Research Article

Remarkable Phytochemical Characteristics of Chi-Nan Agarwood Induced from New-Found Chi-Nan Germplasm of *Aquilaria sinensis* Compared with Ordinary Agarwood

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Wild Chi-Nan agarwood is regarded as the highest quality agarwood from *Aquilaria* spp. However, the comprehensive research on chemical composition of wild Chi-Nan agarwood is limited. An integrated strategy using SHS-GC-MS and UPLC-Q/ToF-MS was applied to explore the phytochemical characteristics of a kind of agarwood induced from a newly identified germplasm of Chi-Nan *A. sinensis*. Progenesis QI and MS-Dial were used to preprocess the UPLC-Q/ToF-MS and GC-MS raw data, respectively. Principle component analysis (PCA) and orthogonal partial least squares to latent structure-discriminant analysis (OPLS-DA) models were built to discriminate Chi-Nan agarwood from ordinary agarwood and to screen potential distinguishing components between them. In this study, we clarified the distinguishing differences between Chi-Nan agarwood and ordinary agarwood. The difference is mainly manifested in the average contents of 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxybenzene)ethyl] chromone, which are 170 and 420 times higher in Chi-Nan agarwood than in ordinary agarwood, respectively, while the contents of 5,6,7,8-diepoxy-2-(2-phenylethyl)chromones (DEPECs), 5,6-epoxy-2-(2-phenylethyl)chromones (EPECs), and 5,6,7,8-tetrahydro-2-(2-phenylethyl)chromones (THPECs) such as agarotetrol are extremely low. The content of the main sesquiterpenes in Chi-Nan agarwood was higher than that in ordinary agarwood, especially in regard to guaiane and eudesmane derivatives. In addition, there were significant differences in the contents of low-molecular-weight aromatic compounds such as 2-methyl-4H-1-benzopyran-4-one, 4-methoxybenzaldehyde, and 2-hydroxybenzaldehyde between Chi-Nan agarwood and ordinary agarwood. All the mentioned main chemical characteristics of this new Chi-Nan agarwood were coincident with those of the rare wild Chi-Nan agarwood from *A. malaccensis*, *A. sinensis*, and *A. crassna*. We reported differences in 2-(2-phenylethyl)chromones, sesquiterpenes, and low-molecular-weight aromatic compounds between Chi-Nan agarwood and ordinary agarwood from *A. sinensis* for the first time; it is necessary to evaluate the agarwood from the new-found Chi-Nan germplasm.

1. Introduction

Agarwood is resinous wood obtained from wounded *Aquilaria* tree, which is a genus belonging taxonomically to the Thymelaeaceae family [1]. The main constituents in agarwood are volatile constituents and semi-volatile components [2]. The former include low-molecular-weight aromatic compounds and sesquiterpene derivatives, and the
latter principally consist of 2-(2-phenylethyl)chromone derivatives [3]. Sesquiterpenes, including agarofurans, cadinanes, eudesmanes, eremophilanes, guianes, and agarospiranes, are considered to be the prominent contributors to agarwood aroma [4]. 2-(2-phenylethyl)chromone monomers can be divided into four categories based on the A ring of 2-(2-phenylethyl)chromones, namely, 5,6,7,8-tetrahydro-2-(2-phenylethyl)chromones (THPECs), diepoxyl-tetrahydro-2-(2-phenylethyl)chromones (DEPECs), epoxy-tetrahydro-2-(2-phenylethyl)chromones (EPECs), and flindersia type 2-(2-phenethyl)chromones (FTPECs) [2, 5].

Chi-Nan agarwood (CNA), a high-quality agarwood, is also called Qi-Nan or Jar-Nan in China, Kanankoh or Kyara in Japan, and Tagara in India [6]. In China, wild CNA is considered representative of high-quality agarwood, whose price is increasing up to thousands of RMB yuan per gram [7, 8]. Wild Chi-Nan agarwood is valued for its mysterious and elegant oriental odour that could be obviously smelt without heating, which make it discriminate from other kinds of agarwood. Investigations on wild CNA have rarely been reported, in contrast to abundant reports related to ordinary agarwood. In 1985, Hashimoto proved that benzaldehyde and 4-methoxybenzaldehyde originated from the pyrolysis of 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxyphenyl)ethyl]chromone, respectively, in the neutral part of Kyara [9]. Ishihara discovered that the relative contents of 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxyphenyl)ethyl]chromone from Kanankoh (A.agallocha, synonym: A. malaccensis) [10] smoke were much higher than those from Jinkoh [11]. In addition, taking four types of wild CNA (A. sinensis and A. agallocha) as materials, Dai showed that there were abundant 2-(2-phenylethyl)chromones and 2-[2-(4′-methoxyphenyl)ethyl]chromones [6, 12]. It is not difficult to notice that previous studies took only the 2-(2-phenylethyl)chromones into account. In addition, Chinese pharmacologist Xie Zongwan described the morphological characteristics of CNA but failed to provide information on the original plant [13]. Due to the confusion regarding the original plant, relevant information on wild CNA has seldom been described in the mentioned studies.

