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Discovering Fragrance Biosynthesis Genes from *Vanda Mimi* Palmer Using the Expressed Sequence Tag (EST) Approach

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Additional information is available at the end of the chapter

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1. Introduction

1.1. Orchidaceae

Orchidaceae is the largest family of angiosperms with an estimation of 17000 to 35000 species in 880 genera (Chai & Yu, 2007). In Malaysia, more than 230 orchid genera and 4000 species had been discovered (Go *et al.*, 2012). In Peninsular Malaysia, a total of 898 species in 143 genera are currently recognised (Go *et al.*, 2010). The amazing vast diversity of types and forms enable the Orchidaceae to be successfully distributed and colonised almost every habitats worldwide (Arditti, 1992). As a result of selective forces from evolution, orchids are found to be evolved from its ancestral forms and adapted well to their present habitats (Aceto & Gaudio, 2011). Associated with their diverse floral morphology and physiology properties, they have drawn the attention of botanists and scientists for centuries. There are orchids which resemble moths (*Phalaenopsis*), butterflies (*Oncidium papillo*), the slippers of Aphrodite or moccasins (*Paphiopedilum* or *Cypripedium*), dancing ladies (*Oncidium*), spiders (*Brassia*), scorpions (*Arachnis*) and bees (*Ophrys*) (Teoh, 1980).

Similar to other angiosperms, two whorls of perianth segment can also be found in orchids. The outer and inner whorls of the orchid flowers consist of three petals and three sepals. The labellum or lip (one of the petals), is distinctly evolved from the other two morphologically and physically (Arditti, 1992). The lifespan of opened-orchid flowers can range from as short as one day to as long as 270 days (Micheneau *et al.*, 2008).

Most orchids are epiphytic that obtain their support from trees but not for nutrition while the rest are terrestrial plants (Rada & Jaimez, 1992). Orchids' cultivation was famous since 5000 years ago in China where *Cymbidium* was grown as potted plants, and *Vanda* and

Aerides were suspended from baskets (Teoh, 1980). Orchid hybrids can be divided into three groups which are hybrid species (produced by selfing a species or by crossing two plants belonging to the same species), inter-specific hybrid (produced from the crossing between two different species belonging to the same genus or by second crossing with other inter-specific hybrids) and inter-generic hybrid (produced from the crossings of orchids belonging to different genera).

Orchid hybrids cultivation started since 1856 by John Dominy (<http://www/ionopsis.com/hybridization.htm>). *Calanthe dominii* was created from the crossing between *Calanthe masuca* and *Calanthe furcated*. Orchid hybrids made up majority of the commercial orchids in Malaysia and Singapore. There are lots of reasons that led to the wide area coverage of orchid hybrids cultivation, such as the ease of cultivation, free blooming habit, and compactness and fantastic arrays of shapes, colours and flowers (Kishor *et al.*, 2006).

The export of orchids from Malaysia, Thailand and Singapore contributed RM200 million annually in the world floral (Ooi, 2005). According to the Japan Florists' Telegraph Delivery Association, cut flower in Japan constituted around 13.3% of the imported market in the year 2010 with Malaysia having the largest market share of imports, which accounted for 23.4% (7,648 million yen), followed by Columbia at 19.2% and China at 10.4%. Malaysian orchids consisted of 8.1% of the imported cut flower orchid to Japan.

1.2. Vandaceous orchids

Tropical Asia is the native home for approximately 50 vandaceous orchids species. They are distributed in Sri Lanka and southern India to New Guinea, northern Australia and Solomon Islands, and north to China, Taiwan and the Philippines. Thailand was found to be predominated with 11 vanda species. In Thailand, vanda is a vital commercial orchid. Most of them exhibit monopodial growth where their leaves are varies according to habitat. Vandas have many different colours, and majority are yellowish-brown with dark brown spots.

The vanda has been designated as the 'Queen Orchid of the East' due to its robust and large rounded flowers (Teo, 1981). Most of the vandaceous orchids prefer sunlight but some are well adapted to shady areas. Like any other tropical orchids, they require warm temperature with good aeration. The vandaceous orchids take around three and half to 10 years to become mature flowering plants (Kishor *et al.*, 2008). Once matured, this orchid genus blooms every two to three months with the flowers lasting two to three weeks. As a result of land development, 28 orchid species have been listed as endangered species on Appendix II of the Convention on International Trade in Endangered Species (CITES) and prohibited from worldwide export. Among those orchids, two belong to vandas (*Vanda coerulea* and *Vanda sanderiana*).

It is impossible to differentiate or identify an orchid species based on the vegetative parts of the plant alone. Hence, a convenient and flower-independent method to allow quick assessment of a given orchid vegetative specimen for species confirmation can be achieved with the help of molecular markers. To date, approximately 50 species are registered under vandas in the Royal Horticultural Society (RHS) database due to their commercial importance.

1.3. *Vanda Mimi Palmer*

Vanda Mimi Palmer (VMP) (Figure 1) is one of those hybrids known to be highly sought after mainly for its fragrance rather than its flower shape or colour. It is a hybrid of *Vanda Tan Chay Yan* and *Vanda tessellate* (Roxb.) Hk.f. ex G. Don, and was registered on 1st December 1963 by the Gem Nursery, Singapore. It inherited its sweet fragrance and tri-colour flower from *V. tessellate*, while its terete-shaped is inherited from *V. Tan Chay Yan*. It has purplish green, but brown-speckled, petals and sepals with a dark blue lip. This hybrid is a frequent bloomer throughout the year with maximal floral scent emission when the flower is fully bloomed (an average bloom diameter of 5.0-7.0 cm). Fragrance emission is detected only when the flower bud starts to open as none is detected from the closed bud [average size of 0.8-1.4 cm](Chan *et al.*, 2011; Janna *et al.*, 2005). It has been recognised for its extraordinary fragrance as evident with awards won from the competitions organised by the Royal Horticultural Society of Thailand in 1993 and the 17th World Orchid Conference in 2002.



Figure 1. *Vanda Mimi Palmer*, a hybrid crossing between *Vanda Tan Chay Yan* and *Vanda tessellate* (Roxb.) Hk.f. ex G. Don.

Orchids cultivation entails hard work as the orchids can be easily infected by orchid-infecting viruses. More than 50 orchid-infecting viruses have been detected worldwide, with cymbidium mosaic virus (CymMV) and odontoglossum ringspot tobamovirus (ORSV) infections being the most prevalent (Sherpa *et al.*, 2007). Infected orchid cultivars usually exhibit blossoms with brown necrotic streaks and other necrosis symptoms. Infected flowers are smaller in size and poorer in quality. This has caused severe economic damages in the cut flower and potted plant industry (Ajjikuttira *et al.*, 2002; Sherpa *et al.*, 2007; Vejaratpimol *et al.*, 1999). CymMV infection is dominant and extremely stable in Orchidaceae, and it was found to be prevalent in VMP. A screen of our VMP cDNA library revealed a 1.6% contamination rate with CymMV genes (Teh *et al.*, 2011). Markers might be useful in facilitating the screening of virus-infected stock plants to minimise losses incurred in the floral industry. So far, the identification of CymMV infection is done through serological,

bioassay or electron microscopy. Those techniques include enzyme-linked immunosorbent assays (ELISA), dot-blot immunoassay (DBIA), rapid immunofilter paper assay (RIPA), immunosorbent electron microscope (ISEM), DIG-labelled cRNA probes, reverse transcription polymerase chain reaction (RT-PCR), quartz crystal microbalance-based DNA biosensors and TaqMan real-time quantitative RT-PCR (Eun *et al.*, 2002; Eun & Wong, 2000; Hsu *et al.*, 1992; Hu *et al.*, 1998; Khentry *et al.*, 2006; Rani *et al.*, 2010).

2. Importance of floral fragrance

Floral fragrance plays various functions in both the floral and vegetative organs. Fragrance is defined as a highly complex component of floral phenotype for its dynamic patterns of emission and chemical composition (Raguso, 2008). Due to their restriction to specific lineages and interactions in species-specific ecology, these have led to their designation as specialized or secondary metabolites (Pichersky *et al.*, 2006).

