

1 **Metabolic engineering strategies for sesquiterpene production in microorganism**

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28 **Abstract**

29 Sesquiterpenes are a large variety of terpene natural products, widely existing in plants, fungi,
30 marine organisms, insects, and microbes. The value-added sesquiterpenes are extensively used
31 in industries as food, drugs, fragrances and fuels. With increase of the market demands and the
32 price of sesquiterpenes, biosynthesis of sesquiterpenes by microbial fermentation methods
33 from renewable feedstocks acquires increasing attention. The synthetic biology provides robust
34 tools of sesquiterpene production in microorganisms. This review presents a summary of
35 sesquiterpene biosynthesis and metabolic engineering strategies on the host and pathway
36 engineering for sesquiterpene production. The diversity, native producers, and the syntheses of
37 sesquiterpenes are presented. Advances in synthetic biology provide new strategies on creation
38 of the desired hosts for sesquiterpene production. Especially, metabolic engineering strategies
39 for production of sesquiterpenes such as amorphadiene, farnesene, bisabolene, and
40 caryophyllene are emphasized in *Escherichia coli*, *Saccharomyces cerevisiae*, and some other
41 microorganisms. Future perspectives on strain and process improvements for sesquiterpene
42 production are also discussed.

43

44 **Keywords**

45 Sesquiterpene biosynthesis, sesquiterpene synthase, metabolic engineering strategies, MEP
46 pathway, MVA pathway.

47

48 1. Introduction

49 Terpenes are a large and diverse class of organic compounds, with basic C5 isoprene-unit
50 hydrocarbons. According to the number of C5 isoprene-unit, terpenes are classified as hemi-
51 (C5), mono- (C10), sesqui- (C15), di- (C20), sester- (C25), tri- (C30) and tetra- (C40) terpenes
52 (Bohlmann and Croteau, 1999, Chen et al. , 2011). Sesquiterpene, the largest subgroup of
53 terpenes, has several thousands of representatives and more than 300 different skeletons
54 (Breitmaier, 2008, Connolly and Hill, 1991, Sacchettini and Poulter, 1997). Sesquiterpenes
55 mostly consist of hydrocarbons possessing a multitude of different carbon skeletons and some
56 unsaturated bonds, and their structures present acyclic, monocyclic, bicyclic, and tricyclic
57 compounds (Kramer and Abraham, 2011). They present more structures after oxidation or
58 rearrangement based on C15 skeleton. Sesquiterpenes are mostly isolated from a variety of
59 natural sources, including plant, fungi, microorganisms, and insect (Sacchettini and Poulter,
60 1997). In plants and microorganisms, sesquiterpenes are important components of volatile oil,
61 existing in form of alkene, alcohol, ketone, and lactone, emitted respectively for spreading
62 fungal spores, protecting themselves from enemies and communicating with other organisms
63 (Rohlf's and Churchill, 2011).

64 In these natural sources, sesquiterpene synthases are responsible for compound
65 biosynthesis from the common intermediate substrate farnesyl diphosphate (FPP) condensed
66 from isopentenyl diphosphate (IPP) and dimethylallyl pyrophosphate (DMAPP) provided by
67 mevalonate (MVA)-dependent pathway discovered in 1950s and 2-C-methyl-D-erythritol-1-4-
68 phosphate (MEP) pathway disclosed in 1993 (Rohmer, et al. , 1993, Schwender et al. , 1996).
69 The MVA pathway exists in some prokaryotes and almost all eukaryotes including high plants.
70 Many, but not all eubacteria (Rohmer, 1999, Rohmer et al. , 1993, Trutko et al. , 2005), and
71 cyanobacteria (Proteau, 1999) make exclusive use of the MEP pathway. The MVA pathway,
72 located in cytosol, is suggested to be responsible for synthesis of sesquiterpenes (C15),

73 terpenoid aldehydes (C15 and C25) and steroids/triterpenoids (C30). While all MEP-derived
74 isoprenoids are generally considered as typical and exclusive plastidial hemi- (C5), mono-
75 (C10), diterpenes (C20) and carotenoids (C40) (Bouvier et al. , 2005, Opitz et al. , 2014,
76 Pazouki and Niinemets, 2016). Although the MVA pathway and the MEP pathway operate in
77 two different subcellular compartments (cytosol and plastid, respectively), metabolite
78 exchanges or cross-regulation known as crosstalk occur between them (Hemmerlin et al. ,
79 2003).

80 Because many sesquiterpenes have complex molecular structures, inherent bioactivity,
81 and aroma, they could be applied as functional precursors of fragrances, medicines, and
82 fuels (Enserink, 2005, Little et al. , 1950, Nour et al. , 2009, Peralta-Yahya et al. , 2011).
83 Although sesquiterpenes and their derivatives display a broad range of application from
84 medicine to fuels, their production is usually at low levels in native producers (Nour et al. ,
85 2009). The small quantities and poor purities limit the scale-up of sesquiterpene production
86 through natural extraction or chemical synthesis. Given the great market demands and
87 limitations of high production of sesquiterpenes, metabolic engineering strategies are
88 attempted to produce them, such as amorphadiene (Redding-Johanson et al. , 2011, Ro et al. ,
89 2006), farnesene (Meadows et al. , 2016, Su et al. , 2015, Yang et al. , 2016c), bisabolene
90 (Kirby et al. , 2014b), and caryophyllene (Yang et al. , 2016a). Synthetic biology provides an
91 alternative approach to produce plant natural product sesquiterpenes in sufficient quantities
92 using microbial fermentation (Ro et al. , 2006), which is enabled to be an economically feasible
93 choice by recent advances.

94 Even though there have been examples of sesquiterpene biosynthesis (Block et al. , 2019,
95 Kramer and Abraham, 2011) and analytical techniques for them (Merfort, 2002), it is
96 meaningful to summarize and update the advances in sesquiterpene biosynthesis and metabolic
97 engineering strategies. Here, we review the main aspects regarding the applications of

98 sesquiterpenes, natural sesquiterpene biosynthesis and metabolic engineering strategies on
99 sesquiterpene production in engineered microorganisms. First, we briefly characterize
100 diversities and applications of sesquiterpenes. Next, we systematically discuss the natural
101 producers of the compounds and terpene synthases (TPSs). Then, we summarize the strategies
102 on metabolic regulation of isoprenoid pathways as well as other relating engineering networks
103 for increasing sesquiterpene production. Finally, we illustrate the great potentials and
104 challenges of microbial production of sesquiterpenes using metabolic engineering strategies.

105 **2. Sesquiterpenes are widely-existing biosynthetic compounds**

106 The spread applications of sesquiterpenes are systematically summarized here shown in
107 Figure 1. Many sesquiterpenes have been extracted from plants by mankind for different
108 purposes - as fragrances and flavors in cosmetic or food industries, pharmaceutical agents,
109 insecticides and fuels (Agger et al. , 2008). For the natural fragrance of sesquiterpenes and their
110 derivations, they could be used as blends in cosmetic.

111 Various plants are popular traditional Chinese medicines to prevent or treat different
112 diseases for centuries. One of the effective constituents of the pharmacopoeia is sesquiterpene
113 derivatives, and the bioactivities make them to exhibit useful medicinal properties.
114 Sesquiterpene lactones make a huge group of bio-active constituents isolated from plant
115 families. They have been reported for various potential effects of anti-microbial activities,
116 antioxidant and anticancer potentials, served as antifeedants (Knight, 1995, Mori and
117 Matsushima, 1995, Mullin et al. , 1991), antimigraine, anti-inflammatory (Pfaffenrath et al. ,
118 2002, Tassorelli et al. , 2005), antitumor, and antiulcer (Cho et al. , 2000) for treatment of
119 stomach-ache and skin infection (Heinrich et al. , 1998). From sesquiterpenes and their oxides,
120 considerable interests are gained, such as well-known amorphadiene-derived artemisinin for
121 antimalarial activity (Ecksteinludwig et al. , 2003), xanthatin for antitumor activity (Nibret et
122 al. , 2011), parthenolide for curing migraine (Kwok et al. , 2001), and thapsigargin for treating

123 prostate cancer (Drew et al. , 2013). Furanoeudesma-1,3-diene and curzarene are responsible
124 for the analgesic effects of myrrh by interacting with brain opioid receptors (Massoud et al. ,
125 2010). β -2-Himachalen-6-ol, a novel sesquiterpene alcohol unique to Lebanese wild carrot,
126 was demonstrated with potent anti-cancer activity (Taleb et al. , 2016). *Trans*-caryophyllene is
127 a bicyclic sesquiterpene produced by plants in response to herbivore damage (Köllner et al. ,
128 2008). Valerenadiene and valerenic acid have been suggested as the active ingredients
129 responsible for sedative effect in *Valeriana officinalis* (*V. officinalis*) (Pyle et al. , 2012). Some
130 plants synthesize mixture of sesquiterpenes which are believed to be ecologically more
131 effective than single compound, such as the case of attraction of pollinators or parasites of
132 insects (Tholl, 2006). Additionally, some sesquiterpenes are also ingredients of pheromones
133 and juvenile hormones or their precursors (Izquierdo-Bueno et al. , 2018, Lamers, 2003).

134 Sesquiterpenes exist in many natural foods, such as fruits, drinks, and oil. They are
135 comprehensively investigated in essential oil producing plants, such as peppermint, conifers,
136 and citrus fruits (Croteau, 2001, Croteau et al. , 2005, Voo et al. , 2012). Sesquiterpenes
137 accumulate in specialized tissues such as glandular trichomes, oil ducts or secretory cavities
138 (Lange and Turner, 2013). Over 60 sesquiterpenes have been identified in whole grapes or
139 skins (Petronilho et al. , 2014); α -Farnesene was found on natural coating of apples (FE and
140 KE, 1966); (+)-valencene is an aroma compound of citrus fruits, used in flavor foods and drinks
141 (Frohwitter et al. , 2014). Through BLAST analysis, 19 TPSs and 13 sesquiterpenes were
142 identified in unripe fruits of black pepper (*Piper nigrum*), including β -caryophyllene, δ -
143 elemene, α -copaene, cubebene, α -humulene, δ -cadinol, γ -cadinene and δ -cadinene (Jin et al. ,
144 2018). Insects are repelled by subjecting clove oil and patchouli oil, and the functional
145 ingredients are sesquiterpenes and their oxides: seychellene, α -guaiene, α -bulnesene, β - and
146 iso-caryophyllene, β -caryophyllene oxide and ketone, α -humulene, β -patchoulene, α -
147 patchoulene, β -elemene, (+)-longipinene, (-)-isolongifolene, (+)-longifolene, and so on

148 (Zhang et al. , 2011). Therefore, sesquiterpenes may have potential interests as preservative in
149 food. Many sesquiterpenes, β -elemene, γ -elemene, α -caryophyllene, germacrene D and δ -
150 cadinene, exist in fruit of *Zanthoxylum schinifolium*, a kind of flavourful spice and Chinese
151 medicine (Wang et al. , 2011b). α -Humulene and α -caryophyllene are contained in the essential
152 oil of *Humulus lupulus*, an inevitable ingredient for beer brewery (Connolly and Hill, 1991).
153 Interestingly, potent odorants in wine are from some mixed terpenoids, of which the formation
154 mechanism encompasses enzymatic and non-enzymatic steps (Schwab and Wust, 2015).