In recent years, a kind of special agarwood germplasm of A. sinensis has been introduced and domesticated from the wild population and propagated by grafting with ordinary germplasm of A. sinensis. Agarwood with more than 40% of the alcohol soluble extractive content can be obtained after this kind of tree is wounded with the drilling method over one year. However, for ordinary germplasm of A. sinensis, it is almost impossible to obtain agarwood with more than 10% of the alcohol soluble extractive content with the same inducing method [14]. Because of the similarity of characteristics such as appearance, form, texture, color, smell, and taste to wild CNA, this new-found agarwood is also called CNA. To date, there has been only one report by Li [15] in which the chemical properties of this kind of CNA have been tested by TG-FTIR and HS-GC-MS, and these authors named it Kynam. However, the authors did not clarify the specific components to distinguish CNA in terms of fragrance and the chemical characteristics of CNA. Here, gas chromatography-mass spectrometry (GC-MS) and liquid chromatography-mass spectrometry (LC-MS) were used to analyze the differences between CNA induced from new-found germplasm and ordinary agarwood (OA) to obtain a deep knowledge of phytochemical characteristics.

2. Materials and Methods

2.1. Agarwood Materials. Thirteen agarwood samples were analyzed as shown in Table 1 and Figure S1. They are divided into CNA (Chi-Nan agarwood) and OA (ordinary agarwood) group. The seven CNA samples were collected from new-introduced germplasms A. sinensis tree that were provided by three farmers from different planting bases of Hainan and Guangdong province in China. As for the OA group, we chose six ordinary agarwood induced with three kinds of common agarwood-inducing methods, including whole-tree agarwood-inducing technique (Agar-Wit) [16], burning-chisel-drilling (BCD), and wild.

2.2. Chemical Reagent. Al2O3 powder as dispersant (High purity, Tianjin Guangfu Fine Chemical Research Institute, China) and retention index marker alkanes (AccuStandard, USA, C9–C40) were used in GC-MS, as well as anhydrous alcohol (AR, Xilong Science Co., Ltd, China), and acetonitrile (Merck HPLC grade, Darmstadt, Germany).

2.3. Sample Preparation. For all the samples described in Section 1, agarwood material sections, chips, or blocks were ground into powder with liquid nitrogen. Agarwood powder (0.2 g) was extracted with 10 mL of 50% ethanol by means of sonication at room temperature for 30 min. Then, the supernatant was used as the sample reserve solution. Through preexperiments, we found that the content of 2-(2-phenylethyl)chromones and 2-[2-(4′-methoxyphenyl)ethyl]chromones in CNA was too high to analyze, so we diluted the test solution 10 times before analysis.

Agarwood powder was put into a 20 ml headspace bottle, 0.2 g Al2O3 powder was added as the dispersant [17], and the solution was analyzed in the headspace sampler according to the following conditions: headspace heating temperature: 180°C; transmission line temperature: 185°C; injection probe temperature: 190°C; headspace heating time: 40 min; and N2 purging time: 30 s. For each of the samples, 1 mL of headspace gas was used for GC-MS analysis. The retention index marker alkanes (C9–C40) were analyzed by the same program in GC-MS.

2.4. LC-Q/ToF-MS Analysis Condition. LC-MS was performed using an ultra-high-performance liquid chromatography (UPLC) system (H-class, Waters, USA) coupled with a quadrupole time-of-flight tandem spectrometer (Xevo G2- XS, Waters, USA). Separation was performed using a Waters Acquity UPLC BEH C18 column (3.0 mm x 100 mm, 1.7 μm, Waters, USA). The column temperature was 30°C. The detection wavelength was set at 196 nm for all the tested compounds, and the mobile phases were acetonitrile (A) and water (B). A gradient elution was used: 0 min, 20% (v/v) A; 20 min, 50% (v/v) A; 30 min, 20% (v/v) A. The mobile phase
was established at a flow rate of 0.2 mL·min\(^{-1}\), and the injection volume was 5 \(\mu\)L.

