RNA-seq discovery, functional characterization, and comparison of sesquiterpene synthases from *Solanum lycopersicum* and *Solanum habrochaites* trichomes

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**Abstract** *Solanum lycopersicum* and *Solanum habrochaites* (f. *typicum*) accession PI127826 emit a variety of sesquiterpenes. To identify terpene synthases involved in the production of these volatile sesquiterpenes, we used massive parallel pyrosequencing (RNA-seq) to obtain the transcriptome of the stem trichomes from these plants. This approach resulted initially in the discovery of six sesquiterpene synthase cDNAs from *S. lycopersicum* and five from *S. habrochaites*. Searches of other databases and the *S. lycopersicum* genome resulted in the discovery of two additional sesquiterpene synthases expressed in trichomes. The sesquiterpene synthases from *S. lycopersicum* and *S. habrochaites* have high levels of protein identity. Several of them appeared to encode for non-functional proteins. Functional recombinant proteins produced germacrene, β-caryophyllene/α-humulene, viridiflorene and valencene from (E,E)-farnesyl diphosphate. However, the activities of these enzymes do not completely explain the differences in sesquiterpene production between the two tomato plants. RT-qPCR confirmed high levels of expression of most of the *S. lycopersicum* sesquiterpene synthases in stem trichomes. In addition, one sesquiterpene synthase was induced by jasmonic acid, while another appeared to be slightly repressed by the treatment. Our data provide a foundation to study the evolution of terpene synthases in cultivated and wild tomato.

**Keywords** Sesquiterpene synthase · Trichomes · RNA-seq · Wild and cultivated tomato · Jasmonate

**Introduction**

Terpenoids constitute a large class of chemical compounds produced by most, if not all, living organisms. Over 23,000 different terpenoid compounds have been characterized (Sacchettini and Poulter 1997). Plants produce terpenoids that function in primary metabolism such as phytohormones (abscisic acid, gibberellins, cytokinins, and brassinosteroids), are part of photosynthetic pigments (phytol and carotenoids), electron carriers (ubiquinone) or constitute structural components of membranes (phytosterol). However, the majority of plant terpenoids are secondary, or specialized metabolites, present only in a subset of plant lineages. They can be active as direct defensive compounds, such as phytoalexins that accumulate upon pathogen infection (Akram et al. 2008). In addition, volatile terpenoids can function as indirect defensive compounds by attracting predators or parasitoids of the attacking insect.
The emission of different terpenoids is induced by insect herbivory (Kant et al. 2004; Olson et al. 2008; Navia-Gine et al. 2009). Two distinct pathways leading to the universal terpene precursors isopentenyl diphosphate (IPP) and dimethylallyl diphosphate (DMAPP) operate in plants. The mevalonate (MVA) pathway produces IPP in the cytosol, which can be converted to DMAPP by IPP isomerase. The 2-C-methyl-d-erythritol 4-phosphate (MEP) pathway produces plastidial IPP and DMAPP. Head-to-tail elongation of IPP with DMAPP catalyzed by geranyl diphosphate (GPP) synthase or by neryl diphosphate (NPP) synthase lead to the formation of GPP and NPP (the (Z,Z)-isomer of GPP; Croteau and Karp 1979), respectively, which are the precursors for monoterpenes. (E,E)-Farnesyl diphosphate (FPP), and (Z,Z)-Farnesyl diphosphate (Kellog and Poulter 1997) serve as precursors for sesquiterpenes, and are synthesized by (Z,Z)- and (E,E)-FPP synthases, respectively. (E,E,E)-Geranylgeranyl diphosphate (GGPP) synthase catalyzes formation of GGPP, the diterpene precursor. The MEP pathway provides precursors for the synthesis of monoterpenes and diterpenes in plastids, whereas sesquiterpenes are derived from precursors of the MVA pathway in the cytosol. However, exchange of precursor between cytosol and plastids has been reported. Snapdragon flowers can synthesize sesquiterpenes from plastidial isoprenes, a very recently it was shown that several sesquiterpene synthases from wild tomato (Solanum habrochaites) are synthesized in the plastids from plastidial (Z,Z)-FPP (Salauag et al. 2009).

The prenyl diphosphates are converted to terpenes by the action of terpene synthases (TPSs), a group of structurally and evolutionarily related enzymes (Chen et al. 2011). Induced terpenoid synthesis is often correlated with induced expression of terpene synthases (Zulak et al. 2009; Navia-Gine et al. 2009; Herde et al. 2008). Besides regulation at the level of terpene synthases, induction of precursor biosynthetic genes has also been reported (Kant et al. 2004; Ament et al. 2004). In vitro, sesquiterpene synthases can often produce monoterpenes when provided with GPP as substrate, and monoterpenoid synthases can produce sesquiterpenes when provided with FPP. Therefore, subcellular targeting of terpene synthases determines which substrate the terpene synthase encounters. For instance, two nearly identical terpene synthases from snapdragon both catalyze the conversion of GPP to linalool and of FPP to nerolidol in vitro. However, only one of these linalool/nerolidol synthases has a transit peptide, localizing the protein to the plastids (Nagegowda et al. 2008).

To be effective against pests and diseases, defensive terpenoid compounds are often produced at the surface of the plant. Trichomes, which are specialized secretory structures on the surface of leaves and stems, contain high levels of terpenes in many species (Chatzivasileiadis et al. 1999; Besser et al. 2009), as well as other secondary metabolites (van Schie et al. 2007; Fridman et al. 2005; Ben-Israel et al. 2009). Furthermore, many investigations have shown that these compounds, including terpenes, are usually synthesized de novo in the trichomes (Maes et al. 2011; Olsson et al. 2009).

