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3 ANALISIS KANDUNGAN SESQUITERPENE pada GAHARU HASIL
4 INOKULASI DENGAN TEKNIK SIMPORI

5 (14pt)

6
7 SESQUITERPENE CONTENT ANALYSIS OF AGARWOOD INOCULATED USING SIMPORI
8 TECHNIQUE

9 (12 pt)

10
11 **Dian Pratiwi¹, Resti Wahyuni^{1*} (12pt)**

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14 *Corresponding author: resti.wahyuni@brin.go.id (10 pt)

15
16 **Abstract (12 pt)**

17 Gaharu merupakan resin yang terbentuk pada spesies anggota famili *Thymelaeaceae*, termasuk
18 *Gyrinops versteegii* yang mengalami perlukaan atau terinfeksi oleh mikroorganismenya. Kualitas
19 gaharu baik alami maupun budidaya dapat dilihat dari kandungan sesquiterpene. Penelitian ini
20 bertujuan mendeteksi kandungan sesquiterpene pada *G. versteegii* hasil inokulasi dengan teknik
21 simpori pada 3 dosis inokulan berbeda yaitu 3 ml/paku simpori, 9 ml/paku simpori, dan 18 ml/paku
22 simpori dan dipanen saat 18 bulan setelah inokulasi. Gaharu yang telah dipanen kemudian diekstrak
23 menggunakan pelarut n-hexana. Ekstrak n-hexana selanjutnya dianalisis kandungan kimianya
24 menggunakan GC-MS. Data yang digunakan pada hasil adalah deteksi senyawa dengan persentase
25 Similarity Index (SI) di atas 80%. Untuk menguji signifikansi perlakuan maka selanjutnya
26 dilakukan uji statistik dengan taraf signifikansi 5%. Berdasarkan uji Kruskal Wallis, Sig. hitung <
27 0,05 artinya terdapat perbedaan yang signifikan dari tiga perlakuan dosis inokulan terhadap
28 persentase senyawa sesquiterpene yang dihasilkan. Dosis 9ml/paku simpori merupakan dosis
29 terbaik dengan Mean Rank sebesar 16 dan memiliki kandungan sesquiterpene tertinggi yaitu
30 23,69%. Senyawa sesquiterpene yang terdeteksi pada gaharu hasil inokulasi dengan dosis 9ml/paku
31 simpori adalah aromadendrene; α -Selinene; (1aS,4aR)-4a,8,8-Trimethyl-2-methylene-
32 1,1a,2,4a,5,6,7,8-ctahydrocyclopropa[d]naphthalene; 6-Methyl-2-(4-methylcyclohex-3-en-1-
33 yl)hepta-1,5-dien-4-ol; alloaromadendrene; dan eremophilene. Penggunaan dosis inokulan
34 9ml/paku simpori dapat direkomendasikan untuk optimalisasi produksi gaharu yang lebih efisien.

35
36
37 **Kata kunci:** Aromadendrene; Eremophilene; Gaharu; Sesquiterpene; Simpore (11 pt)

38
39 **Abstract (12 pt)**

40 Agarwood is a resin formed in the plants that belong to the *Thymelaeaceae* family, including
41 *Gyrinops versteegii*, as a result of damage or microbial infection. The content of sesquiterpenes in
42 both the wild and cultivated agarwood indicates its quality. Nevertheless, information regarding the
43 comparison between the content has not been provided. This study aimed at identifying the
44 sesquiterpenes in *G. versteegii* inoculated using simpore technique with various dosages of
45 inoculants (3 ml, 9 ml, and 18 ml/porous nail) and harvested at 18 months after inoculation. The
46 harvested agarwood was extracted using n-hexane solvent. The chemical compositions were
47 subsequently analyzed using the GC-MS. Compounds with similarity index above 80% were
48 analyzed, including their presence and percentage in the agarwood. The statistical significance test
49 was carried out with the 5% level of significance. The Kruskal-Wallis test showed p-value < 0.05,
50 indicating the significant differences between the inoculant dosages on the percentage of

