 b      | 172 a        | 807 c                | 0.00 b        |
| Intermediate | 2589 b        | 84.25 a      | 142 b        | 2546 b               | 11.38 b       |
| Optimal   | 3261 a         | 93.08 b      | 104 c        | 2974 a               | 103.08 a      |
| R/S       |                |              |              |                      |               |
| LA/KO     | 1891 b         | 62.61 b      | 140          | 1806 b               | 8.07 b        |
| MM/KO     | 2160 ab        | 52.92 b      | 137          | 2053 ab              | 51.15 a       |
| KO/KO     | 2619 a         | 87.60 a      | 137          | 2467 a               | 55.24 a       |

Statistical significance

Table 2. Effects of three different temperatures (T) and tomato scion (S) ‘Kommeet’ (KO), self-grafted or grafted onto the rootstocks (R) LA 1777 (LA) or ‘Moneymaker’ (MM) on fruit and flower number and fruit/flower ratio (FR/FL).<sup>z</sup>

| Treatment | Fruit (no./plant) | Flowers (no./plant) | FR/FL (no./no.) |
|-----------|-------------------|---------------------|-----------------|
| T         |                   |                     |                 |
| Low       | 8.40 c            | 52.21               | 0.17 c          |
| Intermediate | 15.44 b          | 54.00               | 0.29 b          |
| Optimal   | 19.46 a           | 52.89               | 0.36 a          |
| R / S     |                   |                     |                 |
| LA/KO     | 10.13 b           | 52.71               | 0.19 c          |
| MM/KO     | 13.83 b           | 50.67               | 0.27 b          |
| KO/KO     | 19.34 a           | 53.73               | 0.36 a          |

Statistical significance

<sup>z</sup>Different letters within the same column indicate significant differences between the means according to Duncan’s multiple range test (P ≤ 0.05). ns, *, ** and *** indicate nonsignificant or significant differences at P ≤ 0.05, P ≤ 0.01, and P ≤ 0.001, respectively.
in plants grafted onto LA 1777 (Fig. 3C). Total proteins in leaves were restricted as T decreased without any significant difference between grafting treatments (Fig. 3D).

Exposure of tomato plants to 14.6 °C increased glucose, fructose, sucrose, and starch concentration of leaves when compared with 17 and 19.4 °C (Fig. 3E–H). At suboptimal T, leaf glucose, fructose, and sucrose concentrations were significantly higher in ‘Kommeet’ grafted onto LA 1777 than in self-grafted and plants grafted onto ‘Moneymaker’, other than fructose at 17 °C with no difference between the grafting treatments. Grafting onto LA 1777 reduced starch concentration in plant leaves growing at 14.6 °C. Furthermore, at 14.6 °C, fructose and sucrose concentration in leaves of plants grafted onto ‘Moneymaker’ was significantly higher than in self-grafted plants.

**TOTAL C, N, PROTEIN, AND CARBOHYDRATE CONCENTRATION IN FRUIT.** Total N in fruit of plants grafted onto ‘Moneymaker’ was low at 19.4 and 17 °C but increased appreciably at 14.6 °C, thereby reaching greater total N levels than the other two grafted treatments (Fig. 4A). Total C in fruit decreased as T was reduced from 19.4 to 14.6 °C without significant differences between grafting treatments (Fig. 4B). Reduction of T decreased C/N ratio in fruit of plants grafted onto ‘Moneymaker’ but had no impact on C/N ratio in fruit of self-grafted or plants grafted onto LA 1777 (Fig. 4C). The latter exhibited significantly lower fruit C/N ratios at 17 and 19.4 °C than self-grafted ‘Kommeet’ or plants grafted onto ‘Moneymaker’. When T was reduced from 19.4 to 14.6 °C, total protein concentration increased in tomato fruit from plants grafted onto ‘Moneymaker’ and self-grafted ‘Kommeet’, whereas in the latter, an increase was observed also at 17 °C.

