**Terpene Synthase-b and Terpene Synthase-e/f Genes Produce Monoterpenes for *Phalaenopsis bellina* Floral Scent**

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Orchids are the most species-rich plants and highly interactive with pollinators via visual or olfactory cues. Biosynthesis and emission of volatile organic compounds (VOCs) to the atmosphere facilitate the olfactory cues and ensure successful pollination. *Phalaenopsis bellina* is a scented orchid with monoterpenes as major VOCs, comprising linalool, geraniol, and their derivatives. Comparative transcriptomics analysis identified four terpene synthase-b (TPS-b) genes and two TPS-e/f genes with differential gene expression between scented and scentless *Phalaenopsis* species. Here, we confirmed their differential expression between scented and scentless *Phalaenopsis* orchids and excluded one TPS-b candidate. We analyzed the temporal and spatial expression and functionally characterized these TPSs. Both TPS-b and TPS-e/f genes showed an increased expression on blooming day or 3 days post-anthesis (D+3) before the optimal emission of floral scent on D+5, with especially high expression of PbTPS5 and PbTPS10. The TPS-b genes are expressed exclusively in reproductive organs, whereas the TPS-e/f genes are expressed in both reproductive and vegetative organs. In planta functional characterization of both PbTPS5 and PbTPS10 in tobacco and scentless *Phalaenopsis* plants did not produce terpenoids. Further ectopic expression in scented *Phalaenopsis* cultivar *P. I-Hsin Venus* showed that linalool was the main product, with PbTPS10 displaying 3-fold higher activity than PbTPS5. On in vitro enzyme assay with purified recombinant TPS-b proteins ectopically expressed in *Escherichia coli*, geraniol was the product catalyzed by PbTPS5 and PbTPS9. PbTPS3 was a linalool/β-cis-ocimene synthase and PbTPS4 a linalool synthase. In conclusion, both TPS-b and TPS-e/f enzymes orchestrated floral monoterpane biosynthesis in *P. bellina*.

**Keywords:** *Phalaenopsis*, monoterpane, floral scent, terpene synthase, geraniol, linalool, orchid, recombinant protein

**INTRODUCTION**

Orchidaceae is one of the most species-rich families in angiosperms and highly interactive with their pollinators. Interaction between pollinators and orchids has played a major role in orchid evolution (Schiestl et al., 2000; Sanguinetti et al., 2012). In Orchidaceae, both visual and olfactory captures are crucial for successful pollination. For olfactory capture, orchids emit floral volatile organic
compounds (VOCs), including terpenoids, phenylpropanoids, and benzoids, into the atmosphere. Floral VOCs have been characterized in several orchids, including monoterpenes of linalool and geraniol in *Phalaenopsis bellina* (Hsiao et al., 2006). Emission of the floral scent is developmentally regulated: It is undetectable on blooming day or with flower buds, starts to be detected on 3 days post-anthesis (D + 3), and is optimized on D + 5. It decreases as the flower becomes senescent (Hsiao et al., 2006).

Terpenes represent the largest group of plant secondary metabolites. Terpenoids have many diverse structures, with various carbon skeletons and a large assortment of functional groups. The biosynthesis of a terpene starts from the formation of the basic C5 units, isopentenyl diphosphate (IPP), and its allylic isomer, dimethylallyl diphosphate (DMAPP). IPP is synthesized via two independent pathways: the mevalonate (MVA) pathway in the cytosol and methylerythritol phosphate (MEP) pathway in the plastids. The action of various short-chain prenyltransferases produces direct precursors for terpene synthases (TPSs); monoterpane synthase (MTPS) uses geranyl diphosphate (GDP) as a substrate for producing monoterpenes (C10), sesquiterpene synthase (STPS) uses farnesyl diphosphate (FDP) as a substrate for the biosynthesis of sesquiterpenes (C15), and diterpene synthase (DTPS) uses geranylglycerol diphosphate (GGDP) as a substrate for producing diterpenes (C20) (Spitzer-Rimon et al., 2010; Colquhoun et al., 2011). Both MTPS and DTPS in the MEP pathway are present in the plastids for monoterpene and diterpene biosynthesis, whereas STPS in the MVA pathway is present in the cytoplasm for sesquiterpene biosynthesis (Lichtenthaler et al., 1997).

**TPS** genes are a large family classified into seven major subfamilies, namely, **TPS-a** to **TPS-h** (Chen et al., 2011). The **TPS-c** subfamily is the original subfamily and contains copalyl diphosphate synthases (CPSSs), CPS/kaurene synthases (CPS/KSs), and DTPSSs. **TPS-a** and **TPS-b** subfamilies are the angiosperm-specific subfamilies, and they contain STPSs and MTPSSs, respectively (Bohmann et al., 1998; Chen et al., 2011). The **TPS-d** subfamily is for gymnosperms MTPS and DTPS. **TPS-g** subfamily lacks the RRX1 domain motif for producing acyclic monoterpene and sesquiterpene. The **TPS-h** subfamily is identified as DTPS in *Selaginella moellendorfii* (Chen et al., 2011). The **TPS-e/f** subfamily contains KSs, MTPSSs, STPSs, and DTPSSs. Diverse functions have been reported for **TPS-e/f** enzymes, including linalool synthase in *Clarkia breweri* (Dudareva et al., 1996), geranylinalool synthase in *Arabidopsis thaliana*, and farnesene synthase in *kiwifruit* (Nieuwenhuizen et al., 2009). The **TPS-f** subfamily is probably dicot-specific, whereas the **TPS-e** subfamily is not (Chen et al., 2011).