Floral fragrance has a significant impact in plant reproduction as it is a selective attractant in a variety of animal pollinators especially insects. Pollinators such as bees, butterflies and moths are able to discriminate visitation on plants based on the compositions of the floral scent. This plant-insect interaction has led to many successful pollinations and development of fruits in many crop species (Majetic *et al.*, 2009; Shuttleworth & Johnson, 2009).

Anti-microbial or anti-herbivore properties of floral volatiles could be used by plants to protect their vital floral reproductive parts from potential predators. Two types of plant defences can be characterised based on floral volatiles property, that are direct and indirect defences such as herbivore-induced volatiles signals, and visual and olfactory floral signals to attract pollinators (Schiestl, 2010). Indirect plant defences protect the plants by minimizing damage to plant tissues through attracting arthropods that prey or parasitize the herbivores. This general property has been reported in cabbage (Park *et al.*, 2005), *Lotus japonicas* (Arimura *et al.*, 2004) and cucumber (Agrawal *et al.*, 2002). On the other hand, direct defences take place when herbivore-induced volatiles repel or intoxicate the herbivores and pathogens. For example, some herbivore-induced monoterpenes and sesquiterpenes tend to react with various reactive oxygen species and thus protect the plants from internal oxidative damage (Dudareva *et al.*, 2004). Terp *et al.* (2006) reported that lipoxygenases are produced in *Brassica napus* seeds upon wounding and pathogen attack. Complexity in floral scent chemicals was found to be useful in protecting the plants' reproductive structures from herbivores and ants (Schiestl, 2010).

Through the discovery of pollinator-attracting floral scents as the source of olfactory pleasure since ancient times, humans had figured out unique values in certain types of floral scents and exploited them to cultivate and propagate specific plant species. For centuries, flowers with vibrant colours and scents have been used by people to enhance their beauty and this was seen in almost all major civilizations. Large numbers of aromatic plants have been used as flavourings, preservatives and herbal remedies (Pichersky *et al.*, 2006). Their economic importance also relies on their petals which are found to be the main site of natural fragrances and flavourings in most of the plants (Baudino *et al.*, 2007).

2.1. Fragrance biosynthesis pathways

Plants are known with the capability of synthesizing many volatile metabolites, either primary or secondary metabolites with variety of functions (Pichersky *et al.*, 2006). However, volatile esters formation is not restricted to plant kingdom but also in yeast and fungi especially in the fermentation industry (Beekwilder *et al.*, 2004). Floral scent is made up of a complex mixture of low-molecular-weight lipophilic compounds which are typically liquids with high vapour pressures (Vainstein *et al.*, 2001). With the discovery of novel techniques including gas chromatography-nuclear magnetic resonance (GC-NMR), gas chromatography-mass spectrometry (GC-MS), headspace based techniques in volatiles detection and analyses, the number of identified volatile compounds has increased tremendously (Gonzalez-Mas *et al.*, 2008; Mohd-Hairul *et al.*, 2010; Nojima *et al.*, 2011).

Most of the volatile compounds are derived from three major biosynthesis pathways: phenylpropanoids, fatty acid derivatives and terpenoids. They are thus classified into three major categories: terpenes, lipid derivatives and aromatic compounds. The terpenes are the largest class of plant volatiles, which consist of monoterpene alcohols and sesquiterpenes. There are also other terpene derivatives like ketones that are present in low quantities but have significant contributions to the floral fragrance. The second largest family of plant volatiles is the aromatic compounds. Most of them are derived from the intermediates in the benzenoid biosynthesis pathway that resulted in the synthesis of phenylalanine from the shikimate pathway, followed by a wide range of primary metabolites (eugenol, a lignin precursor, is one of them) and secondary non-volatile compounds (this was well reviewed in Bick & Lange, 2003; Pichersky & Dudareva, 2007).

The third category of plant volatiles is the lipid derivatives from the oxidative cleavage and decarboxylation of a variety of fatty acids which lead to shorter-chain volatiles with aldehyde and ketone moieties formation (reviewed in Baysal & Demirdoven, 2007). There are also other plant volatiles especially those with nitrogen or sulfur, which are produced through the cleavage of modified amino acids or their precursors (Cherri *et al.*, 2007; Pichersky *et al.*, 2006).

2.2. Discovery of fragrance-related genes

Due to the invisibility of floral scents and its dynamic nature, the study on flower scent is limited. There is no convenient plant model system that allows chemical and biochemical studies on floral scents. The well-established *Arabidopsis* as a plant model system failed to serve this purpose as the detection of volatiles production by its flower was barely detectable (Vainstein *et al.*, 2001). To date, the characterization and elucidation of enzymes and genes involved in flower scent production are still not as advanced as the biochemical study on the scent constituents. *Petunia hybrid* is one of the plant model systems that has been used to study the biological importance of floral scent (Spitcer *et al.*, 2007). However, limited information is available for the floral fragrance synthesized in petunia flowers as well as its cell specificity in fragrance production (van Moerkercke *et al.*, 2012).

'Scent' enzymes can be identified through *in vitro* characterization of their enzymatic activities with substrates predicted to be the precursors of known products in the tissues from which they are derived (Dudareva *et al.*, 2004). Besides, they can also be determined by identifying the homologue sequences in genomic or EST databases, expressing the protein in heterologous system like bacterial system followed by biochemical testing of the enzymatic activities with various substrates (Adelene *et al.*, 2012; Bradbury, 2007).

For more than 400 orchids (including both species and hybrids) that were discovered to emit fragrance (Frowine, 2005), in-depth scientific studies on orchid fragrance barely covered 2 percents of the fragrant orchids. Sadly, fragrance study in orchid is not as well established as in other flowers such as rose (Guterman *et al.*, 2002), petunia (Verdonk *et al.*, 2003) and *Clarkia breweri* (Dudareva *et al.*, 1996; Dudareva *et al.*, 1998). The advent of GC-MS technique enabled the evaluation of floral scent components quantitatively and qualitatively (Vainstein *et al.*, 2001). Mohd-Hairul *et al.* (2010) showed that the scent of VMP was predominated by terpenoid, benzenoid and phenylpropanoid compounds through GC-MS analysis (Table 1). It was interesting to note that a comparison of the volatiles captured from VMP with both of its parents revealed that the scent was dissimilar to its fragrance parent, *V. tessellate*. Such biochemical data is useful for subsequent work in the identification of fragrance-related genes in VMP.

3. Expressed sequence tags (ESTs)

Expressed Sequence Tags (ESTs) are partial sequences generated from single-pass sequencing (5'- or 3'- end) from a reverse transcription of mRNA representing tissue of interest or a particular developmental stage of an organism (Adam *et al.*, 1991). In plant, EST approach was first used in the model plant *A. thaliana* for genome and *in vivo* analysis of their gene products (Hofte *et al.*, 1994). Generally, the generated ESTs are highly informative at the middle DNA sequences but its redundancy rate at both ends can reach up to 3% (Nagaraj *et al.*, 2006; Rudd, 2003). Thus, pre-processing of ESTs is required prior to further analyses.

EST libraries for many plant species such as *P. equestris* (Schauer) Rchb.f. (Tsai *et al.*, 2006; Tsai *et al.*, 2011), cycads (Brenner *et al.*, 2003), mints (Lindqvist *et al.*, 2006), apple (Newcomb *et al.*, 2006), *Jatropha curcas* (Chen *et al.*, 2011) and *Prunus mume* (Li *et al.*, 2010) have been developed and deposited into the GenBank database. One of the databases for the submission and deposition of ESTs sequences is the database of expressed sequence tags (dbEST). dbEST is a collection of cDNA sequences and ESTs information from different variety of organisms. From the dbEST released on April 1, 2012 (http://www.ncbi.nlm.nih.gov/dbEST/dbEST_summary.html), the publicly available ESTs data were 72,316,247.