Fig. 2. Effects of three different temperatures (T) and tomato ‘Kommeet’ (KO), self-grafted or grafted onto the rootstocks LA 1777 (LA) or ‘Moneymaker’ (MO), on total nitrogen (N) (A), total carbon (C) (B) and carbon/nitrogen ratio (C/N) (C), total protein (D), glucose (E), fructose (F), sucrose (G), and starch concentration (H) of roots. The values are means ± SE of 10 replications.
In contrast, fruit protein concentration was not influenced by T in plants grafted onto LA 1777. At optimal T, self-grafted ‘Kommeet’ exhibited the lowest fruit protein concentration in comparison with the other grafting treatments but this difference was reversed at intermediate T. Concentrations of glucose, fructose, and sucrose in fruit significantly increased with decreasing T in all grafting treatments (Fig. 4E–G). Fruit glucose, fructose, and sucrose concentrations in plants grafted onto LA 1777 were significantly higher than in the other two grafting treatments at all T levels, except fructose at intermediate and optimal T, which was similar in the other grafting treatment. At optimal T, plants grafted onto ‘Moneymaker’ exhibited lower fruit starch levels than self-grafted ‘Kommeet’ and plants grafted onto ‘Moneymaker’ (Fig. 4H). When T was reduced from 19.4 to 17 °C, fruit starch concentration increased to similar levels in all grafting treatments. However, at 14.6 °C, starch concentration decreased to lower levels than at 17 °C in fruit of self-grafted or plants grafted onto ‘Moneymaker’.

**Oxidation Characteristics, Total Amino Acids, and Antioxidant Enzyme Activities.** Decreasing T increased root MDA concentration when ‘Kommeet’ was either self-grafted or grafted onto ‘Moneymaker’ but had no significant impact when plants were grafted onto LA 1777 (Fig. 5A). As a result, at suboptimal T, root MDA was significantly lower in plants grafted onto LA 1777 than in self-grafted plants or those grafted onto ‘Moneymaker’. In contrast, MDA concentration in leaves was not influenced either by T or by the rootstock genotype (data not shown). GR activity in roots and leaves was not influenced...
by T or rootstock genotype (data not shown). In fruit, GR activity at 19.4 °C was significantly lower in plants grafted onto ‘Moneymaker’ than in the other two grafting treatments, whereas it was not influenced by T decrease from 19.4 to 17 °C (Fig. 5B). However, at 14.6 °C, GR activity in fruit increased in all grafting treatments, the increase being significantly higher in plants grafted onto LA 1777.

Total amino acid concentration in the roots tended to increase with decreasing T (Fig. 5C). The increase was significantly stronger in plants grafted onto LA 1777 than in self-grafted plants and those grafted onto ‘Moneymaker’. At intermediate T, total amino acid concentration in roots of plants grafted onto ‘Moneymaker’ was significantly lower than in those of self-grafted ‘Kommeet’. Total amino acid concentration in leaves was not influenced either by T or by the rootstock genotype (data not shown). Fruit total amino acid concentration increased in all grafting treatments with decreasing T, the increase being more profound as T was reduced from 17 to 14.6 °C (Fig. 5D). At intermediate and optimal T, lowest total amino acid concentration in fruit was measured in self-grafted ‘Kommeet’, whereas at 14.6 °C, fruit total amino acid concentration was significantly higher in plants grafted onto LA 1777 than in those grafted onto S. lycopersicum rootstocks.

Superoxide dismutase activity in roots of plants grafted onto LA 1777 was similar at all T levels (Fig. 6A). However, SOD activity in roots decreased as T was reduced from 19.4 to 17 °C (Fig. 6B). At intermediate and optimal T, lowest SOD root activity in fruit was measured in self-grafted ‘Kommeet’, whereas at 14.6 °C, SOD activity in roots of plants grafted onto LA 1777 was significantly higher than in those grafted onto S. lycopersicum rootstocks.