155 In addition, due to high energy density, comparable cold properties and similar carbon
156 length with cetane, sesquiterpenes reveal great interests as next-generation jet fuel substitutes
157 (Harrison and Harvey, 2017). Especially, the ring and branching portion of sesquiterpenes
158 could increase the gravimetric density and the volumetric energy density of fuel. Some
159 sesquiterpenes have been proposed as jet fuel precursors due to their low hygroscopicity and
160 high energy density. Yoshikuni and coworkers introduced TPS mutants into *E. coli* for terpene
161 production, obtaining eight different terpenes (α -farnesene, sabinene, γ -humulene, sibirene,
162 longifolene, α -longipinene and β -bisabolene), most of which have potential as biofuels
163 (Yoshikuni et al. , 2006). The biosynthesized bisabolene was chemically hydrogenated to
164 generate bisabolane, which was identified as an alternative to D2 diesel by comparing its fuel
165 properties with those of diesel fuels (Peralta-Yahya et al. , 2011). Sesquiterpene thujopsene, α -
166 cedrene and β -cedrene in cedarwood oil, were hydrogenated to generate fuel blends with 12%
167 higher volumetric net heat of combustion than conventional jet fuel (Harrison and Harvey,
168 2017). Sesquiterpene epi-isozizaene, pentalenene and α -isocomene are even proven to have
169 excellent fuel properties by heat of combustion calculation (Liu et al. , 2018).

170 **Figure 1**

171 **3. Sesquiterpene biosynthesis**

172 **3.1 Sesquiterpene producers**

173 In nature, sesquiterpenes widely exist in plants, fungi, marine organisms, and insects
174 (Figure 2) (Connolly and Hill, 1991, Finefield et al. , 2012). Most of sesquiterpenes are
175 essential for the life of their producers, such as steroid hormones in mammals, carotenoids and
176 chlorophylls in plants, and ubiquinone or menaquinone in bacteria (Sacchettini and Poulter,
177 1997, Thomas, 1995). In plants, sesquiterpenes and monoterpenes are important constituents
178 of essential oils. As summarized in Figure 2, they are components of volatile oil in form of
179 alcohol, ketone, lactone or glycoside in Asteraceae, Umbelliferae, Euphorbiaceae, Wedaceae,
180 Leguminous, Cucurbitaceae and Ranunculaceae (Connolly and Hill, 1991). Volatile oil from
181 plants is usually consistent of sesquiterpenes: in oil of *Xanthium strumarium*, germacrene D,
182 guaia-4, -6-diene, β -elemene, β -caryophyllene and β -copaene were detected (Li et al. , 2016a).
183 Sesquiterpenes are modulated by enzymes or other enzymes being induced by environmental
184 stress factors. For example, UV-radiation (Back et al. , 1998), infection (Poole and Llewellyn,
185 2005) and herbivore attack (Yuan et al. , 2008) have been identified to enhance volatile
186 sesquiterpene production. It was reported that H₂O₂ could induce sesquiterpene production by
187 increasing expression of gene *AsTPS10*, *AsTPS16* and *AsTPS19* responsible for sesquiterpene
188 biosynthesis in wounded *Aquilaria sinensis* (Lv et al. , 2019). *Solidago canadensis* majorly
189 produces germacrene D, accounting for more than 90% of the total compounds (Niwa et al. ,
190 2008, Schmidt et al. , 1998, Weyerstahl et al. , 1993). Asteraceae family plant *Parthenium*
191 *hysterophorus* produces guaia-4,6-diene, whose chemical structure was previously confirmed
192 by nuclear magnetic resonance (NMR) studies (Bohlmann et al. , 1977). Sesquiterpene 7-
193 epizingiberene produced and stored in glandular trichomes is toxic and repellent properties
194 (Bleeker et al. , 2012). Through gas-chromatographic-mass spectrometric (GC-MS) analysis
195 of dynamic gas from the native scent of *Solanum tuberosum* L. leaves, the following
196 sesquiterpenes have been identified: *trans*- and *cis*-caryophyllenes, α -ylangene, α -copaene, β -
197 bourbonene, β -elemene, Z- β -farnesene, aromadendrene, α -humulene, α -elemene, β -bisabolene,

198 γ -cadinene and δ -cadinene (Khalilov et al. , 1999). In flowers of *Cananga odorata* var.
199 *fruticosa*, bulk of the 49 VOCs were identified as sesquiterpenes: β -caryophyllene, farnesene,
200 γ -muurolene, humulene, β -germacrene D, cedrene, δ -cadinene, α -patchoulene, α -bergamotene,
201 β -ylangene, β -copaene, and β -cubebene (Jin et al. , 2015).

202 Fungi is another rich resource of terpenes, mostly existing in the format of VOCs (Figure
203 2). Two producers *Ascomycota* and *Basidiomycota* usually produce sesquiterpenes in late
204 growth phase. Many species of the phylum *Ascomycota* have been reported to produce volatile
205 sesquiterpenes (Kramer and Abraham, 2011). Caryophyllene was emitted from *Phialophora*
206 *fastigata*, and α -curcumene was biosynthesized by *Penicillium commune* and *Paecilomyces*
207 *variotii* (Sunesson et al. , 1995). Six TPSs and fifteen sesquiterpenes were respectively
208 characterized and identified in basidiomycete *Coprinus cinereus* (Agger et al. , 2009). Genus
209 *Penicillium*, belonging to phylum *Ascomycota*, is a kind of rich sesquiterpene producer:
210 germacrene A and germacrene B were respectively produced by *P. cyclopium* and *P. expansum*,
211 and β -elemene and β -guaiene were synthesized by *P. clavigerum* (Larsen, 1998). *P. roqueforti*
212 could emit multiply sesquiterpenes β -patchoulene, β -elemene, β -elemene-isomer, diepi-a-
213 cedrene, β -gurjunene, β -patchoulene-isomer, aristolochene, valencene, α -selinene, β -
214 himachalene, α -chamigrene, β -bisabolene, and α -panasinsene (Jeleń, 2002). The fungal
215 phylum Basidiomycota produces multiply sesquiterpenes via transforming and rearranging
216 humulene (Abraham, 2001).

217 Many terpenes from plants and fungi have been described, but relatively fewer microbial
218 sesquiterpenes and TPSs have been functionally summarized. As we summarized here,
219 prokaryote producers of sesquiterpene are mainly identified belonging to Actinomyces, marine
220 bacteria and Cyanobacteria (Figure 2). *Streptomyces* is a rich source of bioactive secondary
221 metabolites, such as antiparasitic avermectins and sesquiterpene-derived antibiotic
222 pentalenolactone (Chou et al. , 2010, Omura et al. , 2001, Quaderer et al. , 2006).

223 Sesquiterpenoid antibiotic pentalenolactone is a commonly occurring metabolite, isolated from
224 more than 30 species of *Streptomyces* (Koe et al. , 1957, Takahashi et al. , 1983).
225 Bioinformatics analysis of *Streptomyces* and *Kitasatospora* genomes identified 20 terpenes,
226 including sesquiterpene β -elemene, α -muurolene, cadinadiene, epi-isozizaene, germacrene A
227 and pentalenene (Yamada et al. , 2015a, Yamada et al. , 2015b). Many marine bacteria are
228 found to produce many sesquiterpenes with great medical values. Four sesquiterpenes—
229 euplotin A, euplotin B, euplotin C and their biogenetic precursor preuplotin—have been
230 isolated from a eukaryotic unicellular marine ciliate *Euplotes crassus* (Dini et al. , 1993).
231 Sesquiterpenes exclusively produced by marine bacteria ciliate protozoon *E. crassus* have the
232 bioactivity of inhibiting growth and killing protozoa *L. major* and *L. infantum* in vitro (Savoia
233 et al. , 2004).

234 In addition, insects were reported to synthesize sesquiterpenes (Finefield et al., 2012,
235 Naya et al. , 1978). Some investigation suggests that insects possess a specific set of C-C
236 cyclases for sesquiterpenoids. Volatile oil from the secretion of a scale insect *Ceroplastes*
237 *ceriferus*, is consisting of numbers of sesquiterpenes: germacrene-D, β -bourbonene, α -copaene,
238 β -bourbonene and β -elemene (Naya et al. , 1978). The presence of some cedrene-type
239 sesquiterpenoids was reported in scale insect *Laeifer lacea* Kerr (Singh et al. , 1974).

240 **Figure 2**

241 **3.2 Sesquiterpene synthases**

242 In nature, terpene synthases and their carbocationic reaction mechanisms are responsible
243 for huge varieties of terpene structures and biosynthesis, and they have garnered attentions of
244 chemists and biologists for many years. Starting from farnesyl diphosphate (FPP),
245 sesquiterpene skeleton is cyclized by different TPSs (Benedict et al. , 2001, Cane, 1990). From
246 analysis of TPSs, DDXXD motifs were found to serve as a crucial site for divalent action and
247 substrate binding (Marrero et al. , 1992, Prosser et al. , 2004). Many TPSs, such as farnesene

248 synthase from *C. junos*, δ -selinene synthase and λ -humulene synthase, were discovered through
249 searching the motifs (Little and Croteau, 2002, Maruyama et al. , 2001).