The nebulization gas was set to 600 L·h\(^{-1}\) at a temperature of 350°C, and the cone gas was set to 50 L·h\(^{-1}\). The source temperature was set to 110°C. The capillary voltage and cone voltage were set to 3500 V and 30 V, respectively. The data acquisition rate was set to 0.3 s with a 0.1 s interscan delay. Data between \(m/z\) 50 and 1200 were recorded in positive ion mode. The quality axis was corrected by sodium formate, and the quality of leucine enkephalin was corrected in real time.

### Table 1: Sample information.

| Num. | Species | Agarwood induction method | Place of production | Description | Sinkage |
|------|---------|----------------------------|---------------------|-------------|---------|
| CNA1 | *A. sinensis* | 3-year-old Chi-Nan germplasm by drilling for 15 months | Ding'an, Hainan province | Irregular pieces, black brown resin bands alternate with yellow white wood stripes, sufficient resin, rich of aromas, acid in taste, soft and glutinous | x |
| CNA2 | *A. sinensis* | 3-year-old Chi-Nan germplasm by drilling for 14 months | Ding'an, Hainan province | Irregular strips, brown resin bands and white wood are distributed alternately, strong fragrance, cool feeling, bitter in taste, hard texture, slightly sticky | x |
| CNA3 | *A. sinensis* | 5-year-old Chi-Nan germplasm by drilling for 18 months | Ding'an, Hainan province | Irregular pieces, black brown resin scatter like spots, adequate resin, the aroma is thick, numb the tongue, hard texture, sticky | √ |
| CNA4 | *A. sinensis* | germplasm by drilling for 12 months | Maoming, Guangdong province | Irregular strips, apparent brown resin spread throughout the surface, intense aroma, taste peppery, hard texture, sticky | √ |
| CNA5 | *A. sinensis* | 3-year-old Chi-Nan germplasm by drilling for 12 months | Maoming, Guangdong province | Irregular pieces, black brown resin scatter like spots, saturated with resin, fragrance is elegant, spicy and numb, tough, sticky | x |
| CNA6 | *A. sinensis* | germplasm by drilling for 18 months | Maoming, Guangdong province | Irregular pieces, black brown resin scatter like spots or stripe, quite strong aroma, little spicy and numb, tough, sticky | √ |
| CNA7 | *A. sinensis* | germplasm by drilling for 12 months | Maoming, Guangdong province | Irregular pieces, obvious black brown resin scatter like spots or stripe, cool feeling, taste peppery and numb, tough, soft and glutinous | √ |
| OA1 | *A. sinensis* | 6-year-old trees induced by Agar-Wit\(^a\) for 18 months | Haikou, Hainan province | Irregular slices, brown resin bands alternate with yellow white wood stripe, pleasant fragrance, crisp | x |
| OA2 | *A. sinensis* | 6-year-old trees induced by Agar-Wit for 8 months | Danzhou, Hainan province | Irregular thin slices, saturated with resin brown resin, sweet fragrance, crisp | x |
| OA3 | *A. sinensis* | Wild agarwood | Hainan province | Irregular pieces, massive protrusions and patches distribute throughout the appearance, slight aroma, soft | x |
| OA4 | *A. sinensis* | Wild agarwood | Hainan province | Irregular pieces, many protrusions and patches distribute throughout the appearance, tawny resin scatter like spots, slight aroma, crisp | x |
| OA5 | *A. sinensis* | 5-year-old trees induced by BCD\(^b\) for 12 months | Maoming, Guangdong province | Irregular pieces or slices, tawny resin and white wood are distributed alternately, many fibers in the cross section, slight aroma, resilient | x |
| OA6 | *A. sinensis* | 5-year-old trees induced by BCD for about 12 months | Maoming, Guangdong province | Irregular pieces, brown resin scatter like spots, cheerful, aroma, crisp | x |

\(^a\)The abbreviation of whole-tree agarwood-inducing technique is Agar-Wit. \(^b\)The abbreviation of burning-chisel-drilling is BCD. "\(\sqrt{\}\)"; the agarwood sample could sink in water. "x": the agarwood sample could not sink in water.

was high purity helium (99.9995%) was used as the carrier gas at a flow rate of 1 mL·min\(^{-1}\). The column temperature program was set as follows: 1 min at the initial temperature of 50°C, which was subsequently ramped to 143°C at a rate of 15°C·min\(^{-1}\); 10 min at 143°C, which was ramped to 155°C at a rate of 1°C·min\(^{-1}\); 0 min at 155°C, which was ramped to 225°C at a rate of 25°C·min\(^{-1}\); 7 min at 225°C, which was ramped to 300°C at a rate of 25°C·min\(^{-1}\); then the temperature was held at 300°C for 5 min. The ratio of gas flowing out of the chromatographic system and gas flowing into the chromatographic column after sample gasification was 5:1.