New terpene synthases are often discovered by homology-based cloning (van Der Hoeven et al. 2000; Portnoy et al. 2008; van Schie et al. 2007; Jones et al. 2008), by random sequencing of cDNAs (van Der Hoeven et al. 2000; Wang et al. 2008) followed by bioinformatic searches of the resulting EST databases (Keeling et al. 2011) or by genome mining (Aubourg et al. 2002; Martin et al. 2010). Terpene synthases that are less abundant in a particular organ or structure, or have low sequence similarity to known terpene synthases might not be identified by these methods. Therefore, we set out to use massive parallel pyrosequencing of the tomato trichome transcriptome in order to find new sesquiterpene synthases present in tomato stem trichomes. The use of massive parallel pyrosequencing of transcripts, termed “RNA-seq” is becoming more widely used (Schilmiller et al. 2010; Wilhelm and Landry 2009). The advantage of RNA-seq above the construction of EST databases is that quantitative expression levels of transcripts are better estimated due to the larger set of data.

Here we describe the identification and characterization of seven sesquiterpene synthase cDNAs from Solanum lycopersicum and six sesquiterpene synthase cDNAs from Solanum habrochaites, a wild tomato species, by using RNA-seq on cDNA from stem trichomes. Functional expression in E. coli provided information on product specificity of the proteins encoded by these cDNAs. Determination of tissue-specific expression of the cultivated tomato sesquiterpene genes showed that the expression of most of them was highest in tissue containing trichomes. Furthermore, one sesquiterpene synthase in the cultivated tomato was induced by jasmonic acid treatment, suggesting its involvement in herbivore-induced terpenoid emission.

Materials and methods

Plant material and mRNA isolation

Tomato plants (Solanum lycopersicum cultivar Money-maker and Solanum habrochaites accession PI127826) were obtained from Enza Zaden (Enkhuizen, The Netherlands) and grown in soil in a greenhouse with day/night temperatures of 23–18°C and a 16/8 h light/dark regime for...
4 weeks. Cuttings were made, which were placed in soil and grown for another 3 weeks. Trichomes from the stem and petioles were collected at the bottom of a 50 ml tube by vortexing frozen petiole and stem segments in liquid nitrogen as described before (van Schie et al. 2007). RNA of trichomes was isolated using the Qiagen RNeasy plant Mini kit according to the manufacturer’s protocol. Messenger RNA was isolated from the pool of total RNA using the PolyAtract mRNA isolation system III from Promega (Madison, Wisconsin, USA) according to the manufacturers’ instructions. Messenger RNA was isolated from total RNA with an efficiency between 0.56 and 0.81%. Terpenes stored in trichomes were sampled by transferring approximately 25 mg trichomes to a 20 ml glass vial containing 2 ml saturated CaCl₂ (5 M) buffered in 100 mM Na-Acetate at pH 4.5. Vials were capped immediately and kept at 8°C prior to sampling with a Solid Phase Micro Extraction fiber (SPME), as described below.

mRNA amplification and double stranded cDNA synthesis

The MessageAmp II aRNA amplification kit from Ambion (Austin, Texas, USA) was used to amplify trichome mRNA. An aliquot of 100 ng of mRNA was used as input for amplification. Amplification was carried out according to the manufacturer’s protocol. The yield of amplified antisense strand RNA was 171 µg for *S. lycopersicum* and 144 µg for *S. habrochaites*. The size ranged from ~400 to 3,000 bases.

First strand cDNA synthesis from the amplified RNA was carried out with random hexamer primers. Synthesis was performed in batches using 10 µg RNA in 20 µl reactions. Random primers at a final concentration of 62.5 nM were combined with amplified RNA and incubated at 70°C for 5 min in a thermo-cycler (Biometra, Göttingen, Germany) after which the sample was transferred to ice. Subsequently, RevertAid M-MulV H⁻ reverse transcriptase from Fermentas Life Sciences (St. Leon-Rot, Germany) in combination with the supplied buffer was added with 1 mM nucleotides. First strand cDNA synthesis was carried out at 42°C for 90 min. Directly thereafter, reactions were transferred to ice and second strand synthesis was performed using Fermentas RNase H and DNA polymerase I from *E. coli*. To each 20 µl reaction tube 8 µl of the supplied DNA polymerase I reaction buffer was added along with 1 unit RNase H and 30 units DNA polymerase I. The reaction volume was increased to 100 µl with water. All components were added cold and the second strand synthesis reaction was carried out at 15°C for 2 h. Double stranded cDNA was purified with the QIAquick PCR purification kit (Qiagen, Germany) and subsequently used as input for massive parallel pyrosequencing.

Massive parallel pyrosequencing and data analysis

The cDNA samples were analyzed by massive parallel pyrosequencing using a Genome Sequencer GS20 sequencing platform (Roche Applied Science). First, 6 µg of each sample was subjected to nebulisation. Further library preparation was performed according to the standard GS20 library preparation protocol as supplied by Roche Applied Bioscience. Emulsion PCR and bead enrichment were carried out according to the standard GS20 protocol. One full picotiterplate (PTP; 70 × 75 mm) with two regions was used. Enriched beads were divided over both regions. Sequencing was performed according to the manufacturer’s instructions (Roche Applied Science). GS20 data processing was performed on-rig using the standard GS software, resulting in an average read length of 89.7 nt, a total number of raw reads of 768,329 and a total number of Passed Filter reads 377,673.