Fig. 4. Effects of three different temperatures (T) and tomato ‘Kommeet’ (KO), self-grafted or grafted onto the rootstocks LA 1777 (LA) or ‘Moneymaker’ (MO), on total nitrogen (N) (A), total carbon (C) (B) and carbon/nitrogen ratio (C/N) (C), total protein (D), glucose (E), fructose (F), sucrose (G), and starch concentration (H) of fruit. The values are means ± ses of 10 replications.
low T, no significant differences were found between these two treatments. SOD activity was significantly greater in leaves of self-grafted ‘Kommeet’ than in those of the other grafting treatments at 19.4 and 17 °C (Fig. 6B). However, at 14.6 °C, leaf SOD activity was similar in all grafting treatments. SOD activity in fruit was not influenced by T decrease from 19.4 to 17 °C and by grafting treatment at 19.4 °C but was significantly reduced by grafting onto ‘Moneymaker’ at 17 °C (Fig. 6C). At 14.6 °C, fruit SOD activity increased in plants grafted onto LA 1777, whereas it was not influenced by T in the other two grafting treatments. In self-grafted and plants grafted onto LA 1777, GPOD activity was raised by low T in comparison with intermediate or optimal T, the increase being more profound with the LA 1777 root genotype (Fig. 6D). In contrast, T had no impact on root GPOD activity in plants grafted onto ‘Moneymaker’. Low T reduced GPOD activity in leaves of ‘Kommeet’ grafted onto S. lycopersicum rootstocks, but not in those grafted onto LA 1777 which exhibited the highest GPOD levels at low T (Fig. 6E). Lowest GPOD activity at 19.4 °C was measured in fruit from plants grafted onto LA 1777. However, at 17 °C, fruit GPOD activity increased in plants grafted onto LA 1777 to similar levels as in plants grafted onto ‘Moneymaker’, whereas the values measured in self-grafted ‘Kommeet’ were significantly lower than in the other two grafting treatments (Fig. 6F). The further T decrease from 17 to 14.6 °C diminished GPOD activity in fruit from self-grafted plants and those grafted onto ‘Moneymaker’ but increased fruit GPOD activity in plants grafted onto LA 1777.

Discussion

In contrast to a previous report (Bloom et al., 2004), but in accordance with later results (Venema et al., 2008), LA 1777, an accession of S. habrochaites, proved to be a compatible rootstock for cultivated tomato (S. lycopersicum). Moreover, in this present study, the growing period after grafting lasted 21 weeks, including a 9-week harvesting period, longer than the 5-week harvest period used in the study of Venema et al. (2008). Thus, the present study is the first report on long-term compatibility of LA 1777 used as a rootstock for cultivated tomato.

As indicated by our data shown in Figure 1A–D, vegetative shoot growth of ‘Kommeet’ is unaffected by intermediate T (17 °C), whereas it was affected by long-term exposure to low T (14.6 °C) depending on rootstock genotype. Mild reductions of vegetative shoot growth of ‘Kommeet’ at low T, when used as a scion in grafted tomato, are in contrast to those previously...
reported. Venema et al. (2008) found strong reduction after a 2-week exposure of tomato grafted onto ‘Moneymaker’, self-grafted or grafted onto LA 1777, at 15 °C compared with 25 °C. One explanation for this difference in responses of tomato to low T is the genotype of the scion. It seems that ‘Kommeet’ as a scion is much more tolerant to suboptimal T than ‘Moneymaker’, which is considered cold-sensitive (Domínguez et al., 2005; Fernández-Muñoz et al., 1995). Genotypic differences in responses of tomato shoot growth to suboptimal T have been reported also by Paul et al. (1984). An additional and evident reason for the contrasting difference in responses of grafted tomato to 14.6 °C between our experiment and that of Venema et al. (2008) is the duration of exposure. It seems that short-term low T stress severely impairs vegetative growth of tomato, but in the long term, growth can partly recover as a result of physiological adaptation to stress conditions (Goulas et al., 2006).

Shortening of root length at low T, regardless of rootstock genotype, indicates that low T imposes formation of thicker roots in tomato, also reported by Kasper and Bland (1992). Nevertheless, our knowledge about key physiological mechanisms underlying the inhibitory effect of suboptimal root-zone T on the cell cycle and cell elongation in roots is currently insufficient (Walter et al., 2009). Rootstock genotype has a strong impact on root growth when the same cultivar is used as a scion, as indicated by a root biomass reduction of 50% when ‘Moneymaker’ was compared with LA 1777 or self-grafted even at optimal T. Optimal T causes stress on LA 1777 resulting in
reduced root mass. LA 1777 is a cold-tolerant accession adapted to T below the optimal treatment. Used as a rootstock, it responds at optimum T with a growth reduction also shown in previous investigations (Venema et al., 2008).