*Phalaenopsis bellina* is a native *Phalaenopsis* species in Borneo, Sarawak, and a common breeding parent for scented cultivars with a monoterpene phenotype. It has linalool, geraniol, and their derivatives as the major floral VOCs (Hsiao et al., 2006). The combination of bioinformatics and genomics identified PbGDPS as the key enzyme in orchid floral scent biosynthesis based on its high enrichment in the scented *P. bellina* but not scentless *Phalaenopsis equestris* and *Phalaenopsis aphrodite* subsp. *formosana* (hereafter *Phalaenopsis aphrodite*) (Hsiao et al., 2008; Chuang et al., 2018a). PbGDPS exhibits a dual prenyltransferase activity, producing GDP and FDP as the products, but lacks the DD(X)−4-D domain (Hsiao et al., 2008). The flower-specific expression profile of PbGDPS is concomitant with the emission pattern of monoterpenes on D + 5 (Hsiao et al., 2006). PbGDPS is the first enzyme with substantially differential expression between *P. bellina* and *Phalaenopsis aphrodite*, followed by several TPS genes in the monoterpene biosynthesis pathway (Chuang et al., 2018b). Comparative transcriptomics revealed the upregulated expression of **TPS-e/f** as well as **TPS-b** genes in *P. bellina* as compared with the scentless *Phalaenopsis aphrodite* (Supplementary Figure 1B).

Here, we confirm the differential gene expression of these **TPS** genes between *P. bellina* and *Phalaenopsis aphrodite*, illustrate the temporal and spatial gene expression, and provide a functional characterization of the **TPSs** in **TPS-b** and **TPS-e/f** subfamilies to further understand floral monoterpene biosynthesis in *Phalaenopsis* orchids. Functional characterization was revealed by both *in planta* and *in vitro* enzyme activity assays with transient overexpression and purified recombinant proteins ectopically overexpressed in *Escherichia coli*, respectively. These results suggest that both **TPS-b** and **TPS-e/f** enzymes contribute to monoterpene biosynthesis in the *P. bellina* floral scent.

**MATERIALS AND METHODS**

**Plant Materials**

*Phalaenopsis bellina* was from the Ming-Hui Orchids Nursery (Changhua, Taiwan) and Quan-Ya Orchid Nursery (Tainan, Taiwan) (Supplementary Figure 2A). *P. lueddemanniana* was selected because it has mild levels of floral linalool (hereafter Venus) was from I-Hsin Biotechnology Corp. (Chiayi, Taiwan) (Supplementary Figure 2B), and *Phalaenopsis amabilis* and *P. Sogo Yukidian “V3”* (hereafter V3) were from Taiwan Sugar Corp. (Tainan, Taiwan). All *Phalaenopsis* plants were grown in the greenhouse at the National Cheng Kung University under natural light and controlled temperature from 27 to 30°C with 80% humidity. Venus, a commercial scented hybrid (Supplementary Figure 2B), was selected because it has mild levels of floral linalool emission. The genetic background of Venus is as follows: *Phalaenopsis amboinensis* (25%), *Phalaenopsis equestris* (25%), *Phalaenopsis venosa* (18.7%), *Phalaenopsis violacea* (15.6%), *Phalaenopsis amabilis* (10.4%), *Phalaenopsis lueddemanniana* (3.1%), and *Phalaenopsis aphrodite* (2.1%) via ORCHIDEYA.CA^1.

Nicotiana tabacum seeds were obtained from Dr. Ching-Chun Chang’s laboratory (Department of Biotechnology and Bioindustry Sciences, NCKU) and germinated directly in soil and grown under a 16-h day/8-h night photoperiod with temperatures set at 27°C. Humidity was adjusted to 70%.

**Phylogenetic Analysis**

For TPS phylogenetic analysis, previously studied TPS sequences were downloaded from NCBI isolated from other plant species, and the full-length sequences were analyzed with the putative

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^1http://www.orchideya.ca/
TPSs isolated from *P. bellina* and *Phalaenopsis aphrodite* transcriptomes. Information on TPSs used for phylogenetic analysis is in Supplementary Tables 1, 2. The alignment was performed by using ClustalW of the Molecular Evolutionary Genetics Analysis (MEGA7) program. The phylogenetic tree was constructed by using the neighbor-joining method with the Jones–Taylor–Thornton model and pairwise deletion with 1,000 bootstrap replicates.

**Quantitative Real-Time PCR and Gene Expression Clustering**

Total RNA was extracted from seven stages of *P. bellina* flowers, starting from D − 1 to D + 17, as described in the study by Chuang et al. (2018b). After DNA contamination was removed by DNase I (New England Biolabs, Beverly, MA, United States), reverse transcription of DNA to cDNA involved using SuperScript III (Invitrogen, Carlsbad, CA, United States). Primers were designed by using Primer Express 3.0 (Applied Biosystems, Foster City, CA, United States). Quantitative Reverse transcription-PCR (RT-PCR) (qRT-PCR) involved using a StepOne Real Time PCR System and SYBR Green kit (Applied Biosystems, Foster City, CA, United States) as described in the study by Chuang et al. (2018b). Several reference genes widely used in *Phalaenopsis* orchids were identified by a local BLASTN search (Chen et al., 2005; Hsiao et al., 2008; Hsieh et al., 2013; Hsu et al., 2015). *Actin1* had equivalent levels in the RNA-seq analysis of the *P. bellina* transcriptome and microarray data for *Phalaenopsis aphrodite* (Chuang et al., 2018b), so *PbActin1* in *P. bellina* was selected as an internal calibrator for further analysis. The expression of all genes was normalized by the reference gene *PbActin1*. Calculation of mean and error bars was based on three replicates. The temporal gene expression experiments for both TPS-e/f and TPS-b were performed in different years with *P. bellina* from different orchid nurseries. *P. bellina* from various orchid nurseries have different scent emission profiles: Some last longer, and others are shorter but with the same volatile compounds. The floral scent emission for *P. bellina* lasted longer for TPS-b gene expression analysis, so we analyzed TPS-b gene expression from bud to D + 17, whereas those used for TPS-e/f gene expression analysis has a shorter scent emission profile, so it was analyzed from 5 days before anthesis (D − 5) to D + 10.