250 Large amount of natural TPSs are extracted from plants. A TPS forming carbocyclic
251 skeleton of δ -guaiene was isolated from *Aquilaria* cells (Kumeta and Ito, 2010). α -Farnesene
252 synthase cDNA from peel tissue of apple fruit was cloned and characterized in *E. coli* (Pechous
253 and Whitaker, 2004). Germacrene A synthase from *Lactuca sativa L.* was expressed in
254 engineered *E. coli*, and β -elemene was extracted by thermally rearrangement (Bennett et al. ,
255 2002, Kraker et al. , 1998). The caryophyllene synthase was characterized from *Artemisia*
256 *annua* (Cai et al. , 2002) and *Mikania micrantha* (Wang et al. , 2009). Multiple TPSs are often
257 expressed in different organs of one organism, and several sesquiterpenes are synthesized
258 through one TPS catalysis. Therefore, several sesquiterpenes are often produced by one
259 organism or one TPS catalysis of substrate FPP. For example, β -humulene synthase and δ -
260 selinene synthase are expressed in *Abies grandis*, each catalyzing at least 52 and 36
261 sesquiterpenes, respectively (Steele et al. , 1998). Chamomile, with great pharmaceutical
262 values and economic essential oil, was excavated five TPSs, four of which MrTPS1, MrTPS3,
263 MrTPS4 and MrTPS5 were exclusively expressed in above-ground organs and respectively
264 catalyzed (-)-(E)- β -caryophyllene, (+)-germacrene A, (E)- β -ocimene and (-)-germacrene D.
265 Enzyme MrTPS2, mainly expressed in roots, catalyzed several Asteraceae-specific
266 sesquiterpenes: (-)-(E)- β -caryophyllene, α -humulene, silphinene, modeph-2-ene, β -isocomene
267 and the major product α -isocomene (Irmisch et al. , 2012). Based on transcriptome sequencing
268 of *Santalum album*, five TPSs were characterized, including two sesquisabinene synthases
269 (SaSQS1, SaSQS2), bisabolene synthase (SaBS), santalene synthase (SaSS) and farnesyl
270 diphosphate synthase (SaFDS) (Srivastava et al. , 2015). In grand fir, δ -selinene synthase and
271 λ -humulene synthase each enzyme synthesizes three major products respectively, and a third
272 TPS catalyze to form only (E)- α -bisabolene (Bohlmann et al. , 1998, Steele et al. , 1998). Six

273 TPSs from *C. cinereus* were described: Cop1 and Cop2 each synthesizes germacrene A as their
274 major product; Cop3 was identified as an α -muurolene synthase but producing β -elemene, δ -
275 muurolene, germacrene D, and δ -cadinene; Cop4 synthesizes δ -cadinene as its major product;
276 Cop5 has not been characterized any compounds; Cop6 catalyzes specific α -cuprenene (Agger
277 et al. , 2009). Two TPSs were identified to synthesize a-farnesene and germacrene D from
278 kiwifruit flowers (Nieuwenhuizen et al. , 2009). In root of *V. officinalis*, two TPSs VoTPS1
279 and VoTPS2 were identified to produce germacrene C/germacrene D and valerenadiene,
280 respectively (Pyle et al. , 2012). In flowers of grapevine *Vitis vinifera* L., 69 putative TPSs were
281 discovered through probe cross-hybridizing, and numbers of sesquiterpenes were identified in
282 heterologous yeast and the plant flowers (Smit et al. , 2019).

283 Recent years, the availability of genome sequences led to the discoveries of some TPSs
284 from actinomycetes and cyanobacteria (Agger et al. , 2008, Cane and Watt, 2003, Giglio et al. ,
285 2008, Gust et al. , 2003, Komatsu et al. , 2008, Lin et al. , 2006a, Zhao et al. , 2008). Based on
286 bioinformatics analysis, a biosynthetic protein (AncCL1) was reconstructed and identified to
287 catalyze major product epi-zizaene and minor product epi-isozizaene (Guzzetti et al. , 2016).
288 In *Streptomyces coelicolor*, epi-isozizaene synthase (EIZS) catalyzes cyclization of FPP to a
289 novel tricyclic epi-isozizaene (Lin et al. , 2006b). Pentalenene synthase (PentS) from
290 *Streptomyces* UC5319 catalyzes to pentalenene (Cane et al. , 1994), and a mixture of
291 pentalenene and caryophyllene is synthesized by site-directed mutagenesis PentS N219D
292 (Seemann et al. , 2002). Similarly, the types or the product distribution of sesquiterpenes could
293 be changed when mutating TPSs, such as MrTPS2 (Irmisch et al. , 2012), bisabolene synthase
294 (*BIS*) (Kirby et al. , 2014a) and amorphadiene synthase (*ADS*) (Li et al. , 2016b).

295 **Figure 3**

296 **4. Metabolic engineering strategies on sesquiterpene production in microorganisms**

297 To explore the potential interest of sesquiterpenes, fermentation-based production process
298 is used to produce diverse sesquiterpenes through engineering metabolic pathways to
299 efficiently drive conversion of the renewable resources. Some industrial model
300 microorganisms are metabolically engineered from isoprenoid pathways, TPSs and other
301 relative metabolic networks for sesquiterpene production. Herein, we describe the general
302 strategies on host and pathway engineering for sesquiterpene production in engineered
303 microorganisms.

304 4.1. Metabolic engineering strategies on sesquiterpene production in *E. coli*

305 4.1.1. Metabolic regulation on MEP pathway for sesquiterpene production in *E. coli*

306 *E. coli* employing the MEP pathway to accumulate FPP is a widely used host for
307 sesquiterpene production. Metabolic engineering and strategies for sesquiterpene production
308 in *E. coli* are summarized in Table 1 and Figure 4. Via the *E. coli* native MEP pathway,
309 sesquiterpene production is generally considerable low when only over-expressing TPSs. For
310 example, *E. coli* BL21 (DE3) harboring only gene farnesene synthase produced 0.325 mg/L α -
311 farnesene (Zhu et al. , 2014). When codon-optimized gene *coEIZS*, *coPentS* and *coMrTPS2*
312 were simply over-expressed in *E. coli* DH1, the titers of tricycle sesquiterpene epi-isozizaene,
313 pentalenene and α -isocomene were 0.54 mg/L, 0.19 mg/L and 0.01 mg/L, respectively (Liu et
314 al. , 2018). In *E. coli* BL21 (DE3), over-expressing wildtype *valerenadiene synthase* (*wvds*)
315 gene rendered 12 μ g/L valerenadiene (Nybo et al. , 2017).

316 For metabolic regulation on the MEP pathway, over-expression of rate-limiting genes and
317 TPSs is the most popular strategy to increase sesquiterpene production. Upon co-expressing
318 gene *Dxs*, *isopentenyl-diphosphate isomerase* (*Idi*) and *ispA* in SOE4 operon, amorphadiene
319 titer increased 3.6-fold comparing to that achieved by only *ADS* gene expression strain (Martin
320 et al. , 2003). Medium optimization and over-expression of gene *Dxs* from *Streptomyces*
321 *avermitilis*, *Idi* from *Bacillus subtilis*, *ispDF* and *ADS* increased amorphadiene titer to 331.70

322 mg/L, 15.5-fold of the initial titer in *E. coli* MG1655 (Wang et al. , 2015). The aim of over-
323 expressing and deleting genes is to enrich the vital MEP pathway precursors. To enrich
324 precursor phosphoenolpyruvate and pyruvate, carbohydrate phosphotransferase system (PTS)
325 genes were deleted, resulting in amorphaadiene titer increased to 182 mg/L in OPT2 medium
326 (Zhang et al. , 2013, Zhang et al. , 2015). To enrich the vital precursor DXP, a novel route of
327 pentose phosphate to it annotated as putative xylose reductase gene *yajO* and *ribB* mutant was
328 screened. Through fusing *ribB* mutant to *Dxr* gene and over-expressing *ispA*, *ispDF* and *Idi*
329 gene, bisabolene titer increased above 4-fold, about 8.8 mg/g DCW (dry cell weight) (Kirby et
330 al. , 2015). Engineering the MEP pathway by cross-lapping in vitro assembly (CLIVA) method
331 and expressing iron sulfur (Fe-S) cluster operon increased amorphaadiene titer to ~300 mg/L
332 (Zou et al. , 2013). Additionally, to increase sesquiterpene production, efflux transporter
333 engineering was carried out: Over-expressing the screened outer membrane protein *tolC*,
334 together with ABC family transporters (*macAB*) or MFS family transporters (*emrAB* or
335 *emrKY*) increased amorphaadiene titer by more than 3-fold, up to 180 mg/L (Zhang et al. , 2016);
336 Over-expressing two copies of *TolC* and *AcrB* improved amorphaadiene yield 118% with 404.83
337 mg/L (Wang et al. , 2013). For valerenadiene titer, besides over-expressing gene *Dxs*, *Idi*, FPP
338 synthase (*FFPS*) and *cvds*, supplying glycerol into fermentation medium increased it to 11.0
339 mg/L (Nybo et al. , 2017).

340 4.1.2. Metabolic regulation on MVA pathway in *E. coli*

341 The low sesquiterpene production may be resulted from the insufficient FPP
342 supplementary via native MEP pathway which is tightly controlled by endogenous regulation
343 (Phulara et al. , 2016). The heterogenous MVA pathway was introduced to *E. coli* for
344 dramatically improving flux to FPP and terpene production (Martin, Pitera, 2003). In *E. coli*
345 DYM1, the synthetic *ADS* gene from *A. annua* and MVA pathway from *S. cerevisiae* were
346 expressed, achieving approximately 112.2 mg/L of amorphaadiene (see Table 1) from LB + 0.8%

347 glycerol culture (Martin, Pitera, 2003). Afterwards, employing a two-phase partitioning
348 bioreactor with dodecane led this engineered strain to produce 0.5 g/L amorphadiene (Newman
349 et al. , 2006). Although introduction of the MVA pathway dramatically increased
350 amorphadiene production, bottlenecks limiting the yield still exist. To investigate them,
351 balancing the heterologous MVA pathway flux was proposed. All the actions of supplementing
352 MVA into medium, increasing copies of mevalonate kinase gene (*mvk*) or plasmid with *ADS*
353 gene and global sensitivity analysis with an equation model for the MVA pathway using kinetic
354 parameters identified the rate-limiting genes *HMGR*, *mvk* and *ADS* (Anthony et al. , 2009,
355 Pitera et al. , 2007, Weaver et al. , 2015). Gene titration studies and metabolomics using liquid
356 chromatography–mass spectrometry (LC-MS) excavated a bottleneck - insufficient *HMGR*
357 leading to HMG-CoA accumulation, being toxic to cells by inhibiting fatty acid biosynthesis
358 and generalizing membrane stress (Kizer et al. , 2008, Pitera et al. , 2007). Sequentially, three
359 strategies were carried out: upregulating *tHMGR* gene, over-expressing *HMGS* and *HMGR*
360 gene from *Staphylococcus aureus* (Tsuruta et al. , 2009), and cofactor engineering using
361 NADH-dependent *HMGR* variants from *Delftia acidovorans* (Ma et al. , 2011). Then, a
362 commercially relevant amorphadiene titer of 27.4 g/L was achieved through a superior fed-
363 batch fermentation process of restricted glucose and nitrogen (Tsuruta et al. , 2009). By
364 supplying more ATP, polyphosphate kinase (PPK)-catalyzed synthesis increased
365 amorphadiene titer 2.5-fold (Shimane et al. , 2012). To characterize strain variation in
366 engineered *E. coli* for increasing the productivity, a multi-omics-based workflow was
367 developed (Brunk et al. , 2016). Especially, targeted proteomics helped to identify *mvk* and
368 *phosphomevalonate kinase (pmk)* as potential bottleneck genes in amorphadiene production.
369 Based on this, codon-optimizing *mvk* and *pmk* gene, and replacing P_{lac} with a stronger P_{trc} were
370 conducted, improving amorphadiene titer 3-fold (>500 mg/L) (Redding-Johanson, Batth, 2011).
371 Assisted by metabolomics and targeted proteomics, metabolic pathway optimization with RBS

372 variants and combinatorial gene assembly reduced toxic intermediate accumulation and
373 increased amorphanthene production about 5-fold (Nowroozi et al. , 2014). Accumulation of
374 toxic intermediates (HMG-CoA, IPP/DMAPP and FPP) of the MVA pathway resulted in cell
375 growth issues. Through toxicity stress and whole-genome transcriptional analysis identifying
376 FPP-responding promoter P_{rstA} and P_{gadE} , dynamic control of the MVA pathway was
377 established, and it doubled amorphanthene titer to 1.4 g/L (Dahl et al. , 2013). Afterwards, this
378 dynamic control strategy was used to produce other sesquiterpenes: dynamic control of the
379 MVA pathway expression and comparative proteome analysis made engineered *E. coli*
380 BW25113 to produce 722.46 mg/L zeaxanthin (Shen et al. , 2016); Together promoter
381 engineering with this dynamic control or inducible control of the MVA pathway, 727.9 mg/L
382 epi-isozizaene, 780.3 mg/L pentalenene and 77.5 mg/L α -isocomene were produced when gene
383 *coEIZS*, *coPentS* and *coMrTPS2* were over-expressed in *E. coli* DH1, respectively (Liu et al. ,
384 2018).