The Passed Filter reads (377,673 ESTs) were cleaned from low quality regions using SeqClean (http://www.tigr.org/tdb/tgi/software/). Four sequences shorter than 40 nucleotides and 422 sequences that consisted mostly of low-complexity successions were discarded. The resulting 377,447 high quality sequences were analyzed using the TGI Clustering (TGICL), a software system for fast clustering of large EST datasets (http://www.tigr.org/tdb/tgi/software/). The parameters used for clustering allowed to group sequences sharing a minimum of 95% identity over at least 40 nucleotides with less than 10 bases of mismatched sequence at either end (Quackenbush et al. 2000). Initially the reads were searched for redundant sequences and after the megablast of the TGICL, one cluster with in total 77,050 ESTs was discharged. The reads of this cluster showed high similarity to chloroplast sequences. Computational limitations did not allow for contig building of a cluster with this redundancy. The resulting sequences were searched for repeats using the dicots repeat databases available at http://www.tigr.org/tdb/e2k1/plant.repeats/. However, the trichome-derived ESTs did not show significant similarity to these repeats, and the repeat masking step of the unigene creation was omitted. A second cycle of unigene creation was performed using the output of the first cycle as input. This process created a set of merged contigs constructed by merging less than 5% of the unigenes from the first run of TGICL. A third cycle of unigene creation did not result in additional merges. The output of the second cycle contained 18,918 unigenes with 2 or more ESTs per unigene and 42,601 singlets.

cDNA library construction and isolation of full length cDNAs of terpene synthases

cDNA libraries were constructed using the HybrizAP – 2.1 XR library construction kit and HybrizAP – 2.1 XR cDNA.
synthesis kit from Stratagene (Cedar Creek, Texas, USA). The size of the primary cDNA library was $3.0 \times 10^6$ pfu for *S. lycopersicum* and $9.9 \times 10^6$ for *S. habrochaites*. The primary libraries were amplified according to the manufacturer’s protocol. The amplified cDNA libraries were excised using the Mass Excision Protocol described by the manufacturer. Excised cDNA libraries were used to PCR amplify 3’ and 5’ fragments of cDNAs of interest identified previously in the pyrosequencing database. When necessary, re-amplification was performed using nested primers. After sequence verification, the resulting full length candidate-sesquiterpene synthases were cloned in-frame into the pGEX-KG expression vector (Guan and Dixon 1991).

In addition to the cDNAs from *S. habrochaites* accession PI127826 as described above, we obtained cDNAs from a different *S. habrochaites* accession (PI126449) using a published EST database (Fridman et al. 2005). Sequence data can be found in the GenBank database (http://www.ncbi.nlm.nih.gov/) under accession numbers: ShTPS9, JN402388; ShTPS12, JN402389; ShTPS14a, JN402390; ShTPS17, JN402391; ShTPS14b, JN402392; ShTPS15b, JN402393; SITPS16, JN402394; SITPS17, JN402395; SITPS31, JN402396; ShTPS15a, JN402397; ShTPS16, JN402398; SITPS15, JN402399.

Functional expression analysis

Expression constructs were transformed to C41 (DE3) electro competent *E. coli* cells (Dumon-Seignovert et al. 2004). From a single colony an overnight culture was grown at 37°C of which 500 μl was inoculated in 50 ml Terrific Broth containing 100 μg ml⁻¹ ampicillin. The culture was grown to an OD₆₀₀ of 0.5–0.6 at 37°C after which they were placed at 4°C for 30 min. Protein expression was induced with 1 mM isopropyl-β-D-1-thiogalactopyranoside (IPTG; Roche, Basel, Switzerland). An empty plasmid was taken along as a negative control. After incubation for 16 h at 16°C at 200 rpm, cells were harvested by centrifugation at 4,500 rpm at 4°C for 15 min. The supernatant was removed and the pellet was resuspended in 2 ml of assay buffer (25 mM HEPES, pH 7.2, 10 mM MgCl₂, 10% (v/v) glycerol) with added lysozyme (1 mg ml⁻¹) and protease inhibitors (Complete, EDTA-free; Roche, Basel Switzerland). An empty plasmid was taken along as a negative control. After incubation for 16 h at 16°C at 200 rpm, cells were harvested by centrifugation at 4,500 rpm at 4°C for 15 min. The supernatant was removed and the pellet was resuspended in 2 ml of assay buffer (25 mM HEPES, pH 7.2, 10 mM MgCl₂, 10% (v/v) glycerol) with added lysozyme (1 mg ml⁻¹) and protease inhibitors (Complete, EDTA-free; Roche, Basel Switzerland). Cells were incubated on ice for 30 min after which they were sonicated. Lysates were centrifuged at 4°C for 25 min at 12,000 g. The supernatant was aliquoted and stored at −80°C.