3.1. Importance of ESTs

EST is a fast, efficient and valuable tool for gene expression, genome annotation and evolutionary studies. Analysis of ESTs provides a platform for functional genomics study as well as uncovering the potentially novel genes, and poses an avenue for genome sequencing

projects (Ayeh, 2008; Kisiel & Podkowinski, 2005; Li *et al.*, 2010; Lindqvist *et al.*, 2006). Early EST projects were focused mostly on economically important plants and crop plants. In subsequent years, more EST projects on plants yet-to-achieve high economical impact started to materialise. The development of ESTs for some of the plants species from the early years until now is summarised in Table 2.

EST also proves to be a beneficial resource for comparative genomic studies in plant. Hsiao *et al.* (2006) deduced monoterpene biosynthesis pathway and identified a few fragrance-related genes in *Phalaenopsis bellina* by making a comparison with the floral EST library of *P. equestris* (scentless species). EST is also useful in the development and mining of microsatellite markers such as simple sequence repeats (SSRs) (Alba *et al.*, 2004). Lindqvist *et al.* (2006) reported the development of mint's EST for the identification of genetic markers for Hawaiian endemic mints. Moreover, EST provides the basis for the understanding of metabolic regulation mechanisms (Chen *et al.*, 2011; Li *et al.*, 2010).

Peak	Relative retention time (min)	Main spectrum fragments (m/z)	Compound name
Monoterpene			
1	8.636	36,41,53,67,79,93,105,121	Ocimene
2	9.147	41,43,59,81,93,112	Linalool oxide
3	9.592	41,43,69,71,93,107,121,136	Linalool
Sesquiterpene			
10	16.492	41,43,69,71,93,107,123,136, 162	Nerolidol
Benzenoid			
4	9.702	51,77,105,136	Methylbenzoate
5	10.030	39,51,65,78,91,105,122	Benzyl acetate
Phenylpropanoid			
6			
8	10.783	39,43,65,79,91,108,150	Phenylethanol
	11.926	39,43,65,78,91,104	Phenylethyl acetate
Indole			
9	13.084	39,50,63,74,90,117	Indole
Formanilide			
7	12.260	39,52,65,76,93,161	Formanilide

Table 1. Volatile compounds emitted by fully open flower of *Vanda Mimi Palmer* with **their relative retention times and spectral fragments**. This table is adapted from Mohd-Hairul *et al.* (2010).

Organism	ESTs
<i>Zea mays</i> (maize)	2,019,114
<i>Arabidopsis thaliana</i> (thale cress)	1,529,700
<i>Glycine max</i> (soybean)	1,461,624
<i>Oryza sativa</i> (rice)	1,252,989
<i>Triticum aestivum</i> (wheat)	1,073,877
<i>Panicum virgatum</i> (switchgrass)	720,590
<i>Brassica napus</i> (oilseed rape)	643,874
<i>Hordeum vulgare</i> + subsp. <i>vulgare</i> (barley)	501,838
<i>Vitis vinifera</i> (wine grape)	446,639
<i>Nicotiana tabacum</i> (tobacco)	334,384
<i>Pinus taeda</i> (loblolly pine)	328,662
<i>Malus x domestica</i> (apple tree)	324,742
<i>Piceaglauca</i> (white spruce)	313,110
<i>Gossypium hirsutum</i> (upland cotton)	297,239
<i>Solanum lycopersicum</i> (tomato)	297,142
<i>Medicago truncatula</i> (barrel medic)	269,238
<i>Solanum tuberosum</i> (potato)	249,761
<i>Lotus japonicas</i>	242,432
<i>Mimulus guttatus</i>	231,095
<i>Raphanus sativus</i> (radish)	110,006
<i>Petunia x hybrid</i>	50,705
<i>Zingiber sp.</i>	38,190
<i>Elaeis guineensis</i>	40,737
<i>Petunia axillaris</i> subsp. <i>axillaris</i>	11,078
<i>Phalaenopsis equestris</i>	5,604
<i>Rosa hybrid cultivar</i>	5,565
<i>Phalaenopsis violacea</i>	2,359
<i>Dendrobium officinale</i>	800
<i>Oncidium hybrid cultivar</i>	280
<i>Phalaenopsis amabilis</i>	103

Table 2. Summary of some of the plants species available in dbEST from the early years until 1 April 2012.

3.2. Discovery of fragrance-related genes from VMPESTs

Expressed sequence tags (ESTs) have been developed from many monocot plant species including *P. equestris* (Tsai *et al.*, 2006), ginger (Chandasekar *et al.*, 2009), oil palm (Low *et al.*, 2008), wheat (Zhang *et al.*, 2004) and maize (Fernandes *et al.*, 2002). To date, more than 1700 floral fragrance compounds have been identified through biochemical analysis with the majority identified from higher plants whilst others are from animals, insects, marine organisms, algae and bacteria. Unfortunately, the developments of the biochemical aspects of fragrance compounds are not in par with the molecular information on fragrance-related genes. More so, reported molecular information from the vandaceous orchids is still extremely scarce. Thus far, the only available fragrance-related molecular works on orchid are from the genera *Phalaenopsis* and *Vanda* (from our research group).

A VMP floral cDNA library was previously constructed from opened flowers at different developmental stages and time-points (Chan *et al.*, 2009). All the cDNA clones with the inserts sizes of 0.5 kb to 1.6 kb were mass excised and single-pass 5'-sequenced. From our attempt, a total of 2,132 ESTs was generated. This VMP dbEST (designated as VMPEST) was clustered, annotated and further classified with Gene Ontology (GO) identifier into three categories: Molecular Functions (51.2%), Cellular Components (16.4%) and Biological Processes (24.6%). Around 3.1% of the VMPEST had hits with other orchid species such as dendrobium, phalaenopsis, oncidium, *Aerides japonica*, and *Aranda Deborah*. A number of fragrance-related transcripts were identified (Table 3; Teh, 2011) by comparing Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways with the three major volatiles biosynthesis pathways (terpenoid, benzenoid and phenylpropanoid) of well-studied scented flowers such as *Antirrhinum majus*, *Clarkia breweri*, *Petunia* hybrid and *Rosa* hybrid (Boatright *et al.*, 2004; Lavid *et al.*, 2002).

From the VMPEST, several fragrance-related transcripts were selected for full-length isolation and expression analysis using real-time quantitative RT-PCR. They were *Vanda Mimi Palmer acetyl-CoA acetyltransferase* (designated as *VMPACA*, in press), *Vanda Mimi Palmer 3-hydroxy-3-methylglutaryl-coenzyme A reductase* (*VMPHMGR*), *Vanda Mimi Palmer 1-deoxy-D-xylulose 5-phosphate synthase* (*VMPDXPS*), *Vanda Mimi Palmer linalool synthase* (*VMP LIS*) and *Vanda Mimi Palmer lipoxygenase* (*VMP Lox*) (Teh, 2011). Gene specific primers were designed and synthesised for 5'-RACE targeting at the incomplete 5'-ends of each of the aforementioned gene transcript. The full length cDNA sequences of those transcripts were deduced, amplified, followed by sequencing and sequence analysis. Real-time quantitative RT-PCR were performed using cDNA templates extracted from different types of VMP tissues (sepals, petals, lips, stems, stalks, columns, roots and leaves), from full-bloom flowers taken at different time points (2-hour intervals within 24-hour duration) and from different flower developmental stages (buds, blooming and full-bloom flower). Overall, the expression profiles revealed that the transcripts exhibited a developmentally regulated pattern in the different flower developmental stages. Majority of the transcripts were highly expressed in the full-bloom stage, with the sepals and petals having the highest expression levels.