However, the most interesting finding of this study regarding vegetative tomato growth is that ‘Kommeet’ grafted onto LA 1777 exhibited improved shoot growth at intermediate and optimal T and root growth at low or intermediate T in comparison with self-grafted or plants grafted onto ‘Moneymaker’. These findings, in combination with the fact that suboptimal T did not restrict root length, while enhancing total root area in ‘Kommeet’ grafted onto LA 1777, indicate that this graft combination provides clear advantages in terms of vegetative tomato growth. Uptake of some nutrients and water is strongly affected by reducing root T in non-grafted S. lycopersicum but not in S. habrochaites but (Bloom et al., 1998, 2004). Change in root morphology may be interpreted as adaptation of nutrient acquisition mechanisms to low T, aimed at extending the absorbing surface area per unit root mass or length (Maduff et al., 1986). Thus, formation of a more extensive root system by tomato plants grafted onto LA 1777 as T is reduced, and the concurrent increase in root/shoot ratio, provides an additional advantage to these plants when exposed to suboptimal T in terms of nutrient and water uptake. An increase in root/shoot ratio has been interpreted by Equiza et al. (2001) as an adaptation aimed at overcoming restrictions in water absorption, which, in case of low T, might be related to increased water viscosity and root resistance to water transport.

Reducions in fruit yield when tomato is exposed to suboptimal T have been already reported by several investigators (Arroca et al., 2001; Korkmaz and Dufault, 2002; Van der Ploeg and Heuvelink, 2005). Suppression of yield at suboptimal T in our study originates from decreased fruit set owing to reduction in pollen quantity, quality (Dominguez et al., 2005), and in pollen quantity, quality (Domı´nguez et al., 2005), which restricts fruit number per plant. Our findings support this hypothesis because suboptimal T decreased total fruit yield by reducing total fruit number, whereas average yield rose. Increase of the average fresh fruit mass as T was lowered indicates that low T impairs more severely the formation and fecundity of pollen than the availability of assimilates after fruit setting, as also suggested by Adams et al. (2001) and Schwarz et al. (2010).

Another interesting finding of the present study is that reducing ambient T restricts the calcium-related physiological disorder BER in fruit. According to Kiegle et al. (2000), low T increases cytosolic Ca\(^{2+}\), whereas according to Knight et al. (1996) and Tanthiarju et al. (1997), Ca\(^{2+}\) could play an essential role in the cold acclimation process. As reported by Ho and White (2005), cytosolic Ca\(^{2+}\) coordinates cell expansion by influencing the incorporation of vesicles containing enzymes required for membrane and wall construction into the plasma membrane. Therefore, an increase in fruit calcium resulting from suboptimal T may prevent morphological defects such as membrane permeability that may result in abnormal leakage of solutes from cells and cell death and concurrently the occurrence of BER. Significant restriction of BER in fruit by grafting onto LA 1777 points to a relationship of Ca\(^{2+}\) transport to fruit. Further research is needed to elucidate the mechanisms underlying this relationship.

Grafting ‘Kommeet’ onto LA 1777 significantly restricted total fruit yield in comparison with grafting onto ‘Moneymaker’ and self-grafting as a result of a strong reduction of fruit numbers per plant at 27% and 48%, respectively. Because the flower number per plant was similar in all grafting treatments, it is concluded that use of LA 1777 as a rootstock impairs pollen fertility. Pollen produced by S. habrochaites is more cold-tolerant than that of S. lycopersicum (Domínguez et al., 2005). However, according to the same authors, S. lycopersicum × S. habrochaites crosses and subsequently segregating generations with S. lycopersicum show reduced pollen fertility as a result of interspecific hybrid sterility. The present study indicates that the combination of S. habrochaites and S. lycopersicum impairs pollen fertility not only if these related species are crossed, but also if the latter is grafted onto the former. These findings point to a role of plant hormones originating from roots and transported to shoot either on pollen viability or on any other function related to pollination and fruit setting (Garay-Arroyo et al., 2012; Sirova et al., 2011). Thus, in terms of yield performance, use of S. habrochaites as a rootstock may pose similar difficulties with those faced when using this wild tomato species for breeding of elite cultivars; i.e., reduced pollen fertility. Further research is needed to confirm this hypothesis.