Activity assays were performed in 20 ml glass vials in a total volume of 500 μl containing 50 mM HEPES, pH 7.2, 100 mM KCl, 7.5 mM MgCl₂, 20 μM MgCl₂, 5% (v/v) glycerol, 5 mM DTT with 50 μl protein extract and either 2 mM (*E*,*E*)-FPP (*E*-*E*-farnesyl diphosphate), (*Z*,*Z*)-FPP (2Z-6Z-farnesyl diphosphate) or GPP (geranyl diphosphate) as substrate (Echelon Biosciences Incorporated, Salt Lake City, USA). Vials were immediately closed with a Teflon lined crimp cap and incubated under moderate shaking for 1 h at 30°C. Enzyme products were sampled with a Solid Phase Micro Extraction fiber (SPME) for 10 min after the vial had been agitated and heated to 50°C. The fiber was desorbed for 1 min in an Optic injector port (ATAS GL Int. Zoeterwoude, NL) which was kept at 250°C. Compounds were separated on a DB-5 column (10 m × 180 μm, 0.18 μm film thickness; Hewlett Packard) in an 6890 N gas chromatograph (Agilent, Amstelveen, NL) with a temperature program set to 40°C for 1.5 min, ramp to 250°C at 30°C per minute and 250°C for an additional 2.5 min. Helium was used as a carrier gas, the column flow set to 3 ml per minute for 2 min, and to 1.5 ml per minute thereafter. Mass spectra were generated with the ion source set to −70 V at 200°C and collected with a Time-of-Flight MS (Leco, Pegasus III, St. Joseph, MI, USA) at 1,850 V, with an acquisition rate of 20 scans per second. Because some terpenes, such as germacrenes, are prone to thermal conversion (Colby et al. 1998; Faraldos et al. 2007) the enzyme assays were also extracted with pentane which was injected at 50°C in the injection port. For this, 500 μl lysate was assayed in the presence of 5 mM DTT and either (*E*,*E*)-FPP, (*Z*,*Z*)-FPP or GPP as a substrate, overlaid with 2 ml pentane (Sigma). After 1 h of incubation at 30°C the pentane layer was transferred to a 2 ml glass vial and concentrated on ice under nitrogen gas, to a final volume of 50 μl. Terpenoids were analysed by injection of 2 μl into the Optic injection port (ATAS GL international) at 50°C, subsequently heated to 275°C at a rate of 4°C s⁻¹ followed by gas-chromatography and mass-spectrometry as described by Bleeker et al. (2009). Boiled protein extract could not convert any of the precursors and was taken along as a negative control. No products were formed with GGPP as substrate. Terpene products were identified using standards when possible or by comparing mass spectra, and Kovats Index (Adams 2002).

The enzymatic assays for ShTPS15 from PI126449 and SITPS31 from cultivar M82 and the analyses of their products were performed as described in Falara et al. (in press).

Expression analysis

RNA was isolated using TRIzol (Invitrogen, Paisley, UK) according to the manufacturer’s instructions, from different tissues of 4-week-old *S. lycopersicum* plants. RNA was isolated from leaves, whole stem (stem pieces with trichomes), bald stem (stem pieces with trichomes removed), stem trichomes, root, fruit and flowers. Additionally, RNA was isolated from stem trichomes of 4-week-old *S. lycopersicum* plants 24 h after spraying with either 1 mM...
jasmonic acid (Ducheva Biochemicals) in tap water containing 0.05% SilwetL-77, or water and SilwetL-77 alone.

The RNA was DNase treated (TURBO DNase kit, Ambion) and the quantity determined with a Nanodrop spectrophotometer (ND-1000, Thermo Scientific). First strand cDNA was synthesized from 1.5 μg of total RNA using the RevertAid kit from Fermentas according to the manufacturers’ instructions. For expression analysis the cDNA equivalent of 100 ng RNA was used as template with the SYBR Green PCR qPCR SuperMix UDG (Invitrogen) and 300 nM of each primer and dispersed as 20 μl on a 96-well optical reaction plate (Applied Biosystems). PCRs were performed in the ABI 7500 Real-Time PCR System (Applied Biosystems). The specificity of the reaction was verified by dissociation analysis. Expression of ACTIN (SGN-U579547) was used to normalize and correct for variance in quality of RNA and quantity of input cDNA. Primer pair efficiencies were estimated by analysis of amplification curves of a standard cDNA dilution range. Three biological replicates were analyzed individually and statistical significance was tested by ANOVA.

Results

Sequences of sesquiterpene synthases from Solanum lycopersicum and Solanum habrochaites show high levels of identity

Massive parallel pyrosequencing of the trichome transcriptomes resulted in 195,377 reads from cDNA derived from S. lycopersicum and 182,386 reads from S. habrochaites accession PI127826, both with an average length of 80 nucleotides. About 0.20% of all reads had high sequence similarity to known sesquiterpene synthases for S. lycopersicum versus 0.12% for S. habrochaites reads (Table 1). This resulted in the identification of multiple TPS sequences belonging to the TPS-a clade according to the phylogeny of the plant TPS family (Bohlmann et al. 1998; Chen et al. 2011).

Recently we have analyzed the nearly completed genomic sequence of S. lycopersicum, and have identified a total of 44 TPS genes (Falara et al. in press). This analysis also showed that the majority of TPS genes in the tomato genome appear to encode sesquiterpene synthases and most of these putative sesquiterpene synthases belong to the TPS-a clade. In the S. lycopersicum database, transcripts for TPS9 (previously shown to encode germacrene C synthase; van Der Hoeven et al. 2000), TPS12 (previously shown to encode β-caryophyllene/α-humulene synthase; van Der Hoeven et al. 2000), TPS15, TPS16 and TPS17 were identified, as well as low numbers of reads for TPS31 (TPS31 was formerly known as LeVS2 (GenBank: AAG09949.1), but the function of LeVS2 was not established). It was noted that the full-length cDNA of TPS15 has a premature stop at position 228 and is predicted to encode a non-functional protein of 76 amino acids.

Multiple putative sesquiterpene synthases from S. habrochaites accession PI127826 were also observed, and full-length cDNAs for these genes were isolated. Based on phylogenetic analysis of their DNA sequences (Fig. 1), these cDNAs came from the TPS9, TPS12, TPS15, and TPS17. Again, TPS15 had several premature stop codons in the 5’ region of the cDNA but could putatively encode a shorter protein of 499 amino acids if the ATG at position 157 is functional as start codon, albeit this protein is unlikely to have sesquiterpene synthase activity. We were able to use the primers for S. lycopersicum TPS16 to amplify the S. habrochaites TPS16 cDNA as well. However, the ShTPS16 cDNA contained a single nucleotide deletion at position 459, a mutation that results in a premature stop at amino acid 153, thus ShTPS16 is predicted to encode a non-functional enzyme.