GenBank Accession	dbEST id	Putative identity	E-value	Score (bits)
GW392501	68671343	<i>3-hydroxy-3-methylglutaryl-coenzyme A reductase 2 [Gossypium hirsutum]</i>	2.00E-108	358
GW392566	68671408	<i>LOX1 (Lipoxygenase 1); lipoxygenase [Arabidopsis thaliana]</i>	1.00E-50	203
GW392695	68671537	<i>linalool synthase 2 [Clarkia breweri]</i>	9.00E-19	97.8
GW392657	68671499	<i>trans-caffeoyl-CoA 3-O-methyltransferase [Populus trichocarpa]</i>	2.00E-52	209
GW392731	68671573	<i>lipoxygenase 1 [Brassica napus]</i>	2.00E-15	85.9
GW392740	68671582	<i>acyltransferase [Vanda hybrid cultivar]</i>	3.00E-70	268
GW392813	68671655	<i>cinnamyl alcohol dehydrogenase [Populus trichocarpa]</i>	2.00E-73	279
GW393688	68672530	<i>carboxyl methyltransferase [Crocus sativus]</i>	5.00E-58	228
GW392895	68671737	<i>cinnamoyl-CoA reductase [Saccharum officinarum]</i>	2.00E-46	189
GW392922	68671764	<i>putative 1-deoxy-D-xylulose 5-phosphate synthase [Hevea brasiliensis]</i>	6.00E-62	241
GW393960	68672802	<i>putative acetyl-CoA C-acyltransferase [Oryza sativa Japonica Group]</i>	4.00E-103	378
GW393331	68672173	<i>lipoxygenase 1 [Brassica napus]</i>	4.00E-27	125
GW393619	68672461	<i>resveratrol O-methyltransferase [Vitis vinifera]</i>	8.00E-56	221
GW393628	68672470	<i>s-adenosylmethionine synthetase, putative [Ricinus communis]</i>	2.00E-59	233
GW394168	68673010	<i>linalool synthase-like protein [Oenothera arizonica]</i>	6.00E-41	171
GW393499	68672341	<i>acetyl-CoA acetyltransferase, cytosolic 1 [Zea mays]</i>	1.00E-136	488

Table 3. Selected putative fragrance-related ESTs generated from *Vanda Mimi Palmer*.

This table is adapted from Teh *et al.* (2011).

Among all the transcripts analysed, *VMPHMGR* was selected for functional analysis. *HMGR* is a very well-studied enzyme in cholesterol synthesis especially in animal system. A functional enzymatic assay performed showed *VMPHMGR* was functionally active in *Escherichia coli*, catalysing the conversion of *HMG-CoA* to mevalonate derivatives, which are commonly used in the metabolic biosynthesis of steroids, terpenoids and carotenoids (Teh, 2011). This *VMPHMGR* shows high sequence identity to the *HMGR* found in other plant species, with 76% sequence similarity with *Oryza sativa* Indica group. Although its N-terminal end differs distinctly in length and amino acids compositions, its C-terminal catalytic domain shows high sequence similarity with other plant species (unpublished data). This highlights the importance of information and evolution of gene of similar function in different organisms.

4. Simple sequence repeat and its importance

Simple Sequence Repeat (SSR) or microsatellite is a short tandem repeats of a unique DNA sequence with one to six nucleotides motif (Jacob *et al.*, 1991). SSR is a famous molecular marker because of its hyper variability, relative abundance, highly reproducible, multiallelic diversity, co-dominantly inherited and extensive coverage of the genome (Mohan *et al.*, 1997).

Owing to its desirable genetic attributes, SSRs have been utilized in genetic and genomic analyses including genetic mapping, marker assisted plant breeding, development of linkage map, and ecology studies (Kalia *et al.*, 2011; Sonah *et al.*, 2011; Yue *et al.*, 2006). Yue *et al.* (2006) reported the usage of SSRs in the protection of new *Dendrobium* varieties.

To date, the used of SSRs markers have been reported in several monocot and dicot species including raspberry and blackberry (Stafne *et al.*, 2005), rice (Chakravarthi & Naravaneni, 2006), common bean (Yu *et al.*, 2000), *Brachypodium* (Sonah *et al.*, 2011) and *Dendrobium* (Yue *et al.*, 2006). However, the effort to develop SSR markers for orchids is limited to several species: Chinese orchid [*Cymbidium* spp.] (Huang *et al.*, 2010), *Phalaenopsis* (Bory *et al.*, 2008; Hsu *et al.*, 2011), Brazilian orchid (*Epidendrum fulgens*) (Pinheiro *et al.*, 2008), *Dendrobium* (Yue *et al.*, 2006) and *Vanda* (Phuekvilai *et al.*, 2009).

So far, the reported SSRs generated from vandaceous orchids were used as selective marker only. Phuekvilai *et al.* (2009) generated SSRs from 33 vandas species for the sole purpose of identifying and evaluating the purity of cultivar in commercial samples. However, the identification and development of SSRs for VMP will be channelled towards facilitating the screening of any potential fragrance-related transcripts from closely related species. Besides, it will be used to determine the extent of inter-species transferability of genes, which had been reported in many plant species (Chapman *et al.*, 2009; Stafne *et al.*, 2005; Wang *et al.*, 2004).

4.1. Data mining of VMPEST-SSR

In recent years, genic microsatellite or EST-SSRs which is less time consuming and relatively easy to develop has replaced the genomics SSRs (Sharma *et al.*, 2007). The publicly available ESTs sequences facilitate the development of SSRs by using the SSR identification tools. These search tools include MISA (MicroSATellite), SSRIT (SSR Identification Tool), SciRoKo, TRF (Tandem Repeat Finder), Sputnik, SSRfinder, SAT (SSR Analysis Tool), Poly and SSR Primer. It is deemed important to choose a search tool which is user-friendly and has unlimited access to a non-redundant database (this was well reviewed by Kalia *et al.*, 2011). The first attempt to develop such EST-SSR marker was in rice by Miyao *et al.* (1996).

SSRIT which is accessible at URL (<http://www.gramene.org>) was used to identify the SSR motifs in our VMPEST. The script assessed the sequences uploaded in FASTA-formatted files and detected the SSR motifs, the number of repeats as well as identified the sequence corresponded to the SSRs. A total of 98 (9.4%) unigenes containing 112 SSRs with motifs length ranging from two to six nucleotides were detected from VMPEST.

The VMPEST-SSRs were classified into 2 groups with 88.4% belonging to Class I ($n \geq 20$ nucleotides) and 11.6% belonging to Class II ($12 \leq n \leq 20$ nucleotides) according to their lengths and genetic marker potential (Teh *et al.*, 2011). Such groupings had been reported in rice by Temnykh *et al.* (2001). In their study, they revealed that higher rates of polymorphism occurred in transcripts with longer SSR sequences. Likewise, Song *et al.* (2010) reported that SSR which covered longer portion of sequence (repeat number less than 35) were deemed better for development of genetic markers because they associated with expressed portion of the genome, thus facilitate better understanding of associated protein(s). From our study, majority of the EST-SSR sequences were categorised as 'ideal' repeat ($n \geq 20$ nucleotides), which correlated well to the SSRs mined from tea [*Camellia sinensis* L.] (Sharma *et al.*, 2009).

Sharma *et al.* (2009) stressed that the length of repeats and the tools used in the EST-SSRs mining play a significant role in EST-SSR occurrences. The di-nucleotide motif (AT/TA) was present with the most abundance (33.9%) in our VMPEST-SSR. Such observation of A- and T-rich signatures being the most common repeat motifs in the VMPEST-SSRs is also reflected in the early findings of Chagne *et al.* (2004), Lagercrantz *et al.* (1993), and Morgante & Olivieri (1993). Interestingly, Blair *et al.* (2009) found that this motif occurred mainly in the 3'-end of the common bean cDNA clones.

Nevertheless, whatever mined results we obtained from our study, each SSR needs to be validated with further analyses such as selection of SSR primers and screening for utility in vandaceous orchids.