### 2.5. SHS-GC-MS Analysis Conditions.

GC-MS analysis was performed with a TSQ GC system (Agilent Technologies, USA). The chromatographic separation was conducted with a DB-5MS capillary column (30 m \(\times\) 0.25 mm i.d., 0.25 mm film thickness, Agilent Technologies, USA). The GC/MS interface temperature was maintained at 250°C. Ultrahigh purity helium (99.9995%) was used as the carrier gas at a flow rate of 1 mL·min\(^{-1}\). The column temperature program was set as follows: 1 min at the initial temperature of 50°C, which was subsequently ramped to 143°C at a rate of 15°C·min\(^{-1}\); 10 min at 143°C, which was ramped to 155°C at a rate of 1°C·min\(^{-1}\); 0 min at 155°C, which was ramped to 225°C at a rate of 25°C·min\(^{-1}\); 7 min at 225°C, which was ramped to 300°C at a rate of 25°C·min\(^{-1}\); then the temperature was held at 300°C for 5 min. The ratio of gas flowing out of the chromatographic system and gas flowing into the chromatographic column after sample gasification was 5:1.

### 2.6. Data Preprocessing and Statistical Analysis.

UPLC-Q/ToF-MS raw data were imported into Progenesis QI (Waters, USA) for preprocessing, while GC-MS raw data were converted and preprocessed by MS-Dial (NSF-JST, Japan). Statistical analysis was carried out by SIMCA-P.
(version 14.1, Umetrics, Umea, Sweden) on the basis of data matrixes. Unsupervised principal component analysis (PCA) was used to identify similarities or latent differences between groups. Data plotted using PC revealed intersample relationships via their spatial proximity. Orthogonal projection to latent structure discriminant analysis (OPLS-DA) was then implemented to detect maximum information from the data set and to distinguish the components induced by different groups. Distinguished compounds were screened by analyzing the Variable Importance in the Projection (VIP) and S-plot. Finally, Student’s t-test (p < 0.05) was used to screen the significant variables with SPSS R26 (IBM, USA).

3. Results

3.1. Phytochemical Characteristics of CNA Based on LC-MS. Based on the high-resolution mass spectrometry and fragment ion information obtained in the experiment, thirty 2-(2-phenethyl)-chromones in thirteen batches of samples were identified according to the methods used in previous studies [18–20], as shown in Table S1. They were distributed into three parts of the total ion chromatograms in Figure 1. It is noteworthy that, between 0 and 10 min (zone a), the main components detected were THPECs, while EPECs/DEPECs flowed out from 10–15 min (zone b). The FTPECs and dimers flowed out after 15 min of the elution gradient (zone c). Historically, the composition of CNA is relatively simple, especially before 15 min, which indicates that FTPECs are the predominant type of 2-(2-phenethyl)chromones in CNA.

Thus, we carried out PCA to obtain an objective understanding of LC-MS data, and the results are shown in Figure 2(a). The two PCs described 72.9% variance, of which the first PC (PC1) accounted for 57.3% and PC2 accounted for 15.6% of the total variance. The CNA mainly gathered on the positive side of PC1, and the OA was distributed on the negative side of PC1 which proved that the two groups had obvious differences in the composition of 2-(2-phenethyl)chromones. Then, supervised OPLS-DA was subsequently used. As depicted in Figure 2(b), the CNA samples are clearly separated from the OA samples [R2X (cum) = 0.727; R2Y (cum) = 0.977; Q2 (cum) = 0.952]. One orthogonal component was calculated. The validation plot obtained from 200 permutation tests confirmed the validity of this OPLS-DA model. The criteria for validity include the following: all the permuted R2 and Q2 values to the left are lower than the original points to the right, and the regression line of the Q2 points intersects the vertical axis (on the left) at or below zero.