**Transient Expression in Nicotiana tabacum**

The modified pCAMBIA1304 vector containing *PcMYB2*, which contributes to the red color formation in *Phalaenopsis* spp. flowers (Hsu et al., 2015), was used as a binary vector for the construction of transient expression in *N. tabacum* and *Phalaenopsis* orchids. TPS genes were constructed into the modified binary vector, and then the recombinant vectors containing inserted TPS genes (dubbed pCAMBIA1304-TPS) were transformed into *Agrobacterium tumefaciens* strain GV3101 by electroporation. The transformed *Agrobacteria* were spread on LB agar plates containing rifampicin (10 µg/ml), gentamycin (25 µg/ml), and kanamycin (50 µg/ml). The plates were incubated for 2 days at 28°C, and colonies were tested for insertion by colony PCR with ProTaq DNA Polymerase (Protech Technology Enterprise, Taiwan). A transformed single colony was used to inoculate a 5-ml culture of LB medium plus appropriate antibiotics, and cells were grown for 24 h at 28°C and 220 rpm. This pre-culture was used to inoculate a 100-ml culture and grown for about 4 h at 28°C to OD600 = 0.8–1.0. *Agrobacteria* were harvested by centrifugation at 5,000 × g for 4°C for 15 min, then resuspended in infiltration medium (Murashige and Skoog medium, 1 mM acetosyringone) to OD600 = 0.4, and incubated for 2–4 h at room temperature. Diluted *Agrobacteria* was injected into the abaxial side of the three youngest *N. tabacum* leaves that were > 1 cm by using a 1-ml syringe. Transformed plants were maintained in a growth chamber at 28°C under long-day conditions (16 h/8 h for light/dark), and gas chromatography–mass spectrometry (GC-MS) analysis was performed at 5–7 days post-infiltration.

**Transient Overexpression in Phalaenopsis Orchids**

The expression plasmid, pCAMBIA1304-TPS as described above, was transformed into *Agrobacterium tumefaciens* strain EHA105 by electroporation. The growth of agrobacteria was as described above. The transformed *Agrobacteria* was injected into the adaxial side of sepal and petals at different floral stages (flower bud or just bloomed flower) by using a 0.5-ml insulin syringe. Transformed plants were maintained at 25°C, and GC-MS analysis was performed at 5 dpi. Because the constructs containing *PcMYB2* gene activated the anthocyanin pathway, red color formation in transformed flowers was considered a positive transformation in *Phalaenopsis* orchids.

**Expression and Purification of Recombinant Proteins in Escherichia coli**

For ectopic overexpression in *E. coli*, the open reading frames of PbTPSs were amplified and cloned into the expression vector pET-28b (Novagen, Madison, WI, United States), which has a His-tag on both the N- and C-termini of the inserted fragment. The construct was introduced into *E. coli* C41 (DE3) pLysS strain (Invitrogen, Carlsbad, CA, United States). Sequence analysis and protein structure modeling confirmed that no mutant occurred near the protein active site on DNA amplification. A single colony of the constructs was incubated in 5 ml LB medium with 50 µl/ml kanamycin overnight at 37°C. Then, the culture was transferred to 750 ml LB medium with 50 µl/ml kanamycin, incubated at 37°C to OD600 = 0.4–0.6, and induced with 0.4 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) and grown under 16°C for 22 h. Cells were pelleted by centrifugation and suspended in extraction buffer (0.3 M NaCl, 50 mM HEPES, pH 7.4), then disrupted by sonication (SONICS, VC750, Newtown, CT, United States), Cell extracts were filtered through a 0.45-µm filter. Recombinant proteins were purified by using HisTrap TALON crude (GE Healthcare, Abingdon, United Kingdom) with different concentrations of imidazole solution (0.3 M NaCl, 50 mM HEPES, 5 mM/150 mM imidazole, pH 7.4). Protein expression was verified on Coomassie Blue-stained SDS-PAGE gel and Western blot analysis with 1/10,000 diluted.
mouse anti-histidine monoclonal antibody (Roche Diagnostics, Penzberg, Germany).

**Enzyme Activity Assay of Purified Recombinant Proteins**
The affinity-purified recombinant proteins were concentrated and incubated with assay buffer (HEPES, pH 7.4, 0.1 M KCl, 10 mM MgCl$_2$, 10 mM DTT, 10% glycerol) by using a 30 MWCO PES Vivassin 50-ml concentrator (Sartorius Stedom Biotech, Goettingen, Germany). For functional characterization, recombinant protein was added to a single vial and diluted with assay buffer to a total volume of 500 µl, then incubated with GDP (2.5 µM) for 1 h at 30°C.