Besides EST-SSR, there are other alternative data mining techniques such as expressed sequence tag-single nucleotide polymorphism (EST-SNP) in maize (Batley *et al.*, 2003), barley (Varshney *et al.*, 2007), melon (Deleu *et al.*, 2009), citrus (Jiang *et al.*, 2010), and cocoa (Allegre *et al.*, 2011), expressed sequence tag-rapid amplified polymorphic DNA (EST-RAPD) in oil palm (Balakrishnan & Vadivel, 2012), and expressed sequence tag-sequence tagged site (EST-STS) in wheat (Leonard *et al.*, 2008; Naji *et al.*, 2008). All the aforementioned techniques are useful for functional genetic diversity estimation of GenBank collections and valuable for use in marker-assisted programmes. However, each technique has its good and down sides that might affect its eventual development.

5. Conclusion

The VMPEST dataset is a potential asset in facilitating the molecular biology and cloning of more genes involved in the fragrance biosynthesis pathway(s). Several fragrance-related transcripts were identified from our VMPEST including *VMPACA*, *VMPHMGR*, *VMPDXPS*, *VMPLox* and *VMPLis*. The functional enzymatic assay that was performed on the selected transcript (*VMPHMGR*) proved to be functionally active in its catalysis reaction in a heterologous system. The detected SSRs loci and microsatellite motifs that had hits with fragrance-related genes in the GenBank are believed to be a valuable resource especially to researchers involved in studying diversity to access the functional diversity of fragrant vandaceous orchids and their linkages to other orchids.

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6. References

- Aceto, S. & Gaudio, L. (2011). The MADS and the beauty: genes involved in the development of orchid flowers. *Current Genomics*, 12, 342-356.
- Adam, M. D., Kelley, J. M., Gocayne, J. D., Dubnick, M., Polymeropoulos, M. H., Xiao, H., Merril, C. R., Wu, A., Olde, B., Moreno, R. F., Kerlavage, A. R., McCombie, W. R. & Venter, J. C. (1991). Complementary DNA sequencing: expressed sequence tags and human genome project. *Science*, 252, 1651-1656.
- Adelene, S. A. L., Janna, O. A., Mohd, P., Norazizah, S., Raha, A.R. & Puad, A. (2012). Functional expression of an orchid fragrance gene in *Lactococcus lactis*. *International Journal of Molecular Sciences*, 13, 1582-1597.
- Agrawal, A., Janssen, A., Bruin, J., Posthumus, M. & Sabelis, M. (2002). An ecological cost of plant defence: attractiveness of bitter cucumber plants to natural enemies of herbivores. *Ecological Letters*, 5, 377-385.
- Ajjikuttira, P. A., Woon, M. H., Ryu, K. H., Chang, C. A., Loh, C. S., Wong, S. M. & Ridge, K. (2002). Genetic variability in the coat protein genes of two orchid viruses: cymbidium mosaic virus and odontoglossum ringspot virus. *Archives of Virology*, 147, 1943-1954.
- Alba, R., Fei, Z. J., Liu, Y., Moore, S., Debbie, P., Cohn, J., Ascenzo, M., Gordon, J., Rose, J., Martin, G., Tanksley, S., Bouzayen, M., Jahn, M. & Giovannoni, J. (2004). ESTs, cDNA microarrays and gene expression profiling: tools for dissecting plant physiology and development. *The Plant Journal*, 39, 697-714.
- Allegre, M., Argout, X., Boccara, M., Fouet, O., Roguet, Y., Berard, A., Thevenin, J. M., Chauveau, A., Rivallan, R., Clement, D., Courtois, B., Gramacho, K., Boland-Auge, A., Tahy, M., Umaharan, P., Brunel, D. & Lanaud C. (2011). Discovery and mapping of a new expressed sequence tag-single nucleotide polymorphism and simple sequence repeat panel for large-scale genetic studies and breeding of *Theobroma cacao* L. *DNA Research*, doi: 10.1093/dnares/dsr039
- Arditti, J. (1992). *Fundamentals of orchid biology*, John Wiley & Sons Inc., New York, USA.

- Arimura, G., Ozawa, R., Kugimiya, S., Takabayashi, J. & Bohlmann, J. (2004). Herbivore-induced defense response in a model legume: two-spotted spider mites, *Tetranychus urticae*, induce emission of (*E*)- β -ocimene and transcript accumulation of (*E*)- β -ocimene synthase in *Lotus japonicus*. *Plant Physiology*, 135, 1976-1983.
- Ayeh, K. O. (2008). Expressed sequence tags (ESTs) and single nucleotide polymorphisms (SNPs): emerging molecular marker tools for improving agronomic traits in plant biotechnology. *African Journal of Biotechnology*, 7, 331-341.
- Balakrishnan, V. P. & Vadivel, A. (2012). In silico RAPD priming sites in expressed sequences and iSCAR markers for oil palm. *Comparative and Functional Genomics*, doi: 10.1155/2012/913709
- Batley, J., Barker, G., O'Sullivan, H., Edwards, K. J. & Edwards, D. (2003). Mining for single nucleotide polymorphisms and insertions/deletions in maize expressed sequence tag data. *Plant Physiology*, 132, 84-91.
- Baudino, S., Caissard, J-C, Bergougnoux, V., Jullien, F., Magnard, J-L, Scalliet, G., Cock, J. M. & Hugueney, P. (2007). Production and emission of volatile compounds by petal cells. *Plant Signal Behaviour*, 2, 525-526,
- Baysal, T. & Demirdoven, A. (2007). Lipoxygenase in fruits and vegetables: A review. *Enzyme and Microbial Technology*, 40, 491-496.
- Beekwilder, J., Alvarez-Huerta, M, Neef, E., Verstappen, F., Bouwmeester, H.& Aharoni, A. (2004). Functional characterization of enzymes forming volatile esters from strawberry and banana. *Plant Physiology*, 135, 1865-1878.
- Bick, J. A. & Lange, M. (2003). Metabolic cross talk between cytosolic and plastidial pathways of isoprenoids biosynthesis: unidirectional transport of intermediates across the chloroplast envelope membrane. *Archives of Biochemistry and Biophysics*, 415, 146-154.
- Blair, M. W., Torres, M. M., Giraldo, M. C. & Pedraza, F. (2009). Development and diversity of Andean-derived, gene-based microsatellites for common bean (*Phaseolus vulgaris* L.). *BMC Plant Biology*, 9, 100-113.
- Boatright, J., Negre, F., Chen, X., Kish, C., Wood, B., Peel, G., Orlova, I., Gang, D., Rhodes, D. & Dudareva, N. (2004). Understanding *in vivo* benzenoid metabolism in petunia petal tissue. *Plant Physiology*, 135, 1993-2011.
- Bory, S., Silva, D., Risterucci, A., Grisoni, M., Besse, P. & Duval, M. (2008). Development of microsatellite markers in cultivated vanilla: Polymorphism and transferability to other vanilla species. *Scientia Horticulturae*, 115, 420-425.
- Bradbury, L. (2007). *Identification of the gene responsible for fragrance in rice and characterization of the enzyme transcribed from this gene and its homologs*. Doctor of Philosophy dissertation, Southern Cross University, Australia.
- Brenner, E. D., Stevenson, D. W., McCombie, R. W., Katari, M. S., Rudd, S. A., Mayer, K. F., Palenchar, P. M., Runko, S. J., Twigg, R. W., Dai, G., Martienssen, R. A., Benfey, P. N. & Coruzzi, G. M. (2003). Expressed sequence tag analysis in *Cycas*, the most primitive living seed plant. *Genome Biology*, 4, 78-88.
- Chagne, D., Chaumeil, P., Ramboer, A., Collada, C., Guevara, A., Cervera, M. T., Vendramin, G. G., Garcia, V., Frigerio, J-M, Echt, C., Richardson, T. & Plomion, C. (2004). Cross-species transferability and mapping of genomic and cDNA SSRs in pines. *Theoretical and Applied Genetics*, 109, 1204-1214.