3.2. Phytochemical Characteristics of CNA Based on GC-MS. To obtain a chemical profile of the violate constituents of agarwood, an analytical method based on SHS-GC-MS was developed. The TIC chromatographs of representative samples (OA1 and CNA1) and the plot corresponding to the temperature program are shown in Figure 3. With the application of Agilent MassHunter Quantitative Analysis 10.0, 101 compounds were identified by searching NIST14 in the MS data and retention index. Tentative identification with similarity over 80% and retention index within ±20 is listed in Table S2. Figure 3 shows that low-molecular-weight aromatic compounds, sesquiterpenes, and chromones flow out in order during different temperatures. Among them, mainly low-molecular-weight aromatic compounds were detected when the temperature was above 50°C (zone a). When the temperature ranges from 142°C to 225°C (zone b), sesquiterpenes and their thermal cracking fragments flow out of the column. Finally, low-polarity 2-(2-phenethyl)-chromones can be detected when the temperature rises up to 225°C until 38.6 min (zone c).

Mutual projections of factor scores for the first two PCs are presented in Figure 4(a), and these scores described 35.8% (PC1) and 16.7% (PC2) of the variability in the data when ellipse hoteling was set as 95% (70.1% can be explained by the four components). Thirteen batches of agarwood samples could be readily divided into two different groups, which indicated that the content and distribution of components were different between CNA and OA. The CNA samples were clustered closer to each other than to the OA samples on the negative side of PC1, which indicates that the differences between the groups are relatively minor.

To identify distinguishing compounds, the data were further processed with orthogonal partial least squared discriminant analysis (OPLS-DA). The classification results are shown in Figure 4(b). The CNA group could be clearly separated from the OA group. The values of R2X (cum) and R2Y (cum) are 0.628 and 1, respectively, which is an important parameter to show how much data contributed to creating the model. In addition, Q2 (cum) was 0.956, indicating that the OPLS-DA model had high predictability. To guard against model overfitting, permutation tests with 200 iterations were carried out.

Furthermore, we analyzed the distribution of different types of sesquiterpenes in the samples. Finally, a total of seventeen types of sesquiterpenes were identified in thirteen batches of samples. Among them, seven predominant types of sesquiterpenes, including α-santalanes, agarspirananes, guiananes, eremophilanes, eudesmanes, cadinanes, and agarofurans, were used for successive analysis. Following the logarithmic conversion of the total peak area of various sesquiterpenes, Figure 5 reflects the distribution of different configurations of sesquiterpenes between CNA and OA. The stars on the boxes represent mean values, and the results indicate that CNA possesses a higher concentration of all sesquiterpenes than OA. The length of the box, which indicates the interquartile range, clearly shows that CNA possesses greater dispersion. Obviously, eudesmane derivatives are most abundant in CNA, while eremophilane derivatives comprise the majority of sesquiterpenes in OA. Comparatively, the concentrations of guiananes and eudesmanes were significantly different (p < 0.05) between the two groups.

3.3. The Distinguishing Components between CNA and OA. To determine the differences in the composition of 2-(2-phenethyl)-chromones, 13 potential distinguishing
components were identified, as shown in Table 2. The S-plot derived from the OPLS-DA model comparing the two groups is shown in Figure 6(a). The VIP value reflects the influence of every compound’s ion on the classification. Variables with a VIP value > 1 have an above-average influence on the explanation of the Y matrix. In our research, we adjusted VIP up to 5 to evaluate significant variables according to the practical situation. The qualified variables that meet the above criteria are the best possible distinguishing components labeled by a gray square. Subsequently, these components were used for Student’s t-test.

Table 2 shows seven identified significant components, including two DEPECs, two THPECs, and three FTPECs with higher contents in OA; these components are highlighted with blue stars. The p values of compounds 2045, 3225, and 5685 were less than 0.01, which means that the differences in these constituents were significant. Likewise, all 6 identified significant components were FTPECs and dimeric PECs in CNA. Moreover, all the constituents’ p values were below 0.001. It should be noted that the average peak areas of 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxybenzene)ethyl]chromone are 170 and 420 times higher in CNA than in OA, respectively, as shown in Table S2. We applied the same method to select the distinguishing volatile constituents from agarwood. According to the specific conditions of this experimental variable, the VIP threshold was adjusted to 4. Based on the above conditions, the compounds in the rectangular frame, as shown in Figure 6(b), are highlighted. Student’s t-test was performed in succession, and variables without significant differences between the two groups (p > 0.05) were eliminated. The remaining compounds (marked with stars in Figure 6(b)) were selected for identification. Detailed information on the
twelve distinguishing components is listed in Table 3. The most noteworthy differences between CNA and OA are in low-weight aromatic compounds and 2-(2-phenylethyl)chromones. Regarding low-weight aromatic compounds, compounds \(53, 64, 180, 334, \) and \(570\) were abundant in OA, while compounds \(223, 286, 304, 361, \) and \(614\) were abundant in CNA. Remarkably, the \(p\) values of 2-(2-phenylethyl)-chromone and 2-[2-(4'-methoxyphenyl)ethyl]-chromone were below 0.001, which indicates that the contents of these two compounds are significantly different between CNA and OA.