**Volatile Collection and GC-MS Analysis**
The floral volatiles of the Phalaenopsis orchids were collected by using a scent-extracting apparatus (Chuang et al., 2017). The scent emission pattern of Phalaenopsis orchids was measured from a single flower, with three biological replicates. The volatiles collected were eluted by hexane and identified by gas chromatography/high-resolution mass spectrometry (GC/HRMS) at the NCKU Instrument Center, as described in the study by Hsiao et al. (2006). As a negative control, metabolites originating from the scent-extracting apparatus were analyzed for the background. The volatiles were collected by using a dynamic headspace sampling system and solid-phase microextraction (Aharoni et al., 2003) in a glass container with 200 mg Tenax-TA resin (60/80 mesh; Supelco, Bellefont, PA, United States) packed into a plastic tube. The headspace sampling apparatus inserted a silicone tube to connect to the solid phase extraction columns and another silicone tube connected the column with air pumps at 350 mmHg collecting. The collected compounds were analyzed by GC-MS (QP2010, SHIMADZU, Shimadzu Co., Tokyo). An INNOWAX column (60 cm, 0.3 mm, 0.25 µm phase thickness) was used, and the oven was programmed from 40 to 230°C (held for 5 min) at 5°C/min increments. The pressure of the helium inlet was set at 75.2 kPa with 34.6 cm/s linear velocity (split flow 8.3 µl/min). The injector temperature was kept at 240°C with the injected volume set to 1 µl and the electron energy to 70 eV. Mass spectra and reconstructed chromatograms were obtained by automatic scanning of the samples in the mass range m/z 20–500 Da. Peaks on mass chromatograms with characteristic fragments were checked for homogeneity. The identities of all compounds were determined by comparing retention times and mass fragmentation patterns with the NIST98 database (US Environmental Protection Agency, 1998) and NIST05 database (SHIMADZU, Shimadzu Co., Tokyo). For quantitative analysis, 10 µg/ml ethyl myristate was used as an internal standard.

**Protoplast Isolation**
Protoplasts of floral buds were isolated from Phalaenopsis aphrodite as described in the study by Shen et al. (2017). Briefly, 10 g of 2-cm flower buds was sterilized in 70% alcohol for 1 min, followed by three washes in sterilized distilled water. The floral bud was cut into 1-mm$^3$ pieces and immersed in sterilized water for 1 h. Then, floral buds were transferred to 10 ml enzyme solution (1 ml of 0.2M MES, 0.4 g cellulose RS, 0.2 g macerozyme R-10, 6 ml of 1M sucrose, 100 µl of 2M KCl, 50 µl of 2M CaCl$_2$, 10% BSA) for incubation for 4 h on a shaker (30 rpm) in darkness at room temperature. After incubation, the enzyme–protoplast mixture was passed through steel screens of 100- and 50-µm pore sizes to remove undigested materials. The sample was centrifuged at 700 rpm for 5–10 min. The protoplasts on the surface were collected, resuspended in the wash buffer (cell and protoplast washing salt containing 0.6 M mannitol), and centrifuged at 700 rpm for 5 min; then the supernatant was discarded. After three washes, the pellet was resuspended in 0.5 ml of W5 solution (150 mM NaCl, 125 mM CaCl$_2$, 5 mM KCl, 2 mM MES, pH 5.7, 5 mM glucose). Then, the supernatant was spun, removed, and resuspended in 500 µl of MMg solution (4 mM MES, pH 5.7, 0.6M mannitol, 15 mM MgCl$_2$) before polyethylene glycol (PEG) 4000 treatment.

**Transient Expression of Green Fluorescent Protein Fusion Proteins**
For cellular localization experiments, green fluorescent protein (GFP) was constructed either upstream or downstream from the analyzed genes, such as GFP-PbTPS3 or PbTPS3-GFP, respectively. The Gateway Cloning System (Invitrogen, Carlsbad, CA, United States) was used for generating p2FGW7-fused protein transformation constructs consisting of the open reading frame of PbTPS3 or PbTPS4 cDNA under the control of the cauliflower mosaic virus (CaMV) 35S promoter. The plasmids were transformed into protoplasts of Phalaenopsis aphrodite by PEG 4000 treatment. An empty vector that expresses GFP under the control of the CaMV 35S promoter served as a negative control. GFP fluorescence was observed under a Carl Zeiss LSM780 confocal laser scanner microscope (CLSM-780, Zeiss, Germany) with GFP excited at 488 nm with a krypton/argon laser and chlorophyll autofluorescence excited at 514 nm with a HeNe-Laser. The emissions were collected through a 506- to 530-nm band pass filter (for GFP) and a 680- to 750-nm band pass filter (for chloroplasts). To strengthen the image visibility and resolution of the original images, all images were enhanced to 50% sharpness and 30% contrast by using the built-in chart format of PowerPoint.

**RESULTS**

**Phylogenetic Analysis of Upregulated TPSs in Phalaenopsis bellina During Floral Development**
Comparative transcriptomic analysis of scented P. bellina and scentless Phalaenopsis aphrodite revealed the upregulation of several TPS genes in P. bellina (Chuang et al., 2018b; Supplementary Figure 1B). Among them, PbTPS5-1, PbTPS5-2, PbTPS9-1, PbTPS9-2-1, PbTPS9-2-2, PbTPS10-1, PbTPS10-2, and PbTPS7 are in the TPS-b subfamily, and PbTPS4, PbTPS3-1, and PbTPS3-2 are in the TPS-e/f subfamily. Alternative splicing had occurred and produced different transcripts from the same gene, such as TPS5-2, which has an intron retention between exons 5 and 6 that caused a stop codon and early termination. A similar situation occurred for TPS10-1, in that an intron
Differential Gene Expression of PbTPSs Between Phalaenopsis bellina and Phalaenopsis aphrodite