- Chai, D. & Yu, H. (2007). Recent advances in transgenic orchid production. *Orchid Science and Biotechnology*, 1, 34-39.
- Chakravarthi, B. K. & Naravaneeni, R. (2006). SSR marker based DNA fingerprinting and diversity study in rice (*Oryza sativa* L.). *African Journal of Biotechnology*, 5, 684-688.
- Chan, W. S., Janna, O. A., Parameswari, N. & Maziah, M. (2009). Molecular characterization of a new 1-deoxy-D-xylulose 5-phosphate reductoisomerase (DXR) transcript from *Vanda Mimi Palmer*. *Scientia Horticulturae*, 121, 378-382.
- Chan, W. S., Janna, O. A. & Parameswari, N. (2011). Isolation, cloning and characterization of fragrance-related transcripts from *Vanda Mimi Palmer*. *Scientia Horticulturae*, 127, 388-397.
- Chandasekar, A., Riju, A., Sithara, K., Anoop, S. & Eapen, S. J. (2009). Identification of single nucleotide polymorphism in ginger using expressed sequence tags. *Bioinformatics*, 4, 119-122.
- Chapman, M. A., Hvala, J., Strever, J., Matvienko, M., Kozik, A., Michelmore, R. W., Tang, S., Knapp, S. J. & Burke, J. M. (2009). Development, polymorphism and cross-taxon utility of EST-SSR markers from safflower (*Carthamus tinctorius* L.). *Theoretical and Applied Genetics*, 120, 85-91.
- Chen, M. S., Wang, G. J., Wang, R. L., Wang, J., Song, S. Q. & Xu, Z. F. (2011). Analysis of expressed sequence tags from biodiesel plant *Jatropha curcas* embryos at different developmental stages. *Plant Science*, doi:10.1016/j.plantsci.2011.08.003
- Cherri, M., Jullien, F., Heizmann, P. & Baudino, S. (2007). Fragrance heritability in hybrid tea roses. *Scientia Horticulture*, 113, 177-181.
- Deleu, W., Esteras, C., Roig, C., Gonzalez-To, M., Fernandez-Silva, I., Gonzalez-Ibeas, D., Blanca, J., Aranda, M. A., Arus, P., Nuez, F., Monforte, A. J., Pico, M. B. & Garcia-Mas, J. (2009). A set of EST-SNPs for map saturation and cultivar identification in melon. *BMC Plant Biology*, 9, 90, doi:10.1186/1471-2229-9-90
- Dudareva, N., Cseke, L., Blanc, V. & Pichersky, E. (1996). Evolution of floral scent in *Clarkia*: novel patterns of S-linalool synthase gene expression in the *C. breweri* flower. *Plant Cell*, 8, 1137-1148.
- Dudareva, N., D'Auria, J., Nam, K. H., Raguso, R. & Pichersky, E. (1998). Acetyl-CoA:benzylalcohol acetyltransferase - an enzyme involved in floral scent production in *Clarkia breweri*. *The Plant Journal*, 14, 297-304.
- Dudareva, N., Pichersky, E. & Gershenzon, J. (2004). Biochemistry of plant volatiles. *Plant Physiology*, 135, 1839-1902.
- Eun, A. J. & Wong, S. (2000). Molecular beacons: a new approach to plant virus detection. *Phytopathology*, 90, 269-275.
- Eun, A. J., Huang, L., Chew, F., Li, S. F. & Wong, S. (2002). Detection of two orchid viruses using Quartz crystal microbalance-based DNA biosensors. *Phytopathology*, 92, 654-658.
- Fernandes, J., Brendel, V., Gai, X., Lal, S., Chandler, V. L., Elumalai, R. P., Galbraith, D. W., Pierson, E. A. & Walbot, V. (2002). Comparison of RNA expression profiles based on maize expressed sequence tag frequency analysis and micro-array hybridization. *Plant Physiology*, 128, 896-910.
- Frowine, S. A. (2005). *Fragrant orchids*, Timber Press Inc., Oregon, USA.

- Go, R., Yong, W. S. Y., Unggang, J. & Salleh, R. (2010). *Orchids of Perlis, Jewels in the Forests*, Revised Edition. Perlis Forestry Department and Universiti Putra Malaysia Press, Kuala Lumpur, Malaysia, p152.
- Go., R., Khor, H. E., Tan, M. C., Janna, O. A., Farah Alia, N., Ng. Y. J., Muskhazli, M., Mohd Nazre, S. & Rosimah, N. (2012). Current knowledge on orchid diversity in Peninsular Malaysia including new records and discovery. (*Manuscript submitted and under review*)
- Gonzalez-Mas, M. C., Garcia-Riano, L. M., Alfaro, C., Rambla, J. L., Padilla, A. I. & Gutierrez, A. (2008). Headspace-based techniques to identify the principal volatile compounds in red grape cultivars. *International Journal of Food Science & Technology*, 44, 510-518.
- Guterman, I., Shalit, M., Menda, N., Piestun, D., Dafny, Y., Shalev, G., Bar, E., Davydov, O., Ovadis, M., Emanuel, M., Wang, J., Adam, Z., Pichersky, E., Lewinsohn, E., Zamir, D., Vainstein, A. & Weiss, D. (2002). Rose scent: genomics approach to discovering novel floral fragrance-related genes. *The Plant Cell*, 14, 2325-2338.
- Hofte, H., Desprez, T., Amselem, J., Chiapello, H., Rouze, P., Caboche, M., Moisan, A., Jourjon, M. F., Charpentreau, J. L., Berthomieu, P., Guerrier, D., Giraudat, J., Quigley, F., Thomas, F., Yu, D. Y., Raynal, M., Cooke, R., Grellet, F., Marcillac, P., Gigot, C., Fleck, J., Phillips, G., Axelos, M., Bardet, C., Tremousaygue, D. & Lescure, B. (1994). An inventory of 1152 expressed sequence tags obtained by partial sequencing of cDNAs from *Arabidopsis thaliana*. *Plant Journal*, 4, 1051-1061.
- Hsiao, Y., Tsai, W., Kuoh, C., Huang, T., Wang, H., Wu, T., Leu, Y., Chen, W. & Chen, H. (2006). Comparison of transcripts in *Phalaenopsis bellina* and *Phalaenopsis equestris* (Orchidaceae) flowers to deduce monoterpene biosynthesis pathway. *BMC Plant Biology*, 6, 14-27.
- Hsiao, Y. Y., Pan, Z. J., Hsu, C. C., Yang, Y. P., Hsu, Y. C., Chuang, Y. C., Shih, H. H., Chen, W. H., Tsai, W. C. & Chen, H. H. (2011). Research on Orchid Biology and Biotechnology. *Plant and Cell Physiology*, 52(9), 1467-1486.
- Hsu, C. C., Chung, Y. L., Chen, T. C., Lee, Y. L., Kuo, Y. T., Tsai, W. C., Hsiao, Y. Y., Chen, Y. W., Wu, W. L. & Chen, H. H. (2011). An overview of the *Phalaenopsis* orchid genome through BAC end sequence analysis. *BMC Plant Biology*, 11, 3-13.
- Hsu, H. T., Vongsasitorn, D. & Lawson, R. H. (1992). An improved method for serological detection of *cymbidium* mosaic potexvirus infection in orchids. *Phytopathology*, 82, 491-495.
- Hu, W. W. & Wong, S. M. (1998). The use of DIG-labelled cRNA probes for the detection of *cymbidium* mosaic potexvirus (CymMV) and *odontoglossum* ringspot tobamovirus (ORSV) in orchids. *Journal of Virological Methods*, 70, 193-199.
- Huang, Y., Li, F. & Chen, K. (2010). Analysis of diversity and relationships among Chinese orchid cultivars using EST-SSR markers. *Biochemical Systematics and Ecology*, 38, 93-102.
- Jacob, H. J., Lindpaintner, K., Lincoln, S. E., Kusumi, K., Bunker, R. K., Mao, Y. P., Ganten, D., Dzau, V. J. & Lander, E. S. (1991). Genetic mapping of a gene causing hypertensive rat. *Cell*, 67, 213-224.
- Janna, O. A., Tan, L., Aziz, A., Maziah, M., Puad, A. & Azlan, J. (2005). Dull to behold, good to smell... *Vanda Mimi Palmer*. *BioTech Communications*, Faculty of Biotechnology & Biomolecular Science Research Bulletin, Universiti Putra Malaysia, Malaysia, volume 1, pp21-25, ISSN 1823-3279.