Searching the trichome EST database of S. habrochaites accession PI126449, a line whose trichomes contain high levels of methylketones but also some sesquiterpenes (Fridman et al. 2005), identified a sequence in the TPS-a clade with high similarity to SITPS15 (Suppl. Fig. 1) and additionally a sequence with high similarity to SITPS14 (Falara et al. in press). The full-length cDNA of the ShTPS15 had an open reading frame of 540 codons without any apparent mutations. Apart from the sequence of TPS14 from S. lycopersicum, expressed primarily in roots (Fig. 7), we were able to isolate a full-length cDNA of ShTPS14 from stem trichomes of accession PI127826 and flowers of accession PI126449. These two ShTPS14 cDNAs had an open reading frame putatively encoding proteins of 554 amino acids.

Phylogenetic analysis (Fig. 1) of the newly identified and previously characterized tomato TPSs shows that SITPS9 clusters together with previously identified germacrene synthases from cultivated tomatoes (SIVFNT, SSTLE1, SSTLE2) and ShTPS9 with germacrene synthases from S. habrochaites accession LA1777 (SSTLH1 and SSTLH2). It also indicates that each TPS from

| TPS    | S. lycopersicum | S. habrochaites |
|--------|----------------|----------------|
|        | Reads | %    | Reads | %    |
| TPS9   | 300   | 1.500| 123   | 0.615|
| TPS12  | 10    | 0.050| 54    | 0.270|
| TPS15  | 42    | 0.210| 58    | 0.290|
| TPS16  | 6     | 0.030| ND    | ND   |
| TPS17  | 25    | 0.125| 5     | 0.025|
| TPS31  | 3     | 0.015| ND    | ND   |
| SUM    | 386   | 2.200| 240   | 1.200|
S. lycopersicum has a higher similarity to its respective ortholog in S. habrochaites accession PI127826 than to other tomato TPSs. Both monoterpene synthases TPS5 and TPS4 (formerly MTS1 and 2; van Schie et al. 2007) cluster together in the phylogenetic tree, as well as santalene/bergamotene (sesquiterpene) synthase (ShSBS) from S. habrochaites, which accepts (Z,Z)-FPP as a substrate (Sallaud et al. 2009), and phellandrene (monoterpene) synthase (TPS20, formerly SlPHS1) from cultivated tomato, which mainly uses NPP as substrate (Schilmiller et al. 2009). The relationships between the different TPSs were confirmed by a phylogenetic tree based on the amino acid sequences (Suppl. Fig. 2). Alignment of the deduced amino acid sequences of TPS9, TPS12, TPS16 and TPS17, depicted in Supplemental figure 3, shows how closely related TPS9 and TPS12, and TPS16 and TPS17, respectively, are. All proteins contained the RR/P8xW motive found in most terpene synthases, which is involved in hydrolysis of the pyrophosphate group and the DDxxD motive, involved in cofactor binding.

Enzymatic activity of recombinant sesquiterpene synthases

Proteins isolated from E. coli cells expressing the putative sesquiterpene synthases were assayed for their ability to convert (E,E)-FPP into sesquiterpenes. The protein encoded by the Moneymaker allele of SlTPS9 produced germacrene C, and minor amounts of germacrene A, B and D (Fig. 2a), similar to the protein encoded by this locus in cultivar VFNT Cherry tomato. ShTPS9 catalyzed the formation of mostly germacrene B and minor amounts of A and C (Fig. 2b), similar to the products observed in the reaction catalyzed by a very similar protein, designated SSTLH1 encoded by a cDNA isolated from S. habrochaites accession LA1777 (van Der Hoeven et al. 2000).
recombinant TPS12 proteins had $\beta$-caryophyllene/$\alpha$-humulene synthase activity (Fig. 3a, b).

SITPS17 and ShTPS17, which are 98.6% identical and 99.1% similar on the protein level, produced both mostly valencene from ($E,E$)-FPP and also an unidentified sesquiterpene, besides azulene and $\alpha$- and $\beta$-farnesenes (SITPS17; Fig. 3c) or $\beta$-farnesene (ShTPS17; Fig. 3d). As predicted, SITPS15 and ShTPS15 from accession PI127826 had no activity, but the ShTPS15 allele isolated from accession PI126449 encoded a protein that catalyzed the formation of mostly germacrene A (Fig. 4). We were unable to detect activity of SITPS16 although a soluble protein of the correct size was expressed in E. coli upon IPTG induction (data not shown). Remarkably, all recombinant proteins that could use ($E,E$)-FPP as substrate could also use ($Z,Z$)-FPP as substrate (Suppl. Fig. 4). SITPS9 made mostly germacrene C from both ($E,E$)-FPP as well as ($Z,Z$)-FPP whereas ShTPS9 produced germacrene B and $\alpha$-humulene with ($Z,Z$)-FPP (Suppl. Fig. 4a,b). SITPS12 and ShTPS12 both made curcumene and $\beta$-bisabolene from ($Z,Z$)-FPP (Suppl. Fig. 4c,d). SITPS17 and ShTPS17 made various bisabolenes when activity assays were done with ($Z,Z$)-FPP (Suppl. Fig. 4e,f).

Although the TPS14 proteins only differ in a few amino acids from each other (Suppl. Fig. 5), their capacity to use ($Z,Z$)- or ($E,E$)-FPP as substrate differed substantially. SITPS14 could use either ($E,E$)-FPP or ($Z,Z$)-FPP equally well to make mostly ($Z$)-$\beta$-bisabolenes or $\alpha$-bisabolenes, respectively (Falara et al. in press). Interestingly, ShTPS14 from PI126449 appears to have no clear preference for either ($Z,Z$)-FPP or ($E,E$)-FPP. In the presence of ($Z,Z$)-FPP the enzyme made predominantly ($Z$)-$\beta$-farnesene, $\alpha$- and $\beta$-acordadiene and $\alpha$- and $\gamma$-bisabolenes, while when assayed with ($E,E$)-FPP $\alpha$-cederene, ($Z$)-thujopsene, $\beta$-farnesene, $\beta$-acordadiene and $\beta$-bisabolene were produced (Fig. 5). ShTPS14 from PI127826 on the other hand clearly favored ($E,E$)-FPP to synthesize $\beta$-farnesene and $\alpha$- and $\beta$-bisabolenes (Fig. 5a, b). SITPS31 predominantly made viridiflorene from ($E,E$)-FPP (Fig. 6).