385 α -Farnesene and β -farnesene are isomers, differing by the location of one double-bond.
386 The commercially available farnesene used as perfume and fuel is a mixture of the isomers.
387 Expressing a codon-optimized farnesene synthase gene made *E. coli* to produce 1.2 mg/L α -
388 farnesene. Augmenting the metabolic flux for FPP synthesis through expressing the MVA
389 pathway and over-expressing *Dxs*, *Idi*, and *ispA* gene conferred a 48-fold improvement.
390 Together with protein fusion of FPPS and farnesene synthase, 380.0 mg/L of α -farnesene was
391 produced (Wang et al. , 2011a). Through engineering the heterologous MVA pathway, 8.74
392 g/L β -farnesene was produced in a lab bioreactor (You et al. , 2017). Although up-regulating
393 bottleneck genes and down-regulating competitive pathway improved sesquiterpene
394 production, the ratios of enzymes in the reconstructed pathway are hardly precisely evaluated.
395 Therefore, targeted engineering analysis was employed: the MVA pathway was reconstructed
396 *in vitro* to produce farnesene using purified enzymes, and the molar ratio of

397 AtoB:ERG13:tHMG1:ERG12:ERG8:MVD1:Idi:ispA:AFS was slightly adjusted to
398 1:10:2:5:5:2:5:2:2. Based on this information, a rationally designed strain was constructed *in*
399 *vivo* for a high titer of farnesene, achieving 1.1 g/L after 96 h post induction in *E. coli*
400 BL21(DE3) (Zhu, Zhong, 2014). The *in vitro* biosynthesis of sesquiterpenes starting from
401 acetic acid using the MVA pathway was also investigated: a ten-step cascade plus added
402 cofactor regeneration systems led conversion of farnesyl pyrophosphate >40% and conversion
403 of sesquiterpenes >90% (Dirkmann et al. , 2018).

404 Bisabolene was identified as a biosynthetic precursor of a novel alternative to D2 diesel
405 fuel. *E. coli* expressing the MVA pathway from *S. cerevisiae* and plant-derived bisabolene
406 synthase produced 388 mg/L bisabolene. Codon-optimization of *tHMGR*, *HMGS*, *mvk* and *pmk*
407 genes, and insertion of a P_{trc} in front of *mvk* gene increased bisabolene titer to 912 mg/L
408 (Peralta-Yahya et al. , 2011). Later, this *E. coli* variant was re-engineered to produce 363 µg/L
409 cedrol under shaking flask culture conditions for 72 h (Luo et al. , 2019). To use the industrial
410 biomass pretreated by ionic liquid (IL) for biofuel production, *E. coli* containing plasmid with
411 bisabolene pathway genes was engineered to be resist tolerant to the IL with an inner membrane
412 transporter regulated by an IL-inducible repressor (Ruegg et al. , 2014). Sequentially, principal
413 component analysis of proteomics (PCAP) was proposed to increase bisabolene titer to 1,150
414 mg/L in strain CB with an extra copy of *BIS* (Alonso-Gutierrez et al. , 2015). These above
415 MVA pathway genes and sucrose utilization genes were also integrated into chromosome to
416 address issues associated with *E. coli* as an industrial platform for isoprenoids and growth
417 ability on sucrose. Although a CRISPR-Cas9 system for replacing promoter sequences
418 increased the bisabolene titer 5-fold, low titer of ~435 µg/L was finally resulted probably due
419 to lower enzymes levels from chromosomally integrated pathway than the strains using
420 plasmids to express pathway proteins (Alonso-Gutierrez et al. , 2018). Similar with the
421 dynamic control, quorum sensing (QS) system controlling gene expression in response to

422 population density, was applied to establish an inducer-free system of bisabolene production.
423 The LuxI/R system from *Vibrio fischeri* was employed to construct sensor and response
424 plasmids and variants, of which a QS chromosome-integrated strain produced 1.1 g/L
425 bisabolene without inducers added, increasing 44% comparing to the previous inducible system
426 (Kim et al. , 2017). In addition, simulation including of molecular simulation to study the
427 membrane permeability of terpenoids (Vermaas et al. , 2018), CFD and kinetic-based modeling
428 to optimize the sparger design of a large-scale photobioreactor (Ali et al. , 2019) were also
429 employed to improve bisabolene production.

430 β -Caryophyllene is a bio-based product with economic potentials and a promising
431 precursor candidate for petroleum-based products. Glucose-6-phosphate dehydrogenase
432 (*G6PH*) for increasing NADPH, geranyl diphosphate synthase (*GPPS2*) and β -caryophyllene
433 synthase (*QHS1*) genes, were co-overexpressed with the MVA pathway in engineered *E. coli*
434 YJM59, producing 1,520 mg/L β -caryophyllene in fed-batch fermentation (Yang et al. , 2016a).
435 To convert acetic acid to β -caryophyllene, genes *QHS1*, *GPPS2*, acetyl-CoA synthase (*ACS*)
436 and acetoacetyl-CoA synthase (*AACS*) were co-expressed along with the heterologous MVA
437 pathway in *E. coli* YJM67 and produced up to 1.05 g/L of β -caryophyllene accumulated after
438 72 h (Yang and Nie, 2016). By expressing the MVA pathway and gene *QHS1* in *E. coli*, 406
439 mg/L terpene mix enriched in 100 mg/L caryophyllene was produced (Wu et al. , 2018).

440 For α -humulene production, the MVA pathway genes from *Streptomyces* (from *HMGS* to
441 *Idi2*) and yeast (*Idi1*), and rat gene acetoacetate-CoA ligase (*Aacl*) and α -humulene synthase
442 of shampoo ginger were co-expressed, and the resultant *E. coli* strain produced 958 mg/L α -
443 humulene when supplemented with a lithium acetoacetate (LAA), a 13.6-fold increase
444 compared with strain expressing only humulene synthase (Harada et al. , 2009). These resulted
445 from that exogenous LAA is efficiently converted into acetoacetyl-CoA by mammalian *Aacl*
446 and used as a dominant precursor for isoprenoid biosynthesis. Strategy of a probable

447 mechanism of adaptive evolution was used to engineer *tHMGR* and terpene synthases (λ -
448 humulene synthase, bisabolene synthase, sibirene synthase, longifolene synthase, and a-
449 longipinene synthase) in *E. coli*, and thereby eliminate the bottleneck from inefficient enzymes,
450 increasing the productivity by ~1,000 fold (Yoshikuni et al. , 2008). Protoilludene was
451 successfully produced by overexpressing a hybrid exogenous MVA pathway, endogenous *ispA*,
452 and protoilludene synthase of *Omphalotus olearius*, and was improved to 1199 mg/L by
453 altering promoters and copy numbers (Yang et al. , 2016b). Additionally, 62.0 mg/L
454 valerenadiene was produced via expressing the MVA pathway and *cvds* gene respectively
455 promoted by P_{trc} and P_{T7} (Nybo et al. , 2017). Co-expression of condon-optimized longifolene
456 synthase from *Picea abies*, *FPPS*, and the MVA pathway genes in *E. coli* BL21 (DE3) rendered
457 382 mg/L longifolene in a 5-L bioreactor (Cao et al. , 2019).

458 **Figure 4**

459 **Table 1**

460 4.2. Metabolic engineering strategies for sesquiterpene production in *S. cerevisiae*

461 *S. cerevisiae* harboring the MVA pathway is one of the most favorite microorganisms for
462 industrial production of isoprenoids. The low productivity limits commodity scale production
463 of sesquiterpene. The metabolic engineering strategies, synthetic biology technologies and
464 advanced tools (summarized in Figure 5 and Table 2) have exhibited potential to convert *S.*
465 *cerevisiae* to a high sesquiterpene-producing host.

466 In *S. cerevisiae*, up-regulating the upstream MVA pathway for increasing a precursor FPP
467 synthesis is the most common strategy to increase sesquiterpene production. Over-expressing
468 *tHMGR* gene increased amorphadiene titer approximately 5-fold, and down-regulating *ERG9*
469 encoding squalene synthase using a methionine repressible promoter (P_{MET3}) increased it an
470 additional 2-fold by increasing FPP pool (Paradise et al. , 2008, Ro et al. , 2006). In *S.*
471 *cerevisiae*, a global transcription factor *UPC2* was mutated to regulate biosynthesis of sterols,

472 an additional copy of *tHMGR* was integrated into chromosome, and *ERG20* encoding FPP
473 synthase was over-expressed, generating FPP-biosynthesis platform EPY224 which produced
474 153 mg/L amorphadiene (Ro et al. , 2006). Over-expressing heterologous ADS, replacing
475 promoter by P_{MET3} for repressing *ERG9* gene and enzyme fusion made amorphadiene titer to
476 25.02 mg/L in *S. cerevisiae* MTCC 3157 (Baadhe et al. , 2013). P_{CTR3} repressed by cheap
477 CuSO₄ was slight better to suppress *ERG9* expression to reduce FPP flux to squalene than P_{MET3}
478 repressed by methionine (Paddon et al. , 2013, Peng et al. , 2017). To achieve high
479 sesquiterpene, fermentation process needs to be optimized via such as response surface
480 methodology (Baadhe et al. , 2014) and feeding various carbon sources. Feedback-controlled
481 ethanol pulse feed has several significant advantages over restricted glucose feed. This strategy
482 is successfully applied to amorphadiene with titer of 41 g/L and artemisinic acid production in
483 *S. cerevisiae* CEN.PK2 Y293 (Paddon et al. , 2013, Westfall et al. , 2012). With development
484 of genomics and relevant technologies, comprehensive metabolic models were developed.
485 Model prediction via network models has been applied to sesquiterpene production: OptGene
486 (Burgard et al. , 2003, Patil et al. , 2005), as an extension modeling framework of OptKnock,
487 was subjected to minimization of metabolic adjustment (MOMA) method to identify *GDH1*
488 coding NADPH-dependent glutamate dehydrogenase as the best knockout target. Then,
489 through over-expressing *tHMGI* and replacing NADH-dependent *GDH2* with *GDH1* gene,
490 sesquiterpene titer increased to 30.1 mg/L from 11.3 mg/L (Asadollahi et al. , 2009). Flux
491 balance analysis (FBA) and MOMA methods were used for predicting knockout targets,
492 resulting in 54. 55 mg/L amorphadiene in the highest strain *hvk2D:: his3::ADS* (Sun et al. ,
493 2014). Assisted by 3D model of ADS protein, mutability landscape guided enzyme engineering
494 was performed to improve the catalytic activity, increasing catalytic efficiency ~ 4-fold in
495 mutant H448A and k_{cat} by 5-fold in mutant T399S/H448A (Abdallah et al. , 2018).