- Jiang, D., YE, Q. L., Wang, F. S. & Cao, L. (2010). The mining of citrus EST-SNP and its application in cultivar discrimination. *Agricultural Sciences in China*, 9, 179-190.
- Kalia, R. K., Rai, M. K., Kalia, S., Singh, R. & Dhawan, A. K. (2011). Microsatellite markers: an overview of the recent progress in plants. *Euphytica*, 177, 309-334.
- Khentry, Y., Paradornuwat, A., Tantiwiwat, S., Phransiri, S. & Thaveechai, N. (2006). Incidence of cymbidium mosaic virus and odontoglossum ringpot virus in *Dendrobium* spp. in Thailand. *Crop Protection*, 25, 926-932.
- Kishor, R., ShaValli Khan, P. S. & Sharma, G. J. (2006). Hybridization and *in vitro* culture of an orchid hybrid *Ascocenda 'Kangla'*. *Scientia Horticulturae*, 108, 66-73.
- Kisiel, A. & Podkowinski, J. (2005). Expressed Sequence Tags and their applications for plants research. *Acta Physiologiae Plantarum*, 27, 157-161.
- Kishor, R., Devi, H., Jeyaram, K. & Singh, M. (2008). Molecular characterization of reciprocal crosses of *Aerides vandarum* and *Vanda stangeana* (Orchidaceae) at the protocorm stage. *Plant Biotechnology Reports*, 2, 145-152.
- Lagercrantz, U., Ellegren, H. & Anderson, L. (1993). The abundance of various polymorphic microsatellite motifs differs between plants and vertebrates. *Nucleic Acids Research*, 21, 1111-1115.
- Lavid, N., Wang, J., Shalit, M., Guterman, I., Bar, E., Beuerle, T., Menda, N., Shafir, S., Zamir, D., Adam, Z., Vainstein, A., Weiss, D., Pichersky, E. & Lewinsohn, E. (2002). O-methyltransferases involved in the biosynthesis of volatile phenolic derivatives in rose petals. *Plant Physiology*, 129, 1899-1907.
- Leonard, J., Watson, C., Carter, A., Hansen, J., Zemetra, R., Santra, D., Campbell, K. & Riera-Lizarazu, O. (2008). Identification of a candidate gene for the wheat endopeptidase *Ep-D1* locus and two other STS markers linked to the eyespot resistance gene *Psh1*. *Theoretical & Applied Genetics*, 116, 261-270.
- Li, X., Shanguan, L., Song, C., Wang, C., Gao, Z., Yu, H. & Fang, J. (2010). Analysis of expressed sequence tags from *Prunus mume* flower and fruit and development of simple sequence repeat markers. *BMC Genetics*, 11, 66-76.
- Lindqvist, C., Scheen, A., Yoo, M., Grey, P., Oppenheimer, D. G., Leebens-Mack, J. H., Soltis, D. E., Soltis, P. S. & Albert, V. A. (2006). An expressed sequence tag (EST) library from developing fruits of an Hawaiian endemic mint (*Stenogyne rugosa*, Lamiaceae): characterization and microsatellite markers. *BMC Plant Biology*, 6, 16-30.
- Low, E. T., Alias, H., Boon, S. H., Shariff, E., Tan, C. Y., Ooi, L., Cheah, S. C., Raha, A. R., Wan, K. L. & Singh, R. (2008). Oil palm (*Elaeis guineensis* Jacq.) tissue culture ESTs: Identifying genes associated with callogenesis and embryogenesis. *BMC Plant Biology*, 8, 62-81.
- Majetic, C., Raguso, R. & Ashman, T. (2009). The sweet smell of success: floral scent affects pollinators attraction and seed fitness in *Hesperis matronalis*. *Functional Ecology*, 23, 480-487.
- Micheneau, C., Fournel, J., Gaucin-Bialecki, A. & Paillet, T. (2008). Auto-pollination in a long-spurred endemic orchid (*Jumellea stenophylla*) on Reunion Island (Mascarene Archipelago, Indian Ocean). *Plant Systematics and Evolution*, 272, 11-22.
- Miyao, A., Zhong, H., Monna, L., Yano, M., Yamamoto, K., Havukkala, I. & Minobe, Y. (1996). Characterization and genetic mapping of simple sequence repeats in rice genome. *DNA Research*, 3, 233-238.

- Mohan, M., Nair, S., Bhagwat, A., Krishna, T. G., Yano, M., Bhatia, C. R. & Sasaki, T. (1997). Genome mapping, molecular markers and marker-assisted selection in crop plants. *Molecular Breeding*, 3, 87-103.
- Mohd-Hairul, A. R., Parameswari, N., Gwendoline Ee, C. L. & Janna O. A. (2010). Terpenoid, benzenoid and phenylpropanoid compounds in the floral scent of *Vanda Mimi Palmer*. *Journal of Plant Biology*, 53, 358-366.
- Morgante, M. & Olivieri, A. M. (1993). PCR-amplified microsatellites as markers in plant genetics. *Plant Journal*, 3, 175-182.
- Nagaraj, S., Gasser, R. & Ranganathan, S. (2006). A hitchhiker's guide to expressed sequence tag (EST) analysis. *Briefing in Bioinformatics*, 8, 6-21.
- Naji, A. M., Moghaddam, M., Ghaffari, M. R., Irandoost, H. P., Farsad, L. K., Pirseyedi, S. M., Mohammadim S. A., Ghareyazie, B. & Mardi, M. (2008). Validation of EST-derived STS markers localized on Qfhs.ndsu-3BS for *Fusarium* head blight resistance in wheat using a 'Wangshuibai' derived population. *Journal of Genetics & Genomics*, 35, 625-628.
- NewComb, R. D., Crowhurst, R. N., Gleave, A. P., Rikkerink, E., Allan, A. C., Beuning, L. L., Bowen, J. H., Gera, E., Jamieson, K. R., Janssen, B. J., Laing, W. A., McArtney, S., Nain, B., Ross, G. S., Snowden, K. C., Souleyre, E., Walton, E. F. & Yauk, Y. K. (2006). Analyses of expressed sequence tags from apple. *Plant Physiology*, 141, 147-166.
- Nojima, S., Kiemle, D. J., Webster, F. X., Apperson, C. S. & Schal, C. (2011). Nanogram-scale preparation and NMR analysis for mass-limited small volatile compounds. *PLoS ONE*, 6: e18178. Doi:10.1371/journal.pone.0018178
- Ooi, J. M. (2005). *Screening of a random peptide library with CymMV for potential development of diagnostic kits*. Master dissertation, Malaysia University of Science and Technology, Selangor Darul Ehsan, Malaysia.
- Park, Y. S., Jeong, M. H., Lee, S. H., Moon, J. S., Cha, J. S., Kim, H. Y. & Cho, T. J. (2005). Activation of defense responses in chinese cabbage by a nonhost pathogen, *Pseudomonas syringae* pv. tomato. *Journal of Biochemistry and Molecular Biology*, 38, 748-754.
- Phuekvilai, P., Pongtongkam, P. & Peyachoknagul, S. (2009). Development of microsatellite markers for *Vanda* orchid. *Natural Science*, 43, 497-506.
- Pichersky, E. & Dudareva, N. (2007). Scent engineering: toward the goal of controlling how flowers smell. *TRENDS in Biotechnology*, 25, 105-110.
- Pichersky, E., Noel, J. & Durareva N. (2006). Biosynthesis of plant volatiles: nature's diversity and ingenuity. *Science*, 311, 808-811.
- Pinheiro, F., Santos, M. O., Barros, F., Meyer, D., Salatino, A., Souza, A. P. & Cozzolino, S. (2008). Isolation and characterization of microsatellite loci in the Brazilian orchid *Epidendrum fulgens*. *Conservation Genetics*, 9, 1661-1663.
- Rada, F. & Jaimez, R. (1992). Comparative ecophysiology and anatomy of terrestrial and epiphytic *Anthurium bredemeyeri* Schott in a tropical Andean cloud forest. *Journal of Experimental Botany*, 43, 723-727.
- Raguso, R. (2008). Wake up and smell the roses: The ecology and evolution of floral scent. *Annual Review of Ecology, Evolution and Systematics*, 39, 549-569.
- Rani, P., Pant, R. & Jain, R. (2010). Serological detection of cymbidium mosaic and odontoglossumringspot viruses in orchids with polyclonal antibodies produced against their recombinant coat proteins. *Journal of Phytopathology*, 158, 542-545.