51 sesquiterpenes. Agarwood with *F. solani* at a dosage of 9 ml/porous nail shows the best result with
52 a Mean Rank of 16 and 23.69% of sesquiterpenes. It contains aromadendrene; α -Selinene;
53 (1aS,4aS,8aR)-4a,8,8-Trimethyl-2-methylene-1,1a,2,4a,5,6,7,8-ctahydrocyclopropa[d]naphthalene;
54 6-Methyl-2-(4-methylcyclohex-3-en-1-yl)hepta-1,5-dien-4-ol; alloaromadendrene; and
55 eremophilene. Therefore, this dosage is recommended for optimizing agarwood production.

56
57 **Keywords:** Agarwood; Aromadendrene; Eremophilene; Sesquiterpene; Simpori (11 pt)

58 59 **INTRODUCTION (12 pt)**

60 Agarwood is one of the most sought non-timber forest products due to its high economic and
61 aesthetic values, including for health, religious ceremonies, and aromatic purposes (Batubara et al.,
62 2022) (Tamyiz et al., 2022) (Saputra et al., 2024). Agarwood is a resin formed in the plants that
63 belong to the Thymelaeaceae family as a result of damage or microbial infection (Wahyuni et al.,
64 2020). In Indonesia, the major sources of agarwood are 26 species of seven genera, namely
65 *Aquilaria*, *Gonystylus*, *Gyrinops*, *Aetoxylon*, *Enkelia*, *Phaleria*, and *Wikstroemia*, which grow
66 naturally in the wild (Lukman et al., 2022). The first three genera are the most frequently used
67 (Sutomo et al., 2021). *Gyrinops* dominated the eastern, while *Aquilaria* dominated the western
68 Indonesia (Roemantyo & Partomihardjo T, 2010).

69 *Gyrinops versteegii* is a major source of agarwood, particularly obtained from West Nusa
70 Tenggara, East Nusa Tenggara, and Papua (Wahyuni et al., 2020). It is offered in various forms,
71 from large-sized raw materials to powder, as well as finished products such as perfume and incense
72 (Ceniza et al., 2021). Nevertheless, harvesting in nature has caused its amount to decrease
73 drastically. Consequently, it is listed in the CITES Appendix II to specify that the trade of this
74 species and its derivatives is restricted (Sutomo et al., 2021). Moreover, species belonging to
75 *Aquilaria* and *Gyrinops* are also categorized “Critically endangered”, “Data deficient”,
76 “Endangered”, and “Vulnerable” by the International Union for Conservation of Nature (IUCN)
77 (Syameera et al., 2024).

78 The natural formation of agarwood requires a slow process and occurs only on 10 percent of
79 wild trees due to fungal infection or physical damage (Gogoi et al., 2023). Therefore, cultivation is
80 an alternative to obtain agarwood and to reduce the wild harvest at once. This effort comprises
81 cultivation and inoculation technology to trigger the natural formation of agarwood. A modified
82 inoculation technique called *simpori* has been conducted on *G. versteegii* in West Nusa Tenggara
83 (Wahyuni et al., 2020). The technique combines nailing the tree trunk using porous nails and filling
84 it with *Fusarium solani* in a liquid medium at a certain concentration. Previous study showed that
85 the visual quality of agarwood using the technique with 3, 9, and 6 ml of *F. solani* inoculant per
86 porous nail, and harvested months after inoculation was similar, namely the quality of agarwood
87 with mastic content and weaker aroma (Grade C-medium) (SNI 7631:2011, 2011). Sesquiterpene
88 and chromone compounds were also found in the agarwood with slight variations (Wahyuni et al.,
89 2020).