496 Based on FPP-biosynthesis platform EPY224, many studies were carried out: through
497 replacing *ADS* gene with *BIS* gene, more than 900 mg/L bisabolene was produced (Peralta-
498 Yahya et al. , 2011). Then, this EPY224-derived strain was applied to evolving *BIS*, enriching
499 a mixed population and removing little bisabolene producing strains in presence of selective
500 pressure nonionic surfactants Tween 20 (T20), for the protection of bisabolene against the
501 disruptive action of T20 (Kirby et al. , 2014a). Recently, *S. cerevisiae* EPY224 was re-
502 engineered to produce 344 mg/L pentalenene by replacing gene *BIS* with gene *coPentS* (Liu et
503 al. , 2018). In terpene production, screening color distribution of strains harboring carotenoid-
504 producing pathway is a tool of screening the targeted mutants (Furubayashi et al. , 2014,
505 Wadhwa and Bachhawat, 2019, Yuan and Ching, 2014) and identifying gene deletions
506 contributing to improving isoprenoid production (Ozaydin et al. , 2013). Combination of the
507 deletions and MVA pathway modifications increased bisabolene titer to 5.2 g/L in a
508 fermentation process (Ozaydin et al. , 2013).

509 In yeast, *FS* gene from *Malus domestica* led to higher α -farnesene titer (4 mg/L) than
510 plant-origins of *C. junos* and *A. annua*. Gene *FS* and *tHMG1* were over-expressed in a high-
511 copy plasmid in *S. cerevisiae* CEN.PK113-5D, which produced 170 mg/L farnesene in fed-
512 batch fermentations with a respiratory quotient (RQ)-controlled exponential feeding
513 (Tippmann et al. , 2016). After *FS* gene was replaced by santalene synthase from *Clausena*
514 *lansium*, this engineered *S. cerevisiae* produced 163 mg/L santalene under RQ-controlled
515 exponential feeding (Tippmann et al. , 2016). Before this, dynamic control of gene expression
516 including using P_{HXT1} promoter to control *ERG9* expression, overexpression of gene *tHMGR*
517 and deletion of gene *LPPI* and *DPPI* led strain to produce 92 mg/L α -santalene by diverting
518 the carbon flux FPP towards α -santalene (Scalcinati et al. , 2012a). In yeast, protein fusion
519 strategy could also make multi-enzyme complexes for increasing metabolic flux and compound
520 production. Comparing with that observed in a non-fused control, over-expression of Tya-

521 fused (Tya: Ty1 retrotransposon element) enzymes tHmg1, ispA and FS increased farnesene
522 production 4-fold, yielding 930 mg/L after 7 days in *S. cerevisiae* ATCC200589 (Han et al. ,
523 2018). Specifically, scaffold of FFPS and FS protein through recognizing affibodies to their
524 anti-idiotypic partners, the farnesene yield on glucose was improved by 135% (Tippmann et
525 al. , 2017a). Sequentially, acetoacetyl-CoA synthase (*nphT7*) identified to catalyze the
526 irreversible condensation of malonyl-CoA and acetyl-CoA to acetoacetyl-CoA from
527 *Streptomyces sp.* CL190, was expressed to increase flux, however, it was detrimental for
528 growth and farnesene production (Tippmann et al. , 2017b). Many important insights into
529 microbial metabolism have been achieved using metabolic nodes to disturb metabolic flux and
530 efficiency. Pantothenate (vitamin B5) was utilized as a “metabolic switch” for tuning synthesis
531 rates of molecules relying on CoA intermediates β -farnesene (Sandoval et al. , 2014). To
532 overcome the metabolic inefficiencies including a low ATP/oxygen ratio and challenges of
533 commodity-scale production of anabolic molecules in *S. cerevisiae*, gene aldehyde
534 dehydrogenase acylating (*ADA*), xylulose-5-phosphate specific phosphoketolase (*xPK*),
535 phosphotransacetylase (*PTA*) and NADH-*HMGR* were over-expressed, and DL-glycerol-3-
536 phosphatase (*Rhr2*) was deleted. They rewired the central carbon metabolism, reduced ATP
537 requirement and carbon loss, and improved pathway redox balance of cytosolic acetyl-CoA
538 biosynthesis, leading to farnesene titer exceeded 130 g/L using ~25% less glucose and ~75%
539 less O₂ than previous strains in a fed-batch fermentation (Meadows et al. , 2016, Yu and Chang,
540 2016). Additionally, evolutionary engineering discovered that truncated tTcb3p protein could
541 increase tolerance of replacement jet fuels blend AMJ-700t (10% cymene, 50% limonene, 40%
542 farnesene) in *S. cerevisiae* (Brennan et al. , 2015).

543 Compared with *S. cerevisiae* BY4741, industrial *S. cerevisiae* EG60 showed more
544 advantages on cineole and sesquiterpene production levels (Ignea et al. , 2011). In *S. cerevisiae*
545 EG60, through over-expressing gene *ID11*, *HMG2* (K6R) variant, *ERG20* and *SF* homologue

546 P330 from *Salvia pomifera*, and deleting a single copy of *ERG9* gene, the titers of *trans*- β -
547 caryophyllene, α -cubebene, α -copaene and δ -cadinene were respectively ~10 mg/L, ~7 mg/L,
548 ~12 mg/L and ~38 mg/L (Ignea et al. , 2011). This strain was further engineered to produce
549 125 mg/L caryophyllene through replacing *Salvia fruticosa* caryophyllene synthase Sf126 with
550 *SF*, as well as deleting the ubiquitin ligases *ubc7* and *ssm4/doa10* and endoplasmic reticulum
551 associated degradation resident protein *pho86* (Ignea et al. , 2012). In *S. cerevisiae*, over-
552 expressing gene *tHMG1* and down-regulating gene *ERG9* are two widely-used strategies, and
553 the effect on sesquiterpene biosynthesis was specially studied (Asadollahi et al. , 2008,
554 Asadollahi et al. , 2010). *S. cerevisiae* YZL141 was metabolically engineered to be a
555 sesquiterpene production platform by over-expressing gene *tHMG1*, *ERG20* and *FgJ03939*.
556 Based on this platform, sesquiterpene cyclase and skeletons of fusariumdiene and
557 fusagramineol from *Fusarium graminearum* were characterized, identifying 8 compounds -
558 nerolidol, (-)- α -acorenol, (E)- β -farnesene, (+)- α -bisabolol, (-)-acoradiene, fusariumdiene,
559 epi-fusagramineol and fusagramineol (Bian et al. , 2018). Mutation of *ERG9* gene and uptake
560 of exogenous ergosterol (sue) made *S. cerevisiae* WAT11 to produce about 90 mg/L 5-epi-
561 aristolochene, 90 mg/L premnaspirodiene and 20 mg/L valencene when respectively
562 introducing gene *TEAS* (tobacco epi-aristolochene synthase), *HPS* (*Hyoscyamus muticus*
563 premnaspirodiene synthase) and *CVS* (citrus valencene synthase) (Takahashi et al. , 2007).
564 Thereafter, applying this strategy in *S. cerevisiae* BY4741, co-expressing *HPS* gene with an
565 amino-terminal truncated *tHMGR* gene made it to produce 170.23 mg/L premnaspirodiene
566 (Zhuang and Chappell, 2015). Protein fusion engineering is an efficient strategy in
567 sesquiterpene production both in *E. coli* and yeast. Besides farnesene, it was also applied to
568 germacrene A production increasing 6-fold to 190.7 mg/L by over-expressing *tHMGR* gene
569 and fusing FPPS and germacrene A synthase (Hu et al. , 2017), and hirsutene and Δ 6-
570 protoilludene production by fusion of hirsutene synthase (*HS*) from *Stereum hirsutum* to an

571 HMGS domain (Flynn and Schmidt-Dannert, 2018). Optimizing FPP branch point, modulating
572 MVA pathway, modifying ammonium assimilation pathway and enhancing the transcriptional
573 activator activity as well as a specifically developed continuous fermentation process were
574 combined to produce 0.036 mg/g DCW of α -santalene (Scalcinati et al. , 2012b). Engineering
575 protein degradation of FPPS protein is an effective regulatory strategy to regulate monoterpene
576 and sesquiterpene production in *S. cerevisiae* (Peng et al. , 2018).

577 **Figure 5**

578 **Table 2**

579 4.3. Metabolic engineering strategies for sesquiterpene production in other microorganisms

580 Besides *E. coli* and *S. cerevisiae*, other microorganisms are also engineered to produce
581 sesquiterpenes, such as *Corynebacterium glutamicum*, *B. Subtilis*, *Lactococcus lactis*,
582 cyanobacterium and some eucaryons. The metabolic engineering and strategies for
583 sesquiterpene production in these microorganisms are summarized in Table 3. In *C.*
584 *glutamicum*, due to the low provision of FPP, heterologous expression of *CVS* gene from *Citrus*
585 *sinensis* could not enable (+)-valencene production. Upon deleting two endogenous
586 prenyltransferase genes and expressing either gene *ispA* from *E. coli* or gene *ERG20* from *S.*
587 *cerevisiae*, (+)-valencene production was observed. Employing *CVS* gene from *Nootka cypress*
588 further improved (+)-valencene titer to 2.41 mg/L, increasing 10-fold comparing to *VS* gene
589 from *C. sinensis* (Frohwitter et al. , 2014).

590 *B. subtilis* and *Lactococcus lactis* are two Gram-positive prokaryotes reported to be
591 engineered for sesquiterpene production. *B. subtilis* is a promising microbial host for chemical
592 production purpose because of its fast growth rate and generally regarded as safe status. In *B.*
593 *subtilis*, protein translation engineering was conducted by fusing the highly positive charged
594 six-arginine tag to *ADS* to increase protein expression. Coupled with expressing gene *Dxs* and
595 *Idi* under P_{xyIA} and systematic media optimization, ~20 mg/L amorphadiene was produced in

596 shake flask scale (Zhou et al. , 2013). *L. lactis*, harboring MVA pathway, is also an attractive
597 host for characterizing sesquiterpene production: TPS *VMPSTS* from *Vanda Mimi Palmer* was
598 characterized multiple sesquiterpenes with the dominating germacrene D (Song et al. , 2012).
599 Comparing with the β -sesquiphellandrene titer when only expressing β -sesquiphellandrene
600 synthase, co-expression of *mvk* and *mvaA* gene doubled the amount to 15.5 mg/L (Song et al. ,
601 2014). An industrial bacterium *Rhodobacter sphaeroides* engineered by co-expressing MVA
602 operon from *Paracoccus zeaxanthinifaciens* and *CVS* from *Callitropsis nootkatensis* produced
603 352 mg/L valencene in shaker flasks, 14-fold higher than that of only *CVS* expressing in *S.*
604 *cerevisiae* (Beekwilder et al. , 2014).