All sesquiterpene synthases produce monoterpens from GPP

All sesquiterpene synthases were assayed for monoterpene synthase activity with GPP as substrate. Though not at high efficiency, they all converted the GPP precursor to a range of
most simple monoterpenes (Suppl. Fig. 6). TPS9 and TPS12 produced mostly β-myrcene, limonene and low amounts of terpinolene. In the assays with TPS17 and ShTPS14 of PI126449, additionally the monoterpenes (Z)-β-ocimene, (E)-β-ocimene and linalool were produced (Suppl. Fig. 6).

Expression of *S. lycopersicum* sesquiterpene synthases is differentially regulated

To investigate in which tissues the active *S. lycopersicum* sesquiterpene synthases TPS9, TPS12, TPS14, TPS17 and TPS31 were transcribed, we dissected mature *S. lycopersicum* plants for RNA isolation and subsequent quantitative RT-PCR. Expression of all terpene synthases was lowest in fruits (Fig. 7). TPS9 and TPS17 displayed similar expression patterns, with highest expression in stem trichomes (Fig. 7a, d). TPS12 expression was highest in leaves (Fig. 7b), most likely in the leaf trichomes as shown for SlTPS12 in the cultivar M82 (Schilmiller et al. 2010). In S. lycopersicum, TPS14 is expressed mostly in the roots (Fig. 7c), while in the *S. habrochaites* accession PI127826 TPS14 transcripts were found in trichomes (data not shown). Expression of TPS31 was very low overall, but highest in stem trichomes and leaves (Fig. 7e), most likely in their trichomes.

Differential expression in response to jasmonate treatment

Since jasmonic acid (JA) treatment resulted in the induction of *SIMTS1* (SITPS3) in stem trichomes (van Schie et al. 2007), we tested whether expression of *SITPS9*, *SITPS12*, *SITPS17* or *SITPS31* was induced by JA treatment. Moneymaker plants that were 4 weeks old were treated with 1 mM JA or left untreated as a control, RNA was isolated after 24 h, and the levels of transcripts were measured by qRT-PCR (Fig. 8). Only the trichome-specific expression of *SITPS31* was significantly induced, approximately threefold by JA treatment, just as the positive control *SIMTS1* (SITPS3, data not shown). Interestingly, expression of *SITPS17* appeared to be reduced roughly twofold in JA-treated plants ($P = 0.059$).

Discussion

Large scale transcript sequencing as tool for gene discovery

The presence and biosynthesis of sesquiterpenes in *S. lycopersicum* and *S. habrochaites* had been previously investigated (Colby et al. 1998; Schilmiller et al. 2010; van Der Hoeven et al. 2000). Colby et al. (1998) isolated a cDNA from the cultivar VNFT Cherry and showed that it encodes germacrene C and van der Hoeven et al. (2000) showed that synthesis of germacrene C, β-caryophyllene and α-humulene was controlled by a locus on chromosome...
Using the VFNT cDNA of germacrene C as a probe to screen cDNA libraries, van der Hoeven et al. (2000) also isolated two cDNAs, designated SSTLE1 and SSTLE2, that were very similar to VNFT germacrene C. Based on the sequences of both cDNAs, we can conclude that they are different alleles (or one of them may be a cloning artifact) of SlTPS9, which we now know to be located on chromosome 6 (Falara et al. in press). Van der Hoeven et al. (2000) also identified two alleles of TPS9 from S. habrochaitea accession LA1777, designated SSTLH1 and SSTLH2, whose cDNAs encode germacrene B and germacrene D, respectively.

Subsequently, Schilinger et al. (2010), using a proteomic approach, identified a protein sequence in S. lycopersicum cv. M82 trichomes that catalyzed the formation of \( \beta \)-caryophyllene and \( \alpha \)-humulene, and was therefore designated as CAHS (\( \beta \)-caryophyllene/\( \alpha \)-humulene synthase). Our data indicate that TPS12 encodes CASH. Here we used massive parallel pyrosequencing as an approach for terpene synthase transcript discovery in a specialized tissue type enriched for plant secondary metabolites, the trichomes of

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Fig. 7 Tissue specific expression of SlTPSs. Relative transcript levels for a SlTPS9, b SlTPS12, c SlTPS14, d SlTPS17 and e SlTPS31 as determined by Q-RT-PCR. Mean values (+SE) of 3 biological replicas are shown, normalized for Actin expression. L leaf, WS whole stem, BS bald stem, T stem trichomes, R root, Fr fruit, Fl flower.

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Fig. 8 JA induction of SlTPSs expressed in the stem trichomes. Relative transcript levels for a SlTPS9, b SlTPS12, c SlTPS17 and d SlTPS31 as determined by Q-RT-PCR. Expression in isolated stem trichomes of control (C) and JA-treated plants (JA) are shown as mean values (+SE) of 3 biological replicas, normalized for Actin expression. Asterisks indicate significant difference (\( P < 0.05 \)).
wild and cultivated tomato plants. Using this approach, we were able to identify transcripts of not only TPS9 and TPS12 but also of TPS14, TPS15, TPS16, TPS17 and TPS31 in the trichomes of *S. lycopersicum*, *S. habrochaites*, or both.