90 Sesquiterpene is a substantial chemical component for determining the quality of agarwood
91 (Yu et al., 2023). The higher the sesquiterpene content in agarwood, the higher the quality of the
92 agarwood. In addition, it is relatively stable against environmental changes, including temperatures
93 (Syameera et al., 2024). A study on agarwood in wild *Gyrinops salicifolia* identified that it
94 contained 12 sesquiterpenoids (Shao et al., 2016). Nevertheless, the information about
95 sesquiterpene compound of agarwood in *G. versteegii*, both in the wild and cultivated, is
96 inadequate. After 7 months, *G. versteegii* inoculated using *simpori* technique yielded agarwood
97 with alloaromadendrene and valerenol contents (Wahyuni et al., 2020). Information on
98 sesquiterpenes of agarwood in the cultivated *G. versteegii* is required to provide a comparative
99 overview of quality with the wild ones. This study aims to identify the sesquiterpene content of
100 agarwood in the cultivated *G. versteegii* inoculated using the *simpori* technique at various inoculant
101 concentrations, namely 3, 9, and 18 ml/porous nail, and harvested 18 months after inoculation.

103 MATERIALS AND METHODS (12 pt)

104 Study Site/Location and/or Materials

105 This study was carried out in 2019. The extraction of sapwood was performed at the NTFPs
106 processing laboratory, Research and Development Institute of Technology for Non-timber Forest
107 Products (Balai Litbang Teknologi HHBK) Mataram, West Nusa Tenggara, Indonesia. The GC-MS
108 analysis was done at Research Center for Chemistry-Indonesian Institute of Sciences, Serpong,
109 Indonesia. The characteristics of sapwood agarwood were observed from the cultivated *G.*
110 *versteegii* that was inoculated using the *simpori* technique at various inoculant concentrations,
111 namely 3, 9, and 18 ml/porous nail. The agarwood was harvested 18 months after inoculation.

112

113 Methods

114 Extraction of Sapwood Agarwood

115 The sapwood was harvested, air-dried at room temperature, and grinded using a blender. The
116 powder was extracted using n-hexane repeatedly until the extract liquid was clear. The agarwood
117 powder to solvent ratio was 1:5. Subsequently, the yield was filtered and concentrated using a
118 rotary vacuum evaporator.

119

120 GC-MS analysis 22

121 A sample of 1 μ l was injected into the GC-MS instrument with glass column of 2.5 m long,
122 0.25 m diameter, 0.25 μ m thickness, and P-Sil 5 CB stationary phase. The carrier gas was helium.
123 The device starting temperature was 40 °C held for 1 min and increased at 10 °C per minute until
124 250 °C. The temperature was held for 6 min since the compounds might have different retention
125 times. Each compound's chromatogram and similarity index (SI) were analyzed using NIST MS
126 Search version 2.0.

127

128 Statistical Analysis

129 The compounds with similarity index (SI) above 80% were identified for further analysis. The
130 presence and percentage of each compound on the agarwood were determined. A test of the
131 statistical significance was carried out with the 5% level of significance.

132

133 RESULTS (12 pt)

134 The sesquiterpenes in agarwood from *G. versteegii* inoculated using *simpori* technique are
135 illustrated in Table 1.

136

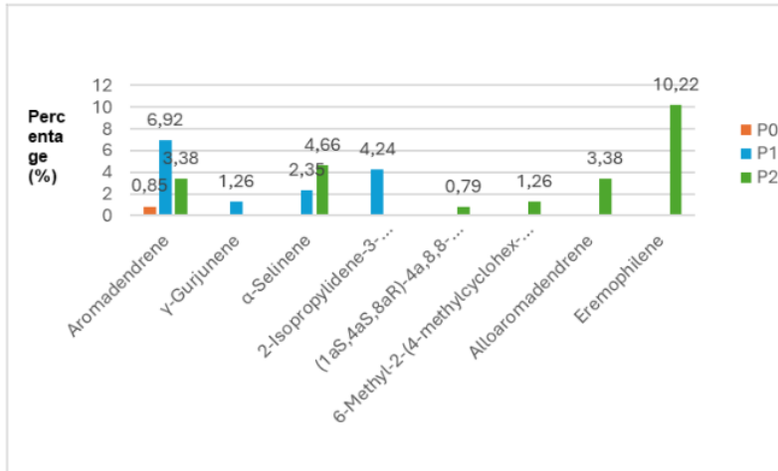
137 **Table 1.** Sesquiterpenes in the agarwood formed in *G. versteegii* inoculated using *simpori*
138 technique with various dosages of inoculant