To further confirm the differential expression of these six PbTPS genes between scented and scentless Phalaenopsis orchids, qRT-PCR was performed. We examined the expression of four TPS-b genes (PbTPS3, PbTPS7, PbTPS9, and PbTPS10) and two TPS-e/f genes (PbTPS3 and PbTPS4) at the flower bud stage and D + 5 (Figure 2). Previously, we found that the emission of monoterpenes in P. bellina peaked at the full-bloom stages (D + 4 and D + 5) but was absent in floral buds (Chuang et al., 2018b). All PbTPS genes in P. bellina showed higher expression at D + 5 than at the bud stage. Except for PbTPS7, the qRT-PCR results were consistent with transcriptomic analysis showing an enhanced expression in the scented orchids. Expression was higher for PbTPS3, PbTPS4, PbTPS5, PbTPS9, and PbTPS10 in P. bellina than in Phalaenopsis aphrodite in floral buds and at D + 5 (Figure 2). These results suggested that PbTPS3, PbTPS4, PbTPS5, PbTPS9, and PbTPS10 were highly associated with floral scent monoterpene biosynthesis; PbTPS7 was removed from the following analysis.

Temporal and Spatial Gene Expression of the PbTPS-b Subfamily

We examined the temporal expression of three TPS-b genes from the flower bud stage to D + 17 of P. bellina by qRT-PCR (Figure 3). Both PbTPS5 and PbTPS10 showed much higher expression than PbTPS9 in P. bellina during floral development (Figure 3A). PbTPS5 and PbTPS10 expressed highly on blooming day (Dd) and D + 3, respectively (Figure 3A). PbTPS9 showed a biphasic pattern, with expression high on D + 5, decreased on D + 7, and then high again on D + 13 (Figure 3A).

The three TPS-b genes were exclusively expressed in reproductive organs, with differential profiles. PbTPS5, PbTPS9, and PbTPS10 expressed in sepal, petal, and lip and to a lesser extent in column (Figure 3B). Especially, PbTPS5 expressed highly in petal, whereas PbTPS10 expressed highly in sepal.

Temporal and Spatial Expression Profiles of the PbTPS-e/f Subfamily

We analyzed the temporal expression of two PbTPS-e/f genes, PbTPS3 and PbTPS4, by qPCR from 5 days before anthesis (D − 5) to D + 10. PbTPS3 showed increasingly high expression from Dd to D + 5, then reduced expression on D + 7, whereas PbTPS4 showed high expression on D + 3 and D + 5, then diminished expression (Figure 3C). The expression was much lower for PbTPS4 than for PbTPS3 in flowers (Figure 3C). PbTPS3 expressed mainly in the leaf, and in the flower, it expressed higher in the lip than in the sepal, petal, or column (Figure 3D). PbTPS4 also showed high expression in the lip and column and very low expression in the sepal and petal; it was expressed in the root and leaf but at a lower level than in the floral organs (Figure 3D).

Transient Overexpression of PbTPS5 and PbTPS10 in Nicotiana tabacum

The temporal expression profiles of PbTPS5 and PbTPS10 were concomitant with the emission of monoterpenes such as linalool and geraniol, so we studied their enzyme functions by ectopic overexpression in N. tabacum. Several compounds were produced when overexpressing PbGDPS to provide GDP as the substrate. A monoterpene myrcene derivative, tetrahydromyrcenol, was produced when transiently overexpressing PbTPS10 or PbGDPS + PbTPS10 in N. tabacum (Supplementary Figure 5). However, neither linalool nor geraniol was produced in N. tabacum overexpressing PbTPS5 or PbTPS10. N. tabacum may have no other orchid-related genes required for monoterpene biosynthesis and hinder the production of linalool or geraniol.

Transient Overexpression of PbTPS5 and PbTPS10 in Scentless Phalaenopsis

We attempted the transient overexpression of PbTPS5 and PbTPS10 in the scentless species Phalaenopsis aphrodite and a scentless cultivar V3. Overexpression of PbTPS5 and PbTPS10 with or without PbGDPS in Phalaenopsis aphrodite did not induce the terpene production in the emitted volatiles or the internal non-volatiles (Supplementary Figures 6A,C), even though monoterpenes of α-terpineol and α-cedrene were detected in the non-volatile compounds in the overexpression of PbGDPS in Phalaenopsis aphrodite. This observation suggests that other enzymes or factors might be needed to produce volatile monoterpenes in Phalaenopsis aphrodite. Further overexpression of PbTPS5 and PbTPS10 was performed in V3, and the production of tetrahydromyrcenol and sesquiterpenoids was enhanced, but neither linalool nor geraniol was detected in the volatiles and non-volatiles of V3 (Supplementary Figures 6B,D). Previously, we showed that the scentless Phalaenopsis orchids lack PbZIP4 and PbbHLH4 transcription factors for floral volatile biosynthesis (Chuang et al., 2018a,b). Hence, some other factors (such as transcription factors) may be required and are missing in the scentless orchid flowers. Alternatively, the presence of repressors may hinder the monoterpene biosynthesis in the scentless Phalaenopsis orchids, even though the key enzymes are provided.
more than 10 generations of crossing. In contrast to *P. bellina* producing one or two flowers and with strong scent, Venus is floriferous (Supplementary Figure 2B) but with reduced floral scent. Before the overexpression, we analyzed the endogenous gene expression and floral scent emission of Venus. We first detected the scent compounds of Venus and examined the gene expression profiles of endogenous *PbGDPS*, *PbTPS5*, and *PbTPS10*. Venus produced VOCs of linalool, octanoic acid, and benzaldehyde in flowers, with linalool as the major floral scent, yet no geraniol was detected (Figure 4B). The floral scent emission followed a similar pattern as that of *P. bellina* (i.e., the strong scent was detected from D + 3). Concomitantly, *PbGDPS* and *PbTPS5* significantly expressed from Dd and D + 3, respectively (Figure 4A), yet with little or no expression of *PbTPS10* during floral development from flower bud to D + 13 (Figure 4A). This result implied that *PbTPS5* alone can account for linalool biosynthesis.