Over representation of germacrene synthases

Germacrene synthases have been isolated from species such as poplar (Arimura et al. 2004), goldenrod (Prosser et al. 2004), kiwi fruit (Nieuwenhuizen et al. 2009), melon (Portnoy et al. 2008), *Cistus creticus* (Falara et al. 2008) and many others. Often when a gene discovery approach is used to find new terpene synthases, in vitro functional assays show that multiple genes in a species encode germacrene synthases. For example, a degenerate primer strategy to find sesquiterpenes in lettuce resulted in the identification of only two germacrene A synthases (Bennett et al. 2002). Recently five sesquiterpenes from the fungus *Coprinus cinereus* were isolated and two of these recombinant sesquiterpene synthases catalyze the formation of germacrene A (Agger et al. 2009). Similarly, two out of three characterized sesquiterpene synthases from sunflower were identified as germacrene A synthases (Gopfert et al. 2009). The observation that three of the sesquiterpene synthases from tomato that we and others have characterized—STPS9 and ShTPS9, and ShTPS15 from accession PI126449—produce germacrenes with FPP as the in vitro substrate is remarkable, since only minor amounts of volatile germacrenes are present in trichomes (Kang et al. 2010; Paetzold et al. 2010; van Der Hoeven et al. 2000) or emitted from tomato (Ament et al. 2004; Kant et al. 2004). It has been postulated that germacrene A is an intermediate in *epi*-aristolochene synthase catalysed by tobacco 5-epi-aristolochene synthase (Rising et al. 2000). Moreover, certain sesquiterpenes in *Medicago trunculata* were proven to be generated predominantly via protonation of the neutral intermediate, germacrene D, as opposed to synthesis directly via FPP isomerization to nerolidyl diphosphate (Garms et al. 2010). Likewise, Selina-3,7(11)-diene (Table 2) is the result of a C6 protonation of germacrene B (Davis and Croteau 2000). Germacrene A can also be derivatized by other enzymes as is suggested for germacrene A from lettuce (Bennett et al. 2002). Therefore, in planta assays are necessary to determine whether these enzymes are bona fide germacrene synthases.

All sesquiterpene synthases accepted both (E,E)-FPP and (Z,Z)-FPP as substrates

TPS14 synthases from both *S. habrochaites* accessions catalyzed the formation of multiple terpenes from (E,E)-FPP and (Z,Z)-FPP, including farnesenes, acoradienes and a variety of bisabolenes (Fig. 5). However, it appears that TPS14 from accession PI127826 exhibited a preference for the (cytosolic) (E,E) isofom of FPP, whereas intriguingly, the TPS14 from accession PI126449 appeared to have an equally high affinity for (Z,Z)-FPP instead. When each ShTPS14 was assayed with both FPP-isomers present, compounds derived from both isomers were identified for ShTPS14 PI126449 whereas the ShTPS14 PI127826 products were identical to the ones seen when the enzyme was incubated with (E,E)-FPP alone (data not shown). In *S. lycopersicum*, TPS14 accepted (E,E)-FPP and (Z,Z)-FPP equally well, producing mostly β-bisabolenene and α-bisabolene, respectively (Falara et al. in press).

ShTPS14 makes multiple products, ((E)-α-bisabolenene, γ-bisabolenes and β-bisabolenes and β-farnesenes; Fig. 5) which, most likely, are direct products of deprotonation of the bisabolyl cation (C6-ring) formed from FPP via nerodiol diphosphate, without other intermediates (Davis and Croteau 2000). Interestingly, the TPS14 proteins appear to contain two aspartate-rich DDXxD Mg2+-binding motifs (305 and 527; Suppl. Fig. 5) as well as a ‘protonation-initiated’ mechanism catalytic DxDD motif (in ShTPS14) (104; Suppl. Fig. 5), both of which are common in di- and tri-terpene cyclases. The presence of a second DDXxD motif could be responsible for alternate orientations of the substrate and therefore synthesis of multiple products (Davis and Croteau 2000). TPS9, 12 and 17 also accepted (Z,Z)-FPP as substrate (Suppl. Fig. 4). Our observations of sesquiterpene synthases reacting with substrates other than the canonical (E,E)-FPP adds additional weight to previous intimations of the flexibility of these enzymes. For example, Jones et al. (2011) reported a cytosolic sesquiterpene synthase from sandalwood able to use both (E,E) and (Z,Z) isomers of FPP to produce similar compounds. Such observations from multiple species suggest that these properties of the enzyme are not in vitro artifacts but might have in vivo relevance. However, whether (Z,Z)-FPP is available in the cytosol of the *Solanum* trichomes is not yet known, although evidence has been presented that *S. habrochaites* accession LA1777 produces (Z,Z)-FPP in the plastids (Sallaud et al. 2009).

Albeit with low efficiency, all TPSs described here were able to convert GPP to (mono) terpene products (Suppl. Fig. 6) indicating a level of plasticity of the active pocket of the protein. While some sesquiterpene synthases that are restricted to the use of a single substrate even with regard to precursors of the same size ((E,E)-FPP or (Z,Z)-FPP; Besser et al. 2009), there are examples of terpene synthases that can accommodate both GPP as well as (E,E)-FPP in a productive manner (van Schie et al. 2007) or even GGPP as a third possible substrate (Martin et al. 2010; Arimura et al. 2008). It has been proposed that trace amounts of GPP are present in the cytosol, and minor amounts of (E,E)-FPP are
available to plastid-localised terpene synthases (Aharoni et al. 2006; Wu et al. 2006). Hence, the active site plasticity of some TPS to accommodate isoprenyl diphosphates of different chain length may be biologically relevant.