No.	Compound	Presence in the agarwood			139
		P0	P1	P2	
1.	Aromadendrene	+	+	+	141
2.	γ -Gurjunene	-	+	-	142
3.	α -Selinene	-	+	+	143
4.	2-Isopropylidene-3-methylhexa- 3,2-dienal	-	+	-	144
5.	(1aS,4aS,8aR)-4a,8,8-Trimethyl-2- methylene-1,1a,2,4a,5,6,7,8- 3,3-tetrahydrocyclopropa[d]naphthalene	-	-	+	145
6.	6-Methyl-2-(4-methylcyclohex-3- en-1-yl)hepta-1,5-dien-4-ol	-	-	+	146
7.	Alloaromadendrene	-	-	+	147
8.	Remophilene	-	-	+	148
					149
					150

151 *P0 = *F. solani* at a dosage of 3 ml/porous nail; P1 = *F. solani* at a dosage of 18 ml/porous nail; P2 = *F. solani* at a dosage of 9
152 ml/porous nail; + = detected, - = not detected

153

154 The percentage of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori*
155 technique is shown in Figure 1.



157

158 **Figure 1.** Percentage of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori*
 159 technique with various dosages of inoculant

160 *P0 = *F. solani* at a dosage of 3 ml/porous nail; P1 = *F. solani* at a dosage of 18 ml/porous nail; P2 = *F. solani* at a dosage of 9
 161 ml/porous nail

162 Subsequently, based on the data in Figure 1, statistical analysis was done to test the effect of
 163 inoculant dosage on the amount of sesquiterpenes. The result of the normality test (Figure 2)
 164 shows p-value is less than 0.05, implying that the data is not normally distributed and the Kruskal-
 165 Wallis test is required. Meanwhile, the result of the Kruskal-Wallis test shows p-value is less than
 166 0.05, indicating the differences between the inoculant dosages on the percentage of sesquiterpenes
 167 in the agarwood are statistically significant (Figure 3). *F. solani* at a dosage of 9 ml/porous nail
 168 shows the best yield with a Mean Rank of 16 (Figure 3).

169

170

171

Tests of Normality							
Senyawa	Dosis	Kolmogorov-Smirnov ^a			Shapiro-Wilk		
		Statistic	df	Sig.	Statistic	df	Sig.
	P0	.513	8	<.001	.418	8	<.001
	P1	.325	8	.013	.665	8	<.001
	P2	.455	8	<.001	.566	8	<.001

a. Lilliefors Significance Correction

172

173 **Figure 2.** Normality test on the percentage of sesquiterpenes in *G. versteegii* agarwood inoculated
 174 using *simpori* technique with various dosages of inoculant

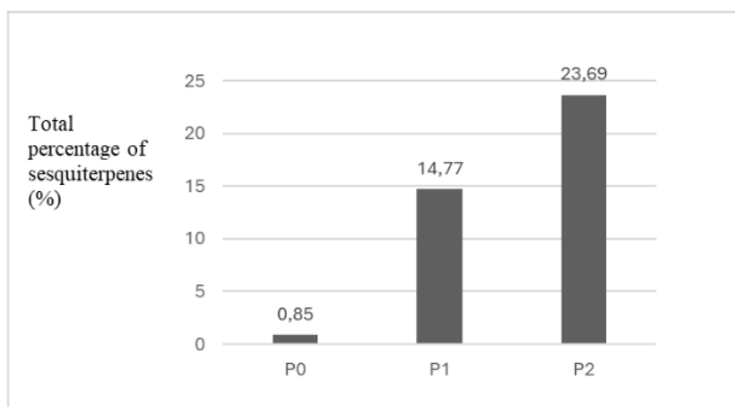
175 *P0 = *F. solani* at a dosage of 3 ml/porous nail; P1 = *F. solani* at a dosage of 18 ml/porous nail; P2 = *F. solani* at a dosage of 9
 176 ml/porous nail