605 Cyanobacterium, as a photosynthetic prokaryote, was engineered to convert CO₂ and H₂O
606 directly into high-valued biofuel sesquiterpenes (Lin and Pakrasi, 2019). Filamentous
607 cyanobacterium *Anabaena sp.* PCC 7120 produced a farnesene photosynthetic productivity of
608 69.1 $\mu\text{g}\cdot\text{L}^{-1}\cdot\text{OD}^{-1}\cdot\text{d}^{-1}$ through its endogenous MEP pathway and *coFS* gene using CO₂,
609 mineralized water and light (Halfmann et al. , 2014). Unicellular cyanobacterium
610 *Synechococcus elongatus* PCC 7942 produced 4.6 mg/L farnesene or 19.8 mg/L amorphadiene
611 when over-expressing MEP pathway gene *Dxs*, *Idi* and *ispA*, with *FS* gene from apple or *ADS*
612 gene from *A. annua* with a dodecane overlay in 7 days (Choi et al. , 2016, Lee et al. , 2017).
613 With *QHS1* gene from *A. annua* integrated into cyanobacterium *Synechocystis sp.* PCC6803
614 genome, $\sim 4 \text{ mg gDCW}^{-1}$ β -caryophyllene was measured by a methanol: chloroform extraction
615 method (Reinsvold et al. , 2011). *Synechococcus sp.* PCC 7002 engineered by expressing *A.*
616 *grandis* *BIS* gene and deleting *glgC* gene produced more than 0.6 mg/L bisabolene with a
617 dodecane overlay (Davies et al. , 2014). Recently, through genetic engineering, serial enzyme
618 loading and artificial micro RNA-based repression of competing pathways, green microalga
619 *Chlamydomonas reinhardtii* produced 10.3 mg bisabolene $\cdot \text{g}^{-1}$ CDW, yielding up to 11.0 mg/L
620 by mixotrophic cultivation in 7 days (Wichmann et al. , 2018).

621 For eukaryote microorganism, *Yarrowia lipolytica*, *Rhodospiridium toruloides*,
622 *Streptomyces venezuelae* and *Xanthophyllomyces dendrorhous* are reported to produce
623 sesquiterpenes for their simple industrial scale-up operations. *Y. lipolytica* Po1h, with *tHMG1*,
624 *Idi*, *ERG20* and *coFS* gene integrated into the genome, produced 57.08 mg/L of α -farnesene, a
625 20.8-fold increase over the initial titer in shake flasks with YPD medium. Bioreactor scale-up
626 in PM medium led α -farnesene concentration up to 259.98 mg/L (Yang et al. , 2016c). The
627 carotenogenic yeast *R. toruloides* was reported to utilize lignocellulose hydrolysate to produce
628 bisabolene and amorphadiene. *R. toruloides* showed superior growth in corn stover
629 hydrolysates prepared by two pretreatment methods: biocompatible IL choline α -ketoglutarate
630 resulting in 261 mg/L bisabolene at bench scale; and an alkaline resulting in 680 mg/L
631 bisabolene in a high-gravity fed-batch bioreactor (Yaegashi et al. , 2017). In *S. venezuelae*,
632 10.52 mg/L bisabolene was produced after deleting FPP and GPP consumption gene
633 *SVEN_0269* and *SVEN_7111*, and inserting P_{SV}-coAgBis and P_{ermEp}-FPPS1 at the *aatP*
634 chromosomal site (Phelan et al. , 2015). Comparing α -cuprenene production in *E. coli*, *S.*
635 *cerevisiae* and *X. dendrorhous* expressing *Cop6* gene from *C. cinereus*, α -cuprenene titer in *X.*
636 *dendrorhous* reached to 80 mg/L, far higher than the other mutants (Melillo et al. , 2013).

637 **Table 3**

638 **5. Conclusions and perspective**

639 Biosynthesizing sesquiterpene by engineered microorganisms is a worthy-studied theme
640 for their widely promising applications and values. More and more sesquiterpenes and their
641 synthases are discovered from all kinds of organisms. Although great values contained in
642 sesquiterpenes, minute amount of sesquiterpenes are biosynthesized by native producers.
643 Benefit from the development of metabolic engineering and synthetic biology, microorganisms
644 have become tractable cell factories for sustainable biomanufacturing of the high-value
645 sesquiterpenes. Recent advances in cells and process engineering have made fermentation to

646 be an attractive route for sesquiterpene production from renewable, low-cost biomass
647 feedstocks.

648 Sesquiterpene production depends on metabolic strategies, and metabolic strategies also
649 need biotechnology innovation. Recent progress in synthetic biology of terpenoid synthases
650 has opened new avenues of research by providing diversities of strategies. Biotechnological
651 applications made possible by recent molecular advances including the engineering of
652 industrial microbiology and regulation of two isoprenoid pathways to increase the desirable
653 sesquiterpenes. The designing, engineering and optimizing processes are served to establish a
654 high efficiency fermentation-based process and stable sesquiterpene production hosts.
655 Although there have been successful applications of sesquiterpenes, like amorphadiene
656 (Westfall et al. , 2012) and farnesene (Harvey et al. , 2014), more studies and challenges need
657 to be performed and overcame to expand its transformational efficiency, stability and recovery.
658 For example, the use of solvents reduced sesquiterpene evaporation during fermentation and
659 improved product recovery, but it resulted in higher costs, due to the additional equipment cost
660 for solvent-product separation (Cuesta et al. , 2019). Further understanding of the enzymatic
661 mechanism and optimal expression of isoprenoid pathway enzymes will be necessitated.
662 Challenges on making bio-based sesquiterpene more competitive in applications come from
663 aspects of searching high-activity enzymes, supplying energy and cofactor, and balancing
664 metabolic flux, as well as efficient recovery and purification of sesquiterpenes from
665 fermentation broth. Furthermore, issues concerning microbial biosynthesis for achieving
666 higher production help to develop approaches of identifying regulatory effects in metabolic
667 pathways for redirecting carbon fluxes to terpenoid pathways.

668 **Ethical Statement**

669 This article does not contain any studies with human participants performed by any of the
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680 **Competing interests**

681 The authors declare that they have no competing interests.

682

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Table 1. Metabolic engineering of *E. coli* for sesquiterpene biosynthesis.

Sesquiterpene	Strain	Description	Titer	Reference
Amorphadiene	<i>E. coli</i> DYM1	Over-expressing <i>ADS</i> gene from <i>A. annua</i> and MVA pathway genes from <i>S. cerevisiae</i> , with <i>mvk</i> , <i>pmk</i> , <i>mvaD</i> , <i>idi</i> and <i>ispA</i> gene under control of P _{LAC} (plasmid pMBIS); <i>atoB</i> , <i>HMGS</i> and <i>tHMGR</i> gene under control of P _{LAC} (plasmid MevT)	112.2 mg/L	(Martin et al. , 2003)
	<i>E. coli</i> K12 MG1655 Δ recA, Δ endA	Containing plasmid P _{BAD} -Dxs-Idi-ispDF and P _{ACM} -ADS, over-expressing multidrug-resistant membrane efflux transporter protein TolC and ABC family transporters (macAB) or MFS family transporters (emrAB or emrKY)	~180 mg/L	(Zhang et al. , 2016)
	<i>E. coli</i> K12 MG1655	Containing plasmid P _{BAD} -Dxs-Idi-ispDF and P _{ACM} -ADS, deleting PTS genes, using optimized OPT2 medium	182 mg/L	(Zhang et al. , 2013)
	<i>E. coli</i> MG1655 DE3	Δ pstHIerr::FRT, containing plasmid pETA-TM2-galP-glk and pACM-P _{T7} -dxs-P _{T7} -idi-ADS-ispA	201 mg/L	(Zhang et al. , 2015)
	<i>E. coli</i> DP10	Containing plasmid pMBIS, supplementing 20 mM MVA to medium	~290 mg /L	(Pitera et al. , 2007)
	<i>E. coli</i> DH1	Plasmid pAM4: lacUV5 promoters for codon-optimized MevT operon and MBIS operon, Cm ^R , Plasmid pAM94: pTrc99A derivative containing <i>ADS</i> and codon-optimized <i>mvk</i> , Ap ^R ; fermentation for 75 h.	293 mg/L/OD ₆₀₀	(Anthony et al. , 2009)
	<i>E. coli</i> B121-Gold DE3	Over-expressing <i>Dxs</i> , <i>Idi</i> , <i>ispA</i> and <i>ADS</i> gene	~300 mg/L	(Yuan and Ching, 2014)
	<i>E. coli</i> BL21 (DE3)	Co-expressing <i>Dxs2</i> from <i>S. avermitilis</i> and <i>Idi</i> from <i>B. subtilis</i>	331.7 mg/L	(Wang et al. , 2015)
	<i>E. coli</i> BL21 (DE3)	Harboring pET21c-ADS, P _{T7} -tolC-tolC-acrB operon	404.8 mg/L	(Wang et al. , 2013)
	<i>E. coli</i> W3110 (ATCC 27325)	Containing plasmid pMBIS and MevT, employing a two-phase partitioning bioreactor with a dodecane organic phase	500 mg/L	(Newman et al. , 2006)
<i>E. coli</i> DH1	Codon-optimizing <i>mvk</i> and <i>pmk</i> gene, and replacing P _{lac} with a stronger P _{trc}	>500 mg/L	(Redding-Johanson et al. , 2011)	