4. Discussion

4.1. The Differences in 2-(2-Phenylethyl)chromones between CNA and OA. As the results demonstrate, the 2-(2-phenylethyl)chromones in CNA induced from new-found Chi-Nan germplasm are relatively simple and all belong to FTPECs. Noticeably, the contents of 2-(2-phenylethyl)chromone and 2-[2-(4'-methoxybenzene)ethyl]-chromone are hundreds of times higher in CNA than in OA. Additionally, Japanese researchers discovered that the relative content of two constituents accounted for 5.83% and 1.59%

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**Figure 3:** The total ion chromatograms of typical agarwood samples (OA1 and CNA1) acquired by SHS-GC-MS and the corresponding temperature gradient variety diagram (a zone: 0–10.6 min; b zone: 10.6–34.0 min; c zone: 34.0–45 min).

**Figure 4:** PCA (a) and OPLS-DA (b) scores plot for the first two components of Chi-Nan agarwood (filled triangle) and ordinary agarwood (filled square) data analyzed by GC-MS.
in Kanankoh (wild CNA), while they accounted for only 0.28% and below 0.05% in Jinkoh (OA) [11]. Dai reported that the relative contents of 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxyphenyl)ethyl]-chromone were extremely high in the ether extract of four kinds of wild CNA. Therefore, we considered 2-(2-phenylethyl)-chromone and 2-[2-(4′-methoxyphenyl)ethyl]-chromone as phytochemical characteristic constituents to distinguish CNA. Such characteristics can also account for the high alcohol soluble extractive content, as described in previous studies [15].

However, some 2-(2-phenylethyl)chromones, such as DEPECs, EPECs, and THPECs, exist in agarwood apart from FTPECs, especially THPECs, which are specific compounds that have so far been detected only in agarwood [21]. We found that the amount and content of THPECs and DEPECs in OA, such as compounds 2045, 4531, 3225, and 4729, especially agarotetrol (2045), were higher. Agarotetrol has been designated as an index component in the quality control of content determination in medicinal agarwood according to the Chinese Pharmacopoeia 2020. In contrast, the content of THPECs in CNA was extremely low, especially in agarotetrol. Thus, CNA fails to reach the requirement of the Chinese Pharmacopoeia 2020 on the basis of the content of agarotetrol.

4.2. Distinguishing Differences in Sesquiterpenes and Low-Weight Aromatic Compounds between CNA and OA. Agarwood is processed into perfumes and incenses in addition to being used as a raw material in traditional and modern medicines. Sesquiterpene volatile constituents and low-weight aromatic compounds are the major volatile constituents in agarwood [2]. Therefore, SHS was used as the pretreatment method to analyze the volatile components of agarwood to replicate the fumigation. In our results, 69 sesquiterpenes were detected and identified from CNA and 43 sesquiterpenes from OA. However, it is noteworthy that the contents of guaianes and eudesmanes derivatives in CNA were superior to those in OA. It has been reported that guaiane and eudesmane derivatives accounted for the majority of sesquiterpenes in wild CNA according to investigations of Ishihara et al. [22] and Yang et al. [23]. Even more importantly, many of them were reported to possess a pleasant fragrance [2, 12]. Guaianes and eudesmanes seem to contribute to the gorgeous and elegant character of CNA by contributing to their odoriferous properties [18]. Consequently, guaianes and eudesmanes may be significant contributors to the unique fragrance of CNA. Of course, we need to carry out relevant studies to specify the characteristics in terms of sesquiterpenes so that we can explain the fragrance of the CNA more perspicuously.