177

Test Statistics ^{a,b}		Ranks		
Senyawa		Dosis	N	Mean Rank
Kruskal-Wallis H	6.112			
df	2			
Asymp. Sig.	.047			
a. Kruskal Wallis Test				
b. Grouping Variable: Dosis				
		P0	8	8.50
		P1	8	13.00
		P2	8	16.00
		Total	24	

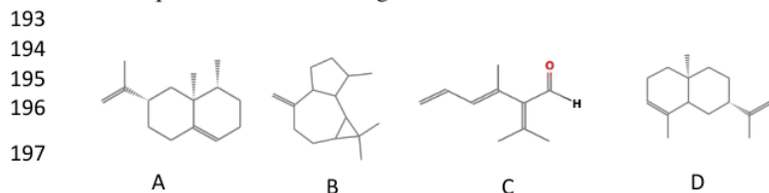
178
179 **Figure 3.** Kruskal-Wallis test on the percentage of sesquiterpenes in *G. versteegii* agarwood
180 inoculated using *simpori* technique with various dosages of inoculant
181 *P0 = *F. solani* at a dosage of 3 ml/porous nail; P1 = *F. solani* at a dosage of 18 ml/porous nail; P2 = *F. solani* at a dosage of 9
182 ml/porous nail

183 The total percentage of sesquiterpenes in agarwood for each treatment is presented in Figure
184 4.
185



186
187 **Figure 4.** Total percentage of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori*
188 technique with various dosages of inoculant
189 *P0 = *F. solani* at a dosage of 3 ml/porous nail; P1 = *F. solani* at a dosage of 18 ml/porous nail; P2 = *F. solani* at a dosage of 9
190 ml/porous nail

191 The main structure of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori*
192 technique is illustrated in Figure 5.



198
199 Source: <https://webbook.nist.gov/>

200 **Figure 5.** The main structure of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori*
201 technique
202 *A: Eremophilene, B: Aromadendrene, C: 2-Isopropylidene-3-methylhexa-3,5-dienal, D: α -Selinene

DISCUSSION (12 pt)

Based on the analysis, the types of sesquiterpene in n-hexane extracts of *G. versteegii* agarwood are diverse as illustrated in Table 1. The use of n-hexane as a solvent has been done previously, showing that sesquiterpenes can be obtained from extraction using nonpolar compounds (Ahmaed & Kulkarni, 2017). The most types of sesquiterpene were found in agarwood inoculated using *simpori* technique at an inoculant dosage of 9 ml/porous nail, namely six types. Subsequently, four types and one type of sesquiterpene were identified in those inoculated with inoculant dosage of 18 ml/porous nail and of 3 ml/porous nail, respectively. Sesquiterpenes indicate the quality of agarwood. The more types of sesquiterpene identified in the agarwood, the higher the quality of the agarwood.

In this study, inoculation using *simpori* technique with an inoculant dosage of 9 ml/porous nail was able to produce aromadendrene; α -Selinene; (1aS,4aR)-4a,8,8-Trimethyl-2-methylene-1,1a,2,4a,5,6,7,8-octahydrocyclopropa[d]naphthalene; 6-Methyl-2-(4-methylcyclohex-3-en-1-yl)hepta-1,5-dien-4-ol; Alloaromadendrene; and Eremophilene. These types of sesquiterpene also affect the aroma of the produced agarwood (Yang et al., 2021). Eremophilene, guaiane, and eudesmane are sesquiterpenes reported to be the main contributors to the aroma of agarwood (Hou et al., 2024). Eudesmane is mainly characterized by fresh, honey, sweet floral, and mint fragrance. The eremophilene is characterized by woody, camphor, amber and other persistent strong aromas. Meanwhile, guaiane is characterized by a slight woody camphor aroma and a powerful honey aroma. Another type of sesquiterpenes, agarospirane, is found in *Aquilaria agallocha* and characterized by a spicy, peppery, woody aroma (Yang et al., 2021).