Cauliflower mosaic virus 35S promoter was used to drive the overexpression of both *PbTPS5* and *PbTPS10*. The emission of linalool was enhanced with transient overexpression of *PbTPS5* and *PbTPS10* in Venus floral tepals (peak 2, Figure 4C). Thus, both *PbTPS5* and *PbTPS10* caused a 7.7- and 29.0-fold increase, respectively, of linalool production as compared with GUS. In addition, emission of tetrahydromyrcenol was detected overexpression of *PbTPS5* and *PbTPS10*, respectively (peak 1, Figure 4C). However, geraniol was not produced on overexpressing *PbTPS5* or *PbTPS10* in Venus.

Functional Characterization With Purified *TPS-b* Recombinant Proteins Ectopically Overexpressed in *Escherichia coli*

To further confirm the enzyme activity, *in vitro* assay was performed with purified recombinant proteins ectopically expressed in *E. coli*. *PbTPS5* was cloned into PET-28b with 6x His at the N- and C-terminus and transformed into *E. coli* C41 (DE3). The affinity-purified *PbTPS5* protein was assayed in a buffer containing substrate GDP, and the products were analyzed by GC-MS. Geraniol was detected as the main product catalyzed by *PbTPS5* (Figure 5A).

The enzyme activity of *PbTPS9* was examined with the purified recombinant proteins as described above. The product of *PbTPS9* was geraniol (Figure 5B).
Functional Characterization of Purified TPS-e/f Recombinant Proteins Ectopically Overexpressed in Escherichia coli

To functionally characterize the two TPS-e/f genes PbTPS3 and PbTPS4, we analyzed the affinity-purified recombinant proteins by using three different prenyl diphosphates as substrates, namely, GDP (C10), FDP (C15), and GGDP (C20). The reaction products were analyzed by GC-MS. PbTPS3 was a dual enzyme as a linalool/($\beta$)-cis-ocimene synthase (Figure 6A). The authentic standard of linalool was used for comparing retention time. PbTPS4 was a linalool synthase and used GDP as the substrate (Figure 6B). Neither FDP nor GGDP could be catalyzed by PbTPS3 and PbTPS4 to produce sesquiterpene or diterpene (Supplementary Figure 7). Overall, these data indicate that PbTPS3 and PbTPS4 are the MTPS that exclusively produced linalool and ($\beta$)-cis-ocimene, or linalool only, respectively.
Subcellular Localization of PbTPS3 and PbTPS4

Monoterpane synthases are mainly present in the plastids. However, both PbTPS3 and PbTPS4 lack typical transit peptides for chloroplast localization (Supplementary Figure 8). We then performed the prediction of the subcellular localizations according to their amino acid sequences with various software packages. Unexpectedly, differential prediction results were obtained. For example, ChloroP 1.1 predicted that both PbTPS3 and PbTPS4 were not localized in the chloroplasts. TargetP suggested that they were localized in other sites and mitochondria, respectively. pSORT II indicated the possible localization of PbTPS3 in the cytosol (69.6%) and nucleus (26.1%) and possible localization of PbTPS4 in the nucleus (34.8%) and cytosol and mitochondria (21.7%). Wolf PSORT indicated that PbTPS3 and PbTPS4 possibly localized in the nucleus and chloroplast, respectively.

Therefore, subcellular localization of PbTPS3 and PbTPS4 was determined with GFP added at the N- or C-terminus of the full-length cDNAs (Figure 7). These constructs were transformed into protoplasts isolated from mature flower buds of Phalaenopsis aphrodite, and the GFP signal and autofluorescence emitted from chloroplasts were examined by confocal laser scanning microscopy. The GFP fluorescence of both PbTPS3 and PbTPS4 fusion proteins was exclusively overlaid with the autofluorescence signal of chloroplasts, which suggests that they are localized exclusively in the chloroplast (Figure 7). In contrast, the controls of no DNA added or empty vector showed no or very low GFP signal, respectively (Figure 7). These results confirmed that both PbTPS3 and PbTPS4 were localized in the plastids even without a typical transit peptide signal.

DISCUSSION

Both TPS-b and TPS-e/f Subfamilies Are Involved in the Biosynthesis of Monoterpenes in Phalaenopsis bellina Flower

In addition to the three TPS genes in the TPS-b subfamily with an enhanced expression in P. bellina as compared with Phalaenopsis aphrodite, two TPS genes in the TPS-e/f subfamily were also enriched. In P. bellina, differential gene expression of PbTPS genes is involved in floral monoterpene biosynthesis in various
parts of a single flower. For example, PbTPS5, PbTPS9, and PbTPS10 were involved in monoterpene biosynthesis in tepal (sepal and petal) and PbTPS3, PbTPS4, PbTPS5, PbTPS9, and PbTPS10 in lip (Figure 8A). In addition, multiple TPS genes are involved in the same enzymatic function, and one TPS gene may catalyze more than one product of monoterpenes. For example, PbTPS3, PbTPS4, PbTPS5, and PbTPS10 produce linalool, and PbTPS5 and PbTPS9 produce geraniol. In addition, PbTPS3 produces (β)-cis-ocimene, with PbTPS3 expressed in both flower and leaf. PbTPS3 has a dual enzyme activity of linalool synthase/(β)-cis-ocimene synthase (Figure 8B).