Correlations between TPS transcript abundance, TPS products and sesquiterpene production in trichomes

Despite the fact that monoterpenes appear to dominate *S. lycopersicum* volatile emissions (Buttery et al. 1987; Bleeker et al. 2009), the majority of TPS genes mined from the tomato genome are sesquiterpene synthases (Falara et al. in press). Based on the in vitro functional characterization of the group of sesquiterpene synthases whose transcripts we detected in stem trichomes in *S. lycopersicum* and *S. habrochaites*, the presence of most, but not all of the sesquiterpenes observed in the respective glands can be explained (Table 2). Most notably, we did not find any terpene synthase that could produce zingiberene, the most abundant sesquiterpene in *S. habrochaites* PI127826 (Bleeker et al. 2009). Also, trichomes of *S. lycopersicum* contain, albeit at minor amounts, the sesquiterpenes α-copaene and cuparene that are unaccounted for in the SIPTS enzymes assayed here.

Since some sesquiterpenes are made by more than one sesquiterpene synthases in the same species, it is difficult to determine the direct contribution of each of them to the observed mixture even when the expression levels of individual sesquiterpene synthases are examined in detail.

Table 2 Sesquiterpenes present in trichomes of *S. lycopersicum* cv. Moneymaker and *S. habrochaites* PI127826

| Sesquiterpene          | *S. lycopersicum* | *S. habrochaites* |
|------------------------|-------------------|-------------------|
| Azulene                | x                 |                   |
| α-Copaene              | x                 |                   |
| β-Elemene              |                   | x                 |
| Caryophyllene          | x                 | x                 |
| γ-Elemene              | x                 | x                 |
| Alpha humulene         | x                 | x                 |
| β-Farnesene            |                   | x                 |
| β-Acoradiene           |                   | x                 |
| Curcumene              |                   | x                 |
| Germacrene D           | x                 | x                 |
| Zingiberene            |                   | x                 |
| Cuparene               | x                 |                   |
| β-Bisabolene           |                   | x                 |
| β-Sesquiphellandrene   |                   | x                 |
| Valencene              |                   | x                 |
| Germacrene C           | x                 |                   |
| Selina-3,7(11)-diene   |                   | x                 |
| Germacrene B           |                   | x                 |

Because we used non-normalized cDNAs in the RNA-seq experiments, we were able to analyze the data output for comparison of transcript levels of the different genes between *S. lycopersicum* and *S. habrochaites*, and the results indicate that overall abundance of sesquiterpene synthase transcripts did not correlate with the total amount of emitted sesquiterpenes, as *S. habrochaites* PI127826 emits over 50 μg g⁻¹ FW sesquiterpenes in 24 h (Bleeker et al. 2009) whereas *S. lycopersicum* plants only emits 0.7 μg sesquiterpenes g⁻¹ FW per day and total read abundance in the RNA-seq was 240 versus 383 respectively (Table 1). Although transcript abundance need not be translated into protein abundance, another explanation for the low sesquiterpene content of *S. lycopersicum* might be found in low precursor biosynthesis. Quantitative RT-PCR on cDNA derived from stem trichomes of *S. lycopersicum* and *S. habrochaites* LA1777 showed that expression of HMG-CoA reductase, a key enzyme in the MEP pathway, is approx. sevenfold higher in *S. habrochaites* (Besser et al. 2009). Our own RNA-seq data also show that transcript levels of HMG-CoA reductase are much higher (180 reads) than in *S. habrochaites* than in *S. lycopersicum* (24 reads). It is possible that the in planta function of terpene synthases may differ from the in vivo function we found in our enzymatic assays though a more likely explanation for the discrepancy between the total sesquiterpene emission and the level of expression of synthases found might be that we have not annotated all sesquiterpene sythase sequences in our dataset. In this study we have confined our analysis of sesquiterpene biosynthesis to sesquiterpene synthases that belong to the TPS-a clade and have no plastid-targeting transit sequences. In *S. habrochaites* accession LA1777, a member of the TPS-e/f clade designated SBS is known to be localized in the plastid and to catalyze the formation of the sesquiterpenes santalene and bergamotene synthase from (Z,Z)-FPP (Sallaud et al. 2009), and although our RNA-seq database contains sequences that are highly similar to SBS, we did not characterize them further.

In addition to divergence in the enzymatic properties of *S. lycopersicum* and *S. habrochaites* alleles of respective TPS genes described above, the two species also differ in the set of sesquiterpene synthase genes that are expressed in their trichomes. For example, TPS14 is expressed in *S. habrochaites* trichomes but not in the *S. lycopersicum* trichomes, whereas TPS31 is expressed in the latter but not in the former, although in both cases the expression of these genes was relatively low. Thus, we can conclude that the differences in the repertoire of sesquiterpenes produced in these two *Solanum* species is due both to evolution of enzymatic activity and regulation of gene expression. Finally, we were also able to show that the major product of SIPTS17 and ShTPS17 is valencene (Fig. 3). This compound is detected in *S. habrochaites* trichomes, but not
in the S. lycopersicum trichomes (Table 2). The major product of SITPS31, viridiflorene (Fig. 6), we could not detect in trichomes of either species (Table 2).

In conclusion, our RNAseq approach has allowed us to identify sesquiterpene synthases that are expressed in stem trichomes of cultivated and wild tomato species. Some of these genes are also expressed elsewhere in the plant, but others appear to be specifically expressed or highly enriched in stem trichomes. The analysis of the enzymatic activity of orthologs revealed that in some case different products are made by orthologous sesquiterpene synthases and that some orthologs differ in their substrate preference. These results provide new tools to study the evolution of terpene synthase activity at the level of protein structure.

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