- Rudd, S. (2003). Expressed sequence tags: alternative or complement to whole genome sequences? *TRENDS IN Plant Science*, 8, 321-328.
- Schiestl, F. P. (2010). The evolution of floral scent and insect chemical communication. *Ecology Letters*, 13, 643-656.
- Sharma, P., Grover, A. & Kahl, G. (2007). Mining microsatellites in eukaryotic genomes. *TRENDS in Biotechnology*, 25, 490-498.
- Sharma, R., Bhardwaj, P., Negi, R., Mohapatra, T. & Ahuja, P. (2009). Identification, characterization and utilization of unigenes derived microsatellite markers in tea (*Camellia sinensis* L.). *BMC Plant Biology*, 9, 53-76.
- Sherpa, A., Hallan, V., Pathak, P. & Zaidi, A. (2007). Complete nucleotide sequence analysis of cymbidium mosaic virus Indian isolate: further evidence for natural recombination among potexviruses. *Journal of Bioscience*. 32, 663-669.
- Shuttleworth, A. & Johnson, S. (2009). A key role for floral scent in a wasp-pollination system in *Eucomis* (Hyacinthaceae). *Annals of Botany*, 103, 715-725.
- Sonah, H., Deshmukh, R. K., Sharma, A., Singh, V. P., Gupta, D. K., Gacche, R. N., Rana, J. C., Singh, N. K. & Sharma, T. R. (2011). Genome-wide distribution and organization of microsatellite in plants: an insight into marker development in *Brachypodium*. *PLoS ONE*, 6, 1-9.
- Song, Q., Jia, G., Zhu, Y., Grant, D., Nelson, R., Hwang, E-Y, Hyten, D. & Cregan, P. (2010). Abundance of SSR motifs and development of candidate polymorphic SSR markers (BARCSOYSSR_1.0) in soybean. *Crop Science*, 50, 1950-1960.
- Spitcer, B., Ben Zvi, M., Ovadis, M., Marhevla, E., Barkai, O., Edelbaum, O., Marton, I., Masci, T., Alon, M., Morin, S., Rogachev, I., Aharoni, A. & Vainstein, A. (2007). Reverse genetics of floral scent: application of tobacco rattle virus-based gene silencing in petunia. *Plant Physiology*, 145, 1241-1250.
- Stafne, E. T., Clark, J. R., Weber, C. A., Graham, J. & Lewers, K. S. (2005). Simple sequence repeat (SSR) markers for genetic mapping of raspberry and blackberry. *Journal of American Horticultural Science*, 130, 1-7.
- Teh, S. L. (2011). *Development of a floral expressed sequence tags resource from and characterization of fragrance-related gene transcripts in Vanda Mimi Palmer*. Master dissertation, Universiti Putra Malaysia, Selangor Darul Ehsan, Malaysia.
- Teh, S. L., Chan, W. S., Janna, O. A. & Parameswari, N. (2011). Development of expressed sequence tag resources for *Vanda Mimi Palmer* and data mining for EST-SSR. *Molecular Biology Reports*, 38, 3903-3909.
- Temnykh, S., DeClerck, G., Lukashova, A., Lipovich, L., Cartinhour, S. & McCouch, S. (2001). Computational and experimental analysis of microsatellites in rice (*Oryza sativa* L.): frequency, length variation, transposon associations and genetic marker potential. *Genome Research*, 1441-1452.
- Teo, C. K. H. (1981). *Tropical orchid hybrids*, FEP International Sdn. Bhd., Singapore.
- Teoh, E. S. (1980). *Asian orchids*, Times Books International, Singapore.
- Terp, N., Gobel, C., Brandt, A. & Feussner, I. (2006). Lipoxygenases during *Brassica napus* seed germination. *Phytochemistry*, 67, 2030-2040.
- Tsai, W., Hsiao, Y., Lee, S., Tung, C., Wang, D., Wang, H., Chen, W. & Chen, H. (2006). Expression analysis of the ESTs derived from the flower buds of *Phalaenopsis equestris*. *Plant Science*, 170, 426-432.

- van Moerkercke, A., Galvan-Ampudia, C. S., Verdonk, J. C., Haring, M. A. & Schuurink, R. C. (2012). Regulators of floral fragrance production and their target genes in petunia are not exclusively active in the epidermal cells of petals. *Journal of Experimental Botany*, doi:10.1093/jxb/ers034
- vanTunen, A. J., Eikelboom, W. & Angenent, G. C. (1993). Floral organogenesis in *Tulipa*. *Flowering Newsletter*, 16, 33-37.
- Vainstein, A., Lewinsohn, E., Pichersky, E. & Weiss, D. (2001). Floral fragrance. New inroads into an old commodity. *Plant Physiology*, 127, 1383-1389.
- Varshney, R. K., Chabane, K., Hendre, P. S., Aggarwal, R. K. & Graner, A. (2007). Comparative assessment of EST-SSR, EST-SNP and AFLP markers for evaluation of genetic diversity and conservation of genetic resources using wild, cultivated and elite barleys. *Plant Science*, 173, 638-649.
- Vejaratpipom, R., Channuntapipat, C., Pewnim, T., Ito, K., Iizuka, M. & Minamiura, N. (1999). Detection and serological relationship of cymbidium mosaic potexvirus isolates. *Journal of Bioscience and Bioengineering*, 87, 161-168.
- Verdonk, J. C., Haring, M. A., van Tunen, A. J. & Schuurink, R. C. (2005). *ODORANT1* regulates fragrance biosynthesis in petunia flowers. *The Plant Cell*, 17, 1612-1624.
- Wang, M. L., Gillaspie, A. G., Newman, M. L., Dean, R. E., Pittman, R. N., Morris, J. B. & Pederson, G. A. (2004). Transfer of simple sequence repeat (SSR) markers across the legume family for germplasm characterization and evaluation. *Plant Genetic Resource*, 2, 107-119.
- Yu, K., Park, S. J., Poysa, V. & Gepts, P. (2000). Integration of simple sequence repeat (SSR) markers into a molecular linkage map of common bean (*Phaseolus vulgaris* L.). *The American Genetic Association*, 91, 429-434.
- Yue, G. H., Lam-Chan, L. T. & Hong, Y. (2006). Development of simple sequence repeat (SSR) markers and their use in identification of *Dendrobium* varieties. *Molecular Ecology Resources*, 6, 832-834.
- Zhang, D., Choi, D., Wanamaker, S., Fenton, R., Chin, A., Malatrasi, M., Turuspekov, Y., Walia, H., Akhunov, E., Kianian, P., Otto, C., Deal, K., Echenique, V., Stamova, B., Ross, K., Butler, G., Strader, L., Verhey, S., Johnson, R., Altenbach, S., Kothari, K., Tanaka, C., Shah, M., Laudencia-Chingcuanco, D., Han, P., Miller, R., Crossman, C., Chao, S., Lazo, G., Klueva, N., Gustafson, J., Kianian, S., Dubcovsky, J., Walker-Simmons, P., Gill, K., Dvorak, J., Anderson, O., Sorrells, M., McGuire, P., Qualset, C., Nguyen, H. & Close, T. (2004). Construction and evaluation of cDNA libraries for large-scale expressed sequence tag sequencing in wheat (*Triticum aestivum* L.). *Genetics*, 168, 595-608.