Both PbTPS3 and PbTPS4 contain a DDXXD and NSE/DTE motif for GDP binding, and they lack the RRX8W motif for cyclic monoterpene synthesis (Supplementary Figure 8). PbTPS3 is the dual function enzyme for producing acyclic monoterpenes, including linalool and (β)-cis-ocimene. It was expressed in flower and to a much higher extent in leaf (Figure 3D). However, GC-MS analysis of leaf VOCs revealed only 1% (β)-cis-ocimene but no linalool (Supplementary Figure 9). The heterologous expression of geraniol synthase from Ocimum basilicum in E. coli and plant systems revealed that heterologous expression affects the amount of geraniol production in the leaf tissue or E. coli.
FIGURE 5 | Functional characterization of purified recombinant protein of (A) PbTPS5 + GDP and (B) PbTP9 + GDP. GC-MS analysis of authentic linalool standard incubated with enzyme assay buffer. As negative controls, reactions were performed with only GDP added without any recombinant proteins.

(Fischer et al., 2013). Therefore, the enzymatic functions of TPS genes depend on the enzyme amino acid sequence or substrate and also the conditions of an enzymatic reaction, especially when comparing plant and bacterial systems (Brillada et al., 2013).

**Functional Characterization of PbTPS5 and PbTPS10**

PbTPS5 showed the enzyme activity of linalool synthase when overexpressed in Venus, and Venus itself produced only linalool and expressed PbTPS5 slightly (Figure 3). Negative regulators of PbTPS10 may be present in the Venus flowers and prevent its gene expression and production of linalool. Previously, differences in biochemical conditions between plant and microbial expression systems produce different products (Kollner et al., 2004). These differences between heterologous systems could result from the protein conformational change in the binding pocket due to different environments (e.g., pH, ion concentrations) or different posttranslational modifications (Fischer et al., 2013). For monoterpane synthesis, GDP is the precursor, and the conversion of GDP into monoterpenols involves the formation of a carbocation intermediate (Ilc et al., 2016). Any subtle differences in protein conformation altered by the environment (chemicals or other proteins) could affect the profile of carbocation intermediate, thus producing different products (Fischer et al., 2013). In addition, the presence of other enzymes that convert one monoterpenol into another monoterpenol cannot be ruled out because of the many enzymes in plants. The other factor that can be taken account in affecting the production of monoterpenes is intracellular pH. Changes in intracellular pH can cause the chemical transformation of monoterpenols. Previous study found that monoterpenols tended to interconvert under acidic condition but are stable at neutral pH (Fischer et al., 2011). Previously, we detected the pH value of various colored Phalaenopsis orchids and found that it increases from red-purple, white, purple-violet to violet blue color from pH 4.77 to 5.54 (Liang et al., 2020). Although the white flowers of Phalaenopsis aphrodite and V3 have similar pH values (5.05
and 5.13, respectively), different compounds were produced by overexpressing TPS genes in these two orchids, which suggests that the pH value was not the major effect on the content of volatiles in *Phalaenopsis* orchids (Supplementary Figure 5).

However, *in vitro* enzyme activity assay revealed that geraniol was the main product catalyzed by purified recombinant *PbTPS5*. With overexpression of *PbTPS5* and *PbTPS10* in Venus, linalool was the only product for both enzymes, with *PbTPS10* displaying high ability for linalool production. In addition, molecular modeling and docking were performed by using SwissDock program with both *PbTPS5* and *PbTPS10* as targets and GDP as a ligand. The ligand was difficult to dock into the active site of *PbTPS5* as compared with *PbTPS10*, which implies that the binding pocket was easier to access with *PbTPS10* than with *PbTPS5*. This finding explains why the activity of *PbTPS10* was better than that of *PbTPS5* in monoterpene synthesis. Within the flower of *P. bellina*, *PbTPS10* mainly expressed in the sepal and slightly in the petal, whereas *PbTPS5* is mainly expressed in the petal but less in the sepal. When combining the gene expression and enzyme activity data, the monoterpene biosynthesis in different floral organs of *P. bellina* could be driven by different genes and for different scent compounds. Recent reports showed that many TPS genes can produce more than one terpenoid, which could be related to the size of the active cavity. Thus, the amino acid sequences of TPS genes are among the factors that affect the TPS functional property (Pazouki and Niinemets, 2016). Also, previous TPS studies indicated that the outcome of *in planta* experiments could be affected by the plants chosen; for instance, when geraniol synthase from *O. basilicum* was expressed in grapevine, citronellol and nerol were produced (Arendt et al., 2016). The products of TPS genes may vary in different host plants because the function of MTPS is based on not only the amino acid sequence but also the cellular background (Fischer et al., 2013).

**Differential Positive and Negative Gene Regulation of *PbTPS5* and *PbTPS10***

Even though the coding sequences of *PbTPS5* and *PbTPS10* are similar, they showed distinct temporal and spatial expression profiles. For example, the temporal expression of *PbTPS5* started on the blooming date, whereas that of *PbTPS10* started on D + 3. Spatially, *PbTPS5* expressed highly in the petal, and *PbTPS10* expressed highly in the sepal, and both expressed similarly in the lip. In addition, these two genes revealed distinct profiles of transcriptional regulation (Chuang et al., 2018b). Both *PbbZIP4* and *PbNAC1* positively regulated the promoter of *PbTPS5* but did not affect the promoter of *PbTPS10* (Chuang et al., 2018b). Conversely, both *PbbHLH4* and *PbERF9* highly positively regulated the promoter of *PbTPS10* but did not affect that of *PbTPS5*. In contrast, *PbbHLH2* and *PbbZIP26* negatively regulated *PbTPS5*, and *PbbHLH5*, *PbbZIP26*, and *PbbMYB22* negatively regulated *PbTPS10* (Chuang et al., 2018b). Even though both *PbTPS5* and *PbTPS10* were negatively regulated by *PbbZIP26*, only *PbTPS10* expression was inhibited in Venus, which suggests that both bHLH5 and MYB22 may function as negative regulators to prevent the expression of *PbTPS10*. 
FIGURE 7 | Subcellular localization of PbTPS3 and PbTPS4. Protoplasts of Phalaenopsis aphrodite tepals were transformed with GFP fusion proteins (GFP-PbTPS3, PbTPS3-GFP, GFP-PbTPS4, and PbTPS4-GFP). Images were taken by confocal laser scanning microscopy. Bright field, the image in bright field; Chloroplast with autofluorescence image false-colored in red; GFP, GFP fluorescence image false-colored in green; Merge, merge of bright field, chloroplast and GFP fluorescence images.
MYB transcription factor has a negative regulation role in floral scent biosynthesis (Reddy et al., 2017). Further assessments will be needed to elucidate the negative regulation of floral scent biosynthesis.