It is meaningful that we found that several low-weight aromatic compounds are similar to the partial structure of FTPECs, such as 4-methoxybenzene-methanol (361), 2-hydroxybenzaldehyde (223), and 1-(2-hydroxyphenyl) ethenone (286). Moreover, the contents of these compounds were higher than those in OA, so it is reasonable for us to speculate that the above low-weight aromatic compounds partly originate from the pyrolysis of 2-(2-phenylethyl)-chromone and 2-[2-(4′-methoxybenzene)ethyl]-chromone. Similarly, the high content of 4-methoxybenzaldehyde (304) in CNA has the same explanation. In fact, Hashimoto found that 2-(2-phenylethyl)chromone and 2-[2-(4′-methoxypyphenyl)ethyl]chromone could pyrolyze at 150°C for 6 h to produce benzaldehyde and 4-methoxybenzaldehyde, respectively [9]. It may be difficult to understand the origin of abundant benzaldehyde in OA. However, Takamatsu proved that agarotetrol could produce pyrolys products such as 4-phenyl-2-butanone and benzaldehyde with HS-SPME-GC-MS analysis when heated to 190°C–200°C [19]. Ishihara et al. found that smoke from agarwood contained a small amount of pulp wood pyrolysis products such as acetic acid, benzaldehyde, and vanillin when Kanankoh (CNA in Japanese) was heated at 180–210°C by an alcohol lamp [11]. Therefore, we can hypothesize that the non-resinous part and agarotetrol in OA results in high concentrations of benzaldehyde (180) and acetic acid (53). In addition, we discovered that the content of 4-phenyl-2-butanone (334) in OA is nearly 1.5 times as high as that in CNA by calculating the peak area shown in Table S2. This is probably related to the biosynthesis of 2-(2-phenylethyl)chromones according to Liao’s research [20]. In conclusion, there were considerable discrepancies in low-weight aromatic compounds between CNA and OA.

4.3. The Phytochemical Characteristics of New-Found CNA Are Consistent with Those of Wild CNA. Considering the chemical composition and application characteristics of agarwood, we comprehensively explored the phytochemical
characteristics of CNA induced from new-found Chi-Nan germplasm of *A. sinensis*. Comparatively, CNA and OA showed dramatic differences in chemical composition. Notably, the phytochemical characteristics of CNA were consistent with those of wild CNA reported by Chinese and Japanese researchers in terms of phytochemical characteristics. In addition, we analyzed a wild CNA sample (CNA8) under the same conditions, and the corresponding results are shown in Fig. S2-S3. Therefore, it is acceptable for us to speculate that CNA in this study is a kind of agarwood that is similar to wild CNA.

Table 2: Thirteen distinguished compounds between CNA and OA in terms of 2-(2-phenylethyl)chromones.

| Variable ID | Molecular formula | p value | Group | Types | Proposed compound |
|-------------|-------------------|---------|-------|-------|-------------------|
| 6455        | C_{19}H_{18}O_{4} | 0.014   | OA    | FTPECs | 1 OCH_{3} 1 OCH_{3} |
| 4531        | C_{18}H_{16}O_{5} | 0.024   | OA    | DEPECs | 1 OCH_{3} |
| 6289        | C_{20}H_{20}O_{5} | 0.028   | OA    | FTPECs | 2 OCH_{3} 1 OCH_{3} |
| 5685        | C_{19}H_{18}O_{6} | 0.001   | OA    | FTPECs | 1 OCH_{3} 1 OH, 1 OCH_{3} |
| 3225        | C_{20}H_{20}O_{6} | 0.005   | OA    | THPECs | 2 OH 1 OH, 1 OCH_{3} |
| 4729        | C_{18}H_{18}O_{4} | 0.034   | OA    | DEPECs | 2 O- |
| 2045        | C_{18}H_{16}O_{6} | 0.005   | OA    | THPECs | Agarotetrol |
| 6964        | C_{11}H_{14}O_{3} | ***     | CNA   | FTPECs | 1 OCH_{3} |
| 7234        | C_{12}H_{14}O_{2} | ***     | CNA   | FTPECs | 2-(2-Phenylethyl)chromone |
| 5255        | C_{14}H_{16}O_{4} | ***     | CNA   | FTPECs | 1 OH, 1 OCH_{3} |
| 6936        | C_{13}H_{14}O_{3} | ***     | CNA   | Dimeric-PECs |
| 7180        | C_{18}H_{18}O_{4} | ***     | CNA   | Dimeric-PECs |
| 5072        | C_{18}H_{18}O_{4} | ***     | CNA   | FTPECs | 1 OH, 1 OCH_{3} |

***indicates a significant difference between two groups ($p < 0.001$).

Figure 6: S-plot at the first component used in potential distinguished components selection based on LC-Q/ToF-MS (a) and GC-MS (b), constituent ions with VIP value > 4 were marked with a black square, compounds marked with yellow stars are richer in CNA, and dark blue marked compounds are richer in OA.