The percentage of sesquiterpenes is presented in Figure 1. It shows that the eremophilene-type has the highest percentage (10.22%) and is only identified in agarwood inoculated using *simpori* technique with an inoculant dose of 9 ml/porous nail. This type is found in natural agarwood from *Gyrinops salicifolia* from Papua New Guinea (Shao et al., 2016). The second highest percentage is the aromadendrene-type. It is a member of the *Guaianes* group frequently found in *Aquilaria crassna* agarwood (Gao et al., 2019). Moreover, it is proposed to be a chemical compound that marks high-quality agarwood because its presence is always detected in high-quality agarwood (Sundaraj et al., 2023). In the present study, aromadendrene was identified in agarwood from all treatments and the highest percentage (6.92%) was obtained in agarwood inoculated using *simpori* technique with an inoculant dose of 18ml/porous nail.

The total percentage of sesquiterpenes in *G. versteegii* agarwood inoculated using *simpori* technique with various inoculant dosages is presented in Figure 4. Based on the figure, the highest total percentage (23.69%) of sesquiterpenes was obtained from that inoculated with an inoculant dosage of 9ml/porous nail. It confirms the findings shown in Table 1 that *simpori* technique with an inoculant dose of 9ml/porous nail produces agarwood with more types of sesquiterpene than other dosages.

Several factors that potentially affect the amount of sesquiterpenes in agarwood are the degree of environmental exposure to sapwood, and volatilization and natural decomposition in trees, and the duration of injury/exposure to microorganisms (Hou et al., 2024). In the present study, all treatments underwent exposure to microorganisms with the same duration (18 months) and under the same environment (one layer). In the present study, the difference in the content of sesquiterpenes was mainly caused by the difference in the inoculant (microorganism) dosage that stimulated the formation of agarwood. A dosage of 9ml/porous nail produced agarwood with the highest number of types and percentage of sesquiterpene. This phenomenon was reinforced through statistical test using the Kruskal-Wallis test (Figure 3), p-value of less than <0.05 (p-value = 0.047), indicating that the administration of different dosages of inoculant produced a significantly different percentage of sesquiterpene. The dosage of 9ml/porous nail produced the highest sesquiterpene with a Mean Rank of 16 (Figure 3).

The chemical structures of aromadendrene, 2-Isopropylidene-3-methylhexa-3,5-dienal, eremophilene, and α -Selinene are shown in Figure 5. Aromadendrene is structurally characterized by a dimethyl cyclopropane ring fused to a hydroazulene skeleton (Lamers, 2003). Aromadendrene

has several functionalities that can be used as a handle for synthetic transformations (Lamers, 2003). Eremophilene is a C₁₅H₂₄ hydrocarbon first isolated from *Petasites officinalis* and *P. albus* (Piers & Keziere, 1969). The structure of eremophilene comes from 1,2,3,4,4a,5,6,7-octahydronaphthalene which is substituted by an isopropenyl group at position 3 and by methyl groups at positions 4a and 5 (the 3R,4aR,5S-diastereoisomer). Subsequently, α -Selinene with double bond in the octahydronaphthalene ring system is endocyclic (2R,4aR,8aR)-configuration while 2-Isopropylidene-3-methylhexa-3,5-dienal has a molecular formula of C₁₀H₁₄O.

CONCLUSION AND SUGGESTION (12 pt)

The administration of different dosages of inoculants, namely 3 ml, 9 ml, and 18 ml/porous nail has been investigated, showing the significant difference to the results of sesquiterpene content with p-value of 0.047. *F. solani* inoculant at a dosage of 9 ml/porous nail obtains the best result with a Mean Rank of 16 and also the highest amount of sesquiterpene (23.69%). The sesquiterpenes identified in agarwood inoculated with a dosage of 9ml/porous nail include aromadendrene; α -Selinene; (1S,4aS,8aR)-4a,8,8-Trimethyl-2-methylene-1,1a,2,4a,5,6,7,8-octahydrocyclopropa[d]naphthalene; 6-Methyl-2-(4-methylcyclohex-3-en-1-yl)hepta-1,5-dien-4-ol; alloaromadendrene; and eremophilene. The treatment of an inoculant dosage of 9ml/porous nail can be recommended for optimizing agarwood production.

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