Functional Characterization of PbTPS3 and PbTPS4
The purified recombinant PbTPS3 had dual enzyme functions as a linalool/(β)-cis-ocimene synthase, whereas that of the purified recombinant PbTPS4 produced only linalool. Both PbTPS3 and PbTPS4 were expressed in reproductive organs, including sepal, petal, and lip. In addition, PbTPS3 was expressed in the column at a similar level as in the sepal, whereas PbTPS4 was expressed much lower in the column. So, the major floral scent of P. bellina in the flower tepal (sepal and petal) is dominated by linalool and geraniol, whereas the lip has the floral scent of (β)-cis-ocimene in addition to linalool, and the column has the floral scent of (β)-cis-ocimene. Emission of (β)-cis-ocimene is 15.8 ng/flower/h, accounting for 2% of the floral scent of P. bellina (Chuang et al., 2018b).

PbTPS3 and PbTPS4 Are New Members of the TPS-e/f Subfamily
TPS enzymes in the TPS-e/f subfamily have dual functions or play different roles. For example, the (β)-cis-ocimene/farnesene synthase of Actinidia delicosa has dual functions, and it produces the (β)-cis-ocimene and farnesene mixture for pollinator attraction (Nieuwenhuizen et al., 2009). The foregoing observations indicate that the TPS genes in the TPS-e/f subfamily have various ecological roles in plants, including floral scent biosynthesis and plant defense. For example, linalool synthase is involved in linalool biosynthesis in Clarkia concinna and C. breweri. The expression of linalool synthase in the stigma of C. breweri leads to the emission of linalool and linalool oxides to serve as pollinator attractants (Dudareva et al., 1996). In addition, the geranyllinalool synthase of A. thaliana can produce the diterpene product of geranyllinalool for plant defense (Herde et al., 2008). Thus, the (β)-cis-ocimene/linalool synthase PbTPS3 and linalool synthase PbTPS4 in P. bellina are new members in the TPS-e/f subfamily.

Both linalool and (β)-cis-ocimene have a defense role against insect herbivores in many plants such as cotton, soybean, and rice (Pare and Tumlinson, 1999). The promoter of PbTPS3 contains three and 10 binding domains of the plant defense transcription factors WBOXATNPR1 and WRKY710S, respectively, and the promoter of PbTPS4 contains three binding domains for WRKY710S (Supplementary Figure 10). Thus, both PbTPS3 and PbTPS4 may play a role in defense in P. bellina. PbTPS3 may play different roles in different organs of P. bellina. In flower, PbTPS3 plays a role in linalool synthesis to attract pollinators but plays a defense role in the leaf.

The Evolution and Diversity of the TPS-e/f Subfamily
Linalool synthases have been identified in two TPS subfamilies, namely, TPS-b and TPS-e/f, so linalool synthase functionality has arisen multiple times independently during evolution (Nieuwenhuizen et al., 2009). The TPS-e/f subfamily is unique in that the members contain a conserved conifer diterpene internal sequence (CDIS) domain. Thus, the CDIS domain exists in a common DTPS ancestor but was lost during the evolution of MTPS and STPS (Brodmann et al., 2008). The TPS-f subfamily is probably dicot-specific, whereas the TPS-e subfamily is not (Chen et al., 2011). The TPS-e/f subfamily has the DDXXD conserved motif and lacks the DXDD conserved motif. From multiple alignment of amino acid sequences of the TPS-e/f subfamily in P. equestris, P. bellina, and other plants, PbTPS3 and PbTPS4 are cladded in the TPS-e/f subfamily (Figure 1; Supplementary Figures 4, 8). Our study indicates that PbTPS3 and PbTPS4 from orchids belong to the TPS-e/f subfamily and enlarge the content of this subfamily.
The subcellular localization analysis indicated that both PbTPS3 and PbTPS4 GFP fusion proteins are located in the plastids. The temporal expression patterns of PbTPS3 and PbTPS4 were concomitant with the volatile emission pattern of *P. bellina*. Taken together, PbTPS3 and PbTPS4 play different ecological roles of \((\beta)-\text{cis}-\text{ocimene/linalool and linalool biosynthesis in } P. \text{ bellina.}

### DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

### AUTHOR CONTRIBUTIONS

HH, Y-WK, and Y-CC performed the experiments of quantitative real-time RT-PCR, did functional characterization, and analyzed the data. Y-PY performed the original screening. L-MH performed the phylogenetic analysis. M-FJ assisted the sequence analysis and molecular docking. W-HC provided the suggestions for plant materials. H-HC conceived research plans and composed the article with assistances of all the authors, completed the writing, and served as the corresponding author and approved the submitted version.

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### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2021.700958/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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