Previous research has indicated that exposure of tomato to low T suppresses carbon export from leaves, thereby imposing accumulation of soluble carbohydrates in these plant organs (Rosa et al., 2009a; Strand et al., 2003). Restriction of sugar export from leaves of cold-stressed plants originates mainly from reduced sink activity resulting from exposure of roots, fruit, and stem apices to low T (Rosa et al., 2009b). Reduced C use in sink organs stressed by low T slows down the rate of sugar unloading, thereby imposing feedback retardation of sugar-loading in leaf phloem (Ainsworth and Bush, 2011). As a result, soluble carbohydrates may accumulate in both photosynthetically active leaves and sink organs such as roots and fruit, as was the case in the present study. Nevertheless, accumulation of soluble sugars in plant tissues stressed by low T may also serve as an adaptive mechanism to stress conditions. Plant metabolism is modulated under abiotic stress conditions and accumulated sugars may act as osmolytes or substrates for cellular respiration (Gupta and Knaur, 2005) or as antioxidative agents (Bogdanovic et al., 2008) that scavenge ROS (Noctor and Foyer, 1998). Thus, a stronger accumulation of soluble sugars in plant tissues exposed to suboptimal T may indicate a higher adaptability to cold-stress conditions. In agreement with this consideration, in our study, soluble carbohydrate levels in roots of plants grafted onto the cold-tolerant rootstock LA 1777 were higher than in roots of the other grafting combinations under low T.

Higher soluble sugar levels in leaves and fruit of plants grafted onto LA 1777 compared with those self-grafted or grafted onto ‘Moneymaker’ at 14.6 °C may predominantly reflect the stronger reduction of leaf biomass in the former compared with the latter two treatments when T dropped (Fig. 1A–D). However, higher soluble sugar levels at 17 °C in leaves and fruit of plants grafted onto LA 1777 compared with the other two grafting treatments were not accompanied by significant reductions in shoot biomass. Thus, it is reasonable to assume that at 17 °C increased levels of glucose and fructose in leaves and fruit of plants grafted onto LA 1777 compared with those from the other two grafting treatments served mainly as an adaptive mechanism to suboptimal T, which is triggered by roots. This consideration is in agreement with previous findings (Cook et al., 2004; Ma et al., 2009; Zhang et al., 2007) indicating that plants subjected to suboptimal T stress use.
sugars and amino acids as protective compounds. Protection of leaf growth and photosynthesis through hormonal signals originating from cucumber roots under cold stress conditions such as abscisic acid has been reported by Zhou et al. (2007). A link between signaling pathways of abscisic acid and sugar metabolism has been reported by several investigators in Arabidopsis thaliana (Arenas-Huertero et al., 2000; Cheng et al., 2002; Rook et al., 2001).

Striking differences in root N and C between ‘Kommee’ grafted onto LA 1777 and plants either self-grafted or onto ‘Moneymaker’ reflect presumably anatomical differences related to allocation of plant biomass into polysaccharides deposited in cell walls and cytosol. Reduced root and leaf protein as T decreased was obviously related to substantial increases of total amino acid concentration in roots and fruit. As reported by Mitchell and Madore (1992) and Zhang et al. (2007), plants subjected to suboptimal T use amino acids, in addition to sugars, as protective compounds. Accumulation of amino acids in plant tissues exposed to suboptimal T originates mainly from up-regulation of nitrate reductase activity (Tucker and Ort, 2002). Thus, it seems that the stronger increase of total amino acid concentration in roots of plants grafted onto LA 1777 and plants either self-grafted or onto ‘Moneymaker’ reflects the higher tolerance of LA 1777 to suboptimal T. Nevertheless, the ability of LA 1777 to increase total amino acid concentration when exposed to suboptimal T does not seem to be governed by root/shoot signaling compounds. Indeed, grafting of ‘Kommee’ onto LA 1777 did not result in a stronger increase of total amino acids in leaves and fruit in comparison with self-grafting or grafting onto ‘Moneymaker’.