	<i>E. coli</i> DH1	Bearing <i>D. acidovorans</i> <i>HMGR</i> by increasing intracellular NADH level using a NADP-dependent formate dehydrogenase (<i>fdh1</i>) from <i>Candida boidinii</i> , along with formate supplementation after 48 h of fermentation.	700 mg/L	(Ma et al. , 2011)
	<i>E. coli</i> DH1	Dynamic control the MVA pathway expression using the identified FPP-responding P_{rstA} and P_{gadE}	1400 mg/L	(Dahl et al. , 2013)
	<i>E. coli</i> DH1	Metabolic pathway optimization using rbs variants and combinatorial gene assembly, pTrc-RBS.ADS-MBIS, using 3% glycerol	3550 mg/L	(Nowroozi et al. , 2014)
	<i>E. coli</i> DH1	Heterologous MVA pathway with <i>HMGS</i> and <i>HMGR</i> from <i>S. aureus</i> , using a fermentation process of restricted glucose and nitrogen with NaOH pH control	29.7 g/L	(Tsuruta et al. , 2009)
α -Farnesene	<i>E. coli</i> DH5a	Codon-optimizing <i>FS</i> gene, augmenting metabolic flux for FPP synthesis by introducing MVA pathway and over-expressing <i>Dxs</i> , <i>Idi</i> and <i>ispA</i> gene, fusing <i>FPPS</i> and <i>FS</i>	380.0 mg/L	(Wang et al. , 2011a)
	<i>E. coli</i> BL21 (DE3)	Developing a reconstituted in vitro farnesene biosynthetic system, rationally designing strain F4 with P_{lac} for <i>ERG12</i> , <i>ERG8</i> , <i>MVD1</i> and <i>Idi</i> gene with pBBR1MCS ori, and P_{T7} for <i>AFS</i> , <i>ispA</i> and <i>Idi</i> gene with pBBR322 ori	1100 mg/L	(Zhu et al. , 2014)
	<i>E. coli</i> BL21 (DE3)	Expressing heterologous MVA pathway, and high-expressing <i>Idi</i> and <i>ispA</i> gene in a lab bioreactor.	8740 mg/L	(You et al. , 2017)
Bisabolene	<i>E. coli</i> BL21 (DE3)	Harboring plasmid pBbA1k-AgBIS-ispA-ispDF-idi, fusing nDXP gene <i>ribB</i> to <i>Dxr</i> gene,	8.8 mg/g DCW	(Kirby et al. , 2015)
	<i>E. coli</i> DH1	F-, λ -, endA1, recA1, gyrA96, thi-1, glnV44, relA1, hsdR17(rK- mK+), DSMBT7, rbsR:: P_{T7} -BS-idi-ispA, poxB:: P_{T7} -MevT	~435 μ g/L	(Alonso-Gutierrez et al. , 2018)
	<i>E. coli</i> DH1	Screening BIS enzyme, codon-optimizing <i>HMGS</i> , <i>tHMGR</i> , <i>mvk</i> and <i>pmk</i> gene, and inserting a strong P_{trc} before <i>mvk</i> gene (pJBEI-2999)	> 900 mg/L	(Peralta-Yahya et al. , 2011)
	<i>E. coli</i> DH1	Using quorum sensing LuxI/R system from <i>Vibrio fischeri</i> to construct sensor plasmid carrying <i>luxI-luxR</i> genes, and response plasmid carrying bisabolene producing pathway genes under control of P_{luxI}	1100 g/L	(Kim et al. , 2017)

	<i>E. coli</i> DH1	Harboring plasmid pBbA5c-CL1-T61-CL2-T21-AgBISppTrc-AgBIS	1150 mg/L	(Alonso-Gutierrez et al. , 2015)
β-Caryophyllene	<i>E. coli</i> DH1	Expressing MVA pathway and <i>QHS1</i> gene	100 mg/L	(Wu et al. , 2018)
	<i>E. coli</i> BL21 (DE3)	Co-expressing <i>ACS</i> , <i>QHS1</i> , <i>GPPS2</i> , <i>AACS</i> gene and heterologous MVA pathway	1050 mg/L	(Yang and Nie, 2016)
	<i>E. coli</i> BL21 (DE3)	Over-expressing MVA pathway and <i>GPPS</i> , <i>G6PH</i> , <i>QHS1</i> genes, fed-batch fermentation for 60 h	1520 mg/L	(Yang et al. , 2016a)
α-Humulene	<i>E. coli</i> BL21 (DE3)	Expressing <i>Streptomyces</i> MVA pathway genes from <i>HMGS</i> to <i>Idi2</i> , yeast <i>Idi1</i> and rat <i>Aacl</i> gene, α-humulene synthase gene of shampoo ginger, supplying with LAA	958 mg/L	(Harada et al. , 2009)
5-Epi-aristolochene	<i>E. coli</i> XL1-Blue	P _{lac} -crtM-crtN, <i>TEAS</i> from <i>Nicotiana tabacum</i> , using the screened TEASmut7 Q481R by scoring their substrate consumption based on color loss of cell harboring carotenoid pathways	160 mg/L	(Furubayashi et al. , 2014)
Protoilludene	<i>E. coli</i> DH5α	Over-expressing a exogenous MVA pathway, endogenous <i>ispA</i> , and protoilludene synthase of <i>Omphalotus olearius</i>	1199 mg/L	(Yang et al. , 2016b)
Valerenadiene	<i>E. coli</i> BL21 (DE3)	Codon-optimizing gene <i>cvds</i> , over-expressing <i>Dxs</i> , <i>Idi</i> and <i>FPPS</i> , optimizing fermentation medium with glycerol as supplementary	11 mg/L	(Nybo et al. , 2017)
Longifolene	<i>E. coli</i> BL21 (DE3)	pTrcLower: pTrcHis2B harboring <i>S. cerevisiae</i> <i>erg19</i> , <i>erg8</i> , <i>erg12</i> and <i>idi1</i> ; pA-mvaESispALFS: pACYCDuet-1 harboring <i>E. faecalis</i> <i>mvaE</i> , <i>mvaS</i> , <i>E. coli</i> <i>ispA</i> and <i>P. abies</i> longifolene synthase	382 mg/L	(Cao et al. , 2019)

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Table 2. Metabolic engineering of *S. cerevisiae* for sesquiterpene biosynthesis.

Sesquiterpene	Strain	Description	Titer	Reference
Amorphadiene	<i>S. cerevisiae</i> WAT11	Using flux balance analysis (FBA) and MOMA methods for predicting knockout targets, strain MATa (leu2-3,112 trp1-1 can1-100 ura3-1 ade2-1 his3-11,15, <i>hvk2D::his3::ADS</i>)	54.6 mg/L	(Sun et al. , 2014)
	<i>S. cerevisiae</i> EPY224	Upregulating <i>HMGS</i> , <i>mvk</i> and <i>pmk</i> gene by <i>upc2-1</i> expression; over-expressing a truncated <i>tHMGR</i> , <i>ispA</i> and <i>ADS</i> gene; <i>erg9</i> : P _{met3} -ERG9	153 mg/L	(Ro et al. , 2006)
	<i>S. cerevisiae</i> CEN.PK2 Y293	Genotype: MATa, <i>erg9Δ::kan^r</i> P _{MET3} -ERG9, leu2-3,112::HIS P _{GAL1} -MVD1, P _{GAL10} -ERG8, his3Δ1::HIS, P _{GAL1} -ERG12, P _{GAL10} -ERG10, <i>ade1Δ::P_{GAL1}-tHMG1</i> , P _{GAL10} -IDI1 ADE1, ura3-52::P _{GAL1} -tHMG1, P _{GAL10} -ERG13, URA3trp1-289::P _{GAL1} -tHMG1, P _{GAL10} -ERG20 TRP1; gal80Δ::nat ^r ; plasmid pAM426: P _{GAL1} -tHMG1. Fermentation with unrestricted ethanol pulse feed with lower oxygen uptake rate (OUR)	41 g/L	(Westfall et al. , 2012)
Bisabolene	<i>S. cerevisiae</i> EPY224	EPY224 platform through replacing <i>ADS</i> gene with <i>BIS</i> gene	900 mg/L	(Peralta-Yahya et al. , 2011)
	<i>S. cerevisiae</i> EPY224	EPY224 platform through replacing <i>ADS</i> gene with <i>BIS</i> gene, enriching a mixed population and removing little or no bisabolene producing strains using selective pressure nonionic surfactants (T20)	375 mg/L	(Kirby et al., 2014a)
	<i>S. cerevisiae</i> CEN.PK2-1C	Genotype: MATa ura3-52 trp1-289 leu2-3,112 his3D 1 MAL2-8C SUC2, YPRCd15::KANMX-PGAL1-C.O.BisSyn~ERG20/P _{GAL10} -tHMG, gal80::HIS5, ypl064w::hph yjl064w::Ca.URA3, harboring pRS-AgBIS	5.2 g/L	(Ozaydin et al. , 2013)
β-Farnesene	<i>S. cerevisiae</i> CEN.PK113-5D	Genotype: MATa, MAL2-8c, SUC2, ura3-52, <i>lpp1Δ::loxP</i> , <i>dpp1Δ::loxP</i> , PERG9Δ::loxP-P _{HXT1} , P _{PGK1} -Z _{IgA} -ERG20, P _{TEF1} -Z _{Taq} -FarnSyn-Cj, P _{TEF1} -anti-Z _{Taq} -anti-Z _{IgA} with (SSSSG) ₄ linker	0.813 mg/g	(Tippmann et al. , 2017a)
	<i>S. cerevisiae</i> CEN.PK113-5D	Genotype: MATa, MAL2-8 ^c , SUC2, ura3-52, <i>lpp1Δ::loxP</i> , <i>dpp1Δ::loxP</i> , P _{ERG9Δ} ::loxP-P _{HXT1} , <i>gdh1Δ::loxP</i> , P _{TEF1} -ERG20, P _{PGK1} -GDH2, over-expressing <i>FS</i> and <i>tHMG1</i> gene in a 2-μm plasmid with respiratory quotient-controlled exponential feeding	170 mg/L	(Tippmann et al. , 2016)