Malondialdehyde is one of several low-molecular-mass end products arising from decomposition of various primary and secondary lipid peroxidation compounds (Demiral and Türkan, 2005). Therefore, an increased MDA level in plant tissue is considered a definite index of oxidative stress resulting in increased formation of ROS. The contrasting differences in root MDA levels between ‘Kommee’ plants grafted onto LA 1777 and those grafted onto S. lycopersicum rootstocks when exposed to 17 and 14.6 °C clearly demonstrate superiority of the former in avoiding oxidative stress caused by suboptimal T. Nevertheless, use of a cold-tolerant rootstock did not reduce the oxidative stress in leaves of tomato when plants were exposed to suboptimal T, as indicated by the similar leaf MDA levels in all grafting treatments at 17 and 14.6 °C. Plants are capable of deploying various enzymatic defense systems to minimize deleterious effects of ROS, which include the enzymes GPOD, GPOD, and SOD (Gill and Tuteja, 2010; Tuteja, 2009). As reported by Foyer and Noctor (2005), antioxidant capacity increases during cold acclimation in several plants as an adaptive mechanism to low T, and Noctor (2005), antioxidant capacity increases during cold-induced oxidative stress in roots of these plants. Furthermore, plants grafted onto LA 1777 exhibited significantly higher levels of soluble carbohydrates, total amino acids, and GPOD activity in leaves and fruit and superoxide dismutase in fruit at 14.6 °C. These results indicate that LA 1777 is capable of improving levels of some antioxidant compounds in tomato shoots, thereby improving its adaptation to 14.6 °C.

Taken together, results of the present study indicate that use of cold-tolerant accessions of S. habrochaites as tomato rootstocks can enhance tolerance of grafted plants to suboptimal T by improving their capacity to scavenge ROS. However, protection conferred by S. habrochaites against cold-induced oxidative stress may not be beneficial in terms of yield under suboptimal T conditions because this wild tomato relative may impair pollen fertility even when used as a tomato rootstock. Future research utilizing microarray analysis of accessions of S. habrochaites as rootstocks in a series of rootstock/scion combinations might elucidate genes controlling root/shoot signaling that influence pollen fertility. Acquired knowledge could be used to establish biomarkers to screen not only wild tomato genotypes serving as rootstocks, but also to test rootstock/scion combinations potentially tolerant to suboptimal T.

**Conclusion**

Grafting tomato ‘Kommeet’ onto the cold-tolerant accession LA 1777 of the wild relative of tomato S. habrochaites increases root biomass at low (14.6 °C) and intermediate T (17 °C), both referring to suboptimal T, and the leaf biomass at intermediate and optimum T (19.4 °C) in comparison with self-grafting and grafting onto the cold-sensitive ‘Moneymaker’. However, grafting onto LA 1777 significantly restricts yield as a result of significant suppression of fruit number per plant in comparison with grafting onto ‘Moneymaker’ and self-grafting while not influencing flower number per plant, which points to a root-induced restriction of fruit set. These results indicate that S. habrochaites impairs pollen fertility not only when used to breed interspecific hybrids with S. lycopersicum, but also when used as rootstock.

Increase of soluble carbohydrates, total amino acids, and GPOD activity in the roots of plants grafted onto LA 1777 and low MDA concentration at suboptimal T point to a much weaker cold-induced oxidative stress in roots of these plants. Furthermore, plants grafted onto LA 1777 exhibited significantly higher levels of soluble carbohydrates, total amino acids, and GPOD activity in leaves and fruit and superoxide dismutase in fruit at 14.6 °C. These results indicate that LA 1777 is capable of improving levels of some antioxidant compounds in tomato shoots, thereby improving its adaptation to 14.6 °C.

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