	<i>S. cerevisiae</i> ATCC200589	Over-expressing structural component Ty1 retrotransposon element Tya-fused enzyme tHmg1, ispA and FS in a two-phase partitioning fed-batch fermentation for 7 days	930 mg/L	(Han et al. , 2018)
	<i>S. cerevisiae</i> CEN.PK2-1C	Genotype: MAT α , erg9 Δ ::P _{GAL4} (OC)-GAL4_syn_P _{MET3} -ERG9, trp1-289::P _{GAL1} -tHMG1_P _{GAL10} -ERG20_T _{RP1} , leu2-3,112::HISM _X _P _{GAL1} -MVD1_P _{GAL10} -ERG8, ade1 Δ ::P _{GAL1} -tHMGR_P _{GAL10} -IDI1_ADE1, his3 Δ 1::HISM _X _P _{GAL1} -ERG12_P _{GAL10} -ERG10_HIS3, ura3-52:: P _{GAL1} -tHMG1_P _{GAL10} -ERG13_URA3, gal80 Δ ::syn_P _{GAL1} -ERG12, erg9 Δ ::P _{GAL4} (OC)-GAL4_syn_P _{MET3} -ERG9, hxt3 Δ ::syn_P _{TDH3} -ERG10-T _{AHP1} -P _{YPD1} -Bj.HMGs-T _{CCW12} -P _{TUB2} -Ca.THL-T _{HXT3} , dit1 Δ ::syn_P _{GAL1} -Aa.FS_A-T _{CYC1} , ndt80 Δ :: P _{GAL1} -Aa.FS_A-T _{CYC1} -P _{HXT3} -ACS2-T _{PGK1} -P _{GAL7} -Zm.PDC-T _{TDH3} , ho Δ ::syn, gal2 Δ ::syn, adh5 Δ :: P _{GAL1} -Aa.FS_B-T _{ADH1} -P _{GAL10} -Aa.FS_A-T _{CYC1} -LEU2_P _{GAL1} -Aa.FS_B-T _{ADH1} -P _{GAL10} -Aa.FS_A-T _{CYC1} ; ald4 Δ ::P _{GAL1} -Dz.eutE-T _{PGK1} -syn_P _{GAL1} -Dz.eutE-T _{PGK1} , ald6 Δ ::syn, rhr2 Δ ::syn_P _{TDH3} -Lm.PK-T _{TDH3} , erg13 Δ ::syn_P _{HXT1} -GAL4*, ygr250c Δ ::syn_P _{ACT3} -PDR3, bdh1 Δ ::P _{TDH3} -Lm.PK-T _{TDH3} -syn_P _{ACT1} -Ck.PTA-T _{PGK1} , adh1 Δ ::P _{FBA1} -ACS2_syn_P _{TDH3} -Sp.HMGr-T _{HMG1} , P _{TDH3} -ERG13*_P _{GAL10} -Sp.HMGr-T _{HMG1} ; 200,000-L industrial bioreactors.	130 g/L	(Meadows et al. , 2016)
α -Santalene	<i>S. cerevisiae</i> CEN.PK113-5D	Genotype: MAT α MAL2-8c SUC2 ura3-52 lpp1 Δ ::loxP dpp1 Δ ::loxP P _{ERG9} Δ ::loxP-P _{HXT1} , plasmid pICK01 containing a copy of tHMG1 and santalene synthase	92 mg/L	(Scalcinati et al. , 2012a)
	<i>S. cerevisiae</i> CEN.PK113-5D	Genotype: MAT α , MAL2-8 c , SUC2, ura3-52, lpp1 Δ ::loxP, dpp1 Δ ::loxP, P _{ERG9} Δ ::loxP-P _{HXT1} , gdh1 Δ ::loxP, P _{TEF1} -ERG20, P _{PGK1} -GDH2, over-expressing santalene synthase and <i>tHMG1</i> gene in a 2- μ m plasmid with respiratory quotient-controlled exponential feeding	163 mg/L	(Tippmann et al. , 2016)
Caryophyllene	<i>S. cerevisiae</i> EG60	Genotype: Mat a/ α , P _{Gal1} -HMG2-(K6R) x2, P _{Gal1} -HA-ERG20, ura3, trp1, his3, leu2, Δ erg9:: HIS5, pYESmyc/P330, 2 μ , URA3, P _{GAL1} -myc-P330	~ 10 mg/L	(Ignea et al. , 2011)
	<i>S. cerevisiae</i> EG60	Genotype: Mat a/ α , P _{Gal1} -HMG2(K6R):: HOX2, ura3, trp1, his3, P _{TDH3} -HMG2(K6R)X2-::leu2 ERG9/erg9, UBC7/ubc7, SSM4/ssm4, PHO86/pho86	125 mg/L	(Ignea et al. , 2012)
α -Cubebene	<i>S. cerevisiae</i> EG60	Genotype: Mat a/ α , P _{Gal1} -HMG2-(K6R) x2, P _{Gal1} -HA-ERG20, ura3, trp1, his3, leu2, Δ erg9:: HIS5, pYESmyc/P330, 2 μ , URA3, P _{GAL1} -myc-P330	~ 7 mg/L	(Ignea et al. , 2011)

α -Copaene	<i>S. cerevisiae</i> EG60	Genotype: Mat a/ α , P _{Gal1} -HMG2-(K6R) x2, P _{Gal1} -HA-ERG20, ura3, trp1, his3, leu2, Δ erg9:: HIS5, pYESmyc/P330, 2 μ , URA3, P _{GAL1} -myc-P330	~12 mg/L	(Ignea et al. , 2011)
α -Cadinene	<i>S. cerevisiae</i> EG60	Genotype: Mat a/ α , P _{Gal1} -HMG2-(K6R) x2, P _{Gal1} -HA-ERG20, ura3, trp1, his3, leu2, Δ erg9:: HIS5, pYESmyc/P330, 2 μ , URA3, P _{GAL1} -myc-P330	~38 mg/L	(Ignea et al. , 2011)
Pentalenene	<i>S. cerevisiae</i> EPY	Upregulating <i>HMGS</i> , <i>MK</i> and <i>PMK</i> gene by <i>upc2-1</i> expression; over-expressing <i>tHMGR</i> , <i>ispA</i> and <i>PentS</i> gene; <i>erg9</i> : P _{met3} -ERG9	344 mg/L	(Liu et al. , 2018)
5-Epi-aristolochene	<i>S. cerevisiae</i> WAT11	Mutating gene <i>ERG9</i> , uptake of exogenous ergosterol (<i>sue</i>), introducing gene <i>TEAS</i> in plasmid pESC-HIS3-GPD-TEAS	90 mg/L	(Takahashi et al. , 2007)
Premnaspirodiene	<i>S. cerevisiae</i> WAT11	Mutating gene <i>ERG9</i> , uptake of exogenous ergosterol (<i>sue</i>), introducing gene <i>HPS</i> in plasmid pESC-HIS3-GPD-HPS	90 mg/L	(Takahashi et al. , 2007)
	<i>S. cerevisiae</i> BY4741	Expressing <i>HPS</i> gene with an amino-terminal truncated in plasmid pESC-HIS3-GPD-HPS, catalytic form of <i>tHMGR</i> gene, uptake of exogenous ergosterol (<i>sue</i>)	170.23 mg/L	(Zhuang and Chappell, 2015)
germacrene A	<i>S. cerevisiae</i> SCIGS22a	MATa MAL2-8c SUC2 ura3-52 <i>lpp1</i> Δ ::loxP <i>dpp1</i> Δ ::loxP PERG9 Δ ::loxP-P _{HXT1} <i>gdh1</i> Δ ::loxP P _{TEF1} -ERG20, P _{GK1} -GDH2 P _{TEF1} - <i>tHMG1</i> , P _{TEF1} -ERG20-GGGGS-LTC2 _{opt} aaaaaaaaaa P _{PGK1} - <i>tHMG1</i>	190.7 mg/L	(Hu et al. , 2017)
Valencene	<i>S. cerevisiae</i> YIP-00-03 (CEN.PK113-5D)	MATa MAL2-8c SUC2 ura3-52, <i>erg9</i> ::P _{MET3} -ERG9, pESC-URA 2 μ URA3 P _{GAL1} - GFTpSD	2 mg/L	(Asadollahi et al. , 2008)
	<i>S. cerevisiae</i> WAT11	Mutating <i>ERG9</i> gene, uptake of exogenous ergosterol (<i>sue</i>), introducing <i>CVS</i> gene	20 mg/L	(Takahashi et al. , 2007)

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Table 3. Metabolic engineering of other organisms for sesquiterpene biosynthesis.

Sesquiterpene	Strain	Description	Titer	Reference
(+)-Valencene	<i>C. glutamicum</i> ATCC 13032	Deleting two endogenous prenyltransferase genes, expressing heterologous gene <i>ispA</i> from <i>E. coli</i> or gene <i>ERG20</i> from <i>S. cerevisiae</i>	2.41 mg/L	(Frohwitter et al. , 2014)
Amorphadiene	<i>B. subtilis</i> 1A1 (BGSC)	Expressing gene <i>ADS</i> under P _{grac} , gene <i>Dxs</i> and <i>Idi</i> under P _{xylA} , protein translation engineering fused six-arginine tag to N terminus of gene <i>ADS</i> , optimizing media by adding pyruvate and dipotassium phosphate	~20 mg/L	(Zhou et al. , 2013)
β-Sesquiphellandrene	<i>L. lactis</i> NZ9000	Introducing β-sesquiphellandrene synthase, co-expressing <i>mvk</i> and <i>mvaA</i> gene	15.5 mg/L	(Song et al. , 2014)
Valencene	<i>R. sphaeroides</i>	Co-expressing MVA operon from <i>Paracoccus zeaxanthinifaciens</i> and <i>CVS</i> from <i>Callitropsis nootkatensis</i> on plasmid (pBBR-MEV) in shaker flasks	352 mg/L	(Beekwilder et al. , 2014)
Farnesene	<i>Anabaena</i> sp. PCC 7120	Expressing a <i>coFS</i> gene, through its endogenous MEP pathway using CO ₂ , mineralized water, and light	69.1 μg·L ⁻¹ ·O ₂ ⁻¹ ·d ⁻¹	(Halfmann et al. , 2014)
	<i>S. elongatus</i> PCC 7942	<i>NSI::Bb1s-dxs-idi-ispA</i> , <i>NSII::Bb1k-AFS</i> direct conversion of CO ₂ , with a dodecane overlay in 7 days	4.6 mg/L	(Lee et al. , 2017)
Amorphadiene	<i>S. elongatus</i> PCC 7942	<i>NSI::Bb1 s-dxs-idi-ispA</i> <i>NSII::Bb2 k-ADS</i> , direct conversion of CO ₂ , with a dodecane overlay in 7 days	19.8 mg/L	(Choi et al. , 2016)
Caryophyllene	<i>Synechocystis</i> sp. PCC6803	Inserting <i>QHSI</i> gene from <i>A. annua</i> via double homologous recombination into genome, converting CO ₂ and H ₂ O, measured by a methanol: chloroform extraction method	~ 4 mg/g DCW	(Reinsvold et al. , 2011)
Bisabolene	<i>Synechococcus</i> sp. PCC 7002	Expressing heterologous <i>A. grandis</i> <i>BIS</i> gene, deleting <i>glgC</i> gene and applying a dodecane overlay	>0.6 mg L ⁻¹	(Davies et al. , 2014)
	<i>C. reinhardtii</i>	Through genetic engineering, serial enzyme loading and artificial micro RNA-based repression of competing pathways	11.0 mg/L	(Wichmann et al. , 2018)
Farnesene	<i>Y. lipolytica</i> Po1h	Integrating <i>tHMG1</i> , <i>Idi</i> , <i>ERG20</i> and <i>coFS</i> gene into genome in a bioreactor scale-up with PM medium	259.98 mg/L	(Yang et al. , 2016c)

Amorphadiene	<i>R. toruloides</i> IFO0880	Mating type A2, P _{GAPDH} -BIS-TNOS, in corn stover hydrolysates prepared by a biocompatible ionic liquid (IL) choline α -ketoglutarate pretreatment	36 mg/L	(Yaegashi et al. , 2017)
Bisabolene	<i>R. toruloides</i> IFO0880	Mating type A2, plasmid P _{GAPDH} -BIS-TNOS, with corn stover hydrolysates prepared by an alkaline pretreatment, in high-gravity fed-batch bioreactor	680 mg/L	(Yaegashi et al. , 2017)
Bisabolene	<i>S. venezuelae</i> ATCC 10712	Deleting gene <i>ΔSVEN_0269</i> and <i>ΔSVEN_7111</i> , inserting P _{SV} -coAgBis and P _{ermEp} -FPPS1 at the <i>aatP</i> chromosomal site	10.52 mg/L	(Phelan et al. , 2015)
α -cuprenene	<i>X. dendrorhous</i>	Integrating gene <i>Cop6</i> in the rDNA from <i>C. cinereus</i>	80 mg/L	(Melillo et al. , 2013)

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