Table 3: Twelve distinguished compounds between CNA and OA in terms of violate constituents.

| Peak | RI  | Molecular formula | p value | Group | Proposed compound |
|------|-----|-------------------|---------|-------|-------------------|
| 570  | 1504| C_{11}H_{14}O_{2} | ***     | OA    | 4-(4-methoxyphenyl)-2-butanone |
| 53   | 733 | C_{18}H_{16}O_{2} | 0.002   | OA    | acetic acid |
| 334  | 1248| C_{10}H_{12}O     | 0.022   | OA    | 4-phenyl-2-butanone |
| 64   | 750 | C_{18}H_{16}O_{2} | 0.008   | OA    | 1-hydroxy-2-propanone |
| 180  | 966 | C_{18}H_{16}O     | 0.047   | OA    | benzaldehyde |
| 614  | 1526| C_{18}H_{16}O_{2} | ***     | CNA   | 2-methyl-4H-1-benzopyran-4-one |
| 304  | 1207| C_{18}H_{16}O_{2} | 0.009   | CNA   | 4-methoxybenzaldehyde |
| 361  | 1290| C_{18}H_{16}O_{2} | ***     | CNA   | 4-methoxybenzenemethanol |
| 223  | 1051| C_{20}H_{20}O_{2} | ***     | CNA   | 2-hydroxybenzaldehyde |
| 286  | 1172| C_{18}H_{16}O_{2} | 0.001   | CNA   | 1-(2-hydroxyphenyl)ethanone |
| 1589 | 2618| C_{18}H_{16}O_{2} | ***     | CNA   | 2-(2-phenethyl)chromone |
| 1557 | 2352| C_{18}H_{16}O_{3} | ***     | CNA   | 2-[2-(4'-methoxyphenyl)ethyl]chromone |

***indicates a significant difference between two groups ($p < 0.001$).
originating from *A. sinensis* or *A. malaccensis*, which is reported by both foreign and domestic researchers [11, 12]. Thus, this is the first study to comprehensively reveal the distinguishing phytochemical characteristics of volatile and semi-volatile components of this newly identified CNA, and the findings indicated that CNA may originate from Chi-Nan germplasm of *Aquilaria* spp. rather than a unique environment, induction method, or species. We also identified the original plant of this CNA via DNA barcoding technology as *A. sinensis*. The relevant conclusions will be published in another paper.

5. Conclusion

We demonstrated the historical phytochemical characteristics of CNA for the first time. Here, we reveal distinguishing differences in terms of 2-(2-phenylethyl)chromones, sesquiterpenes, and low-molecular-weight aromatic compounds between CNA and OA. The main chemical characteristic of CNA is the single composition of 2-(2-phenylethyl)chromones, which is characterized by extremely high contents of 2-(2-phenylethyl)chromone and 2-[2-4’-methoxybenzene]ethyl chromone in CNA. Regarding the different types of 2-(2-phenylethyl)chromones, CNA mainly contains FTPECs, while the contents of DEPECs, EPECs, and THPECs, such as agarotetrol, are extremely low. The content of low-molecular-weight aromatic compounds in the smoke of agarwood detected by heating from the pyrolysis of FTPECs was higher than that in OA, especially for 4-phenyl-2-butanol, benzealdehyde, 4-methoxybenzaldehyde, and 2-hydroxybenzaldehyde. Nevertheless, the concentrations of acetic acid, benzealdehyde, and 4-phenyl-2-butanol were much higher in OA than in CNA. Remarkably, the guaiane and eudesmane derivatives with higher content in CNA were likely to be the crucial contributors to the unique fragrance of CNA when used as incense. Due to the significant differences between CNA and OA, it can be inferred that CNA may originate from Chi-Nan germplasm of *Aquilaria* spp., which can produce CNA in a short period of time without a special environment, place of origin, or method. The unique phytochemical characteristics of CNA may be related to the genetic information of the original plant germplasm or endophytic fungi. However, the biosynthesis mechanism of the abovementioned chemical characteristics needs further research for clarification.

**Abbreviations**

CNA: Chi-Nan agarwood  
OA: Ordinary agarwood  
SHS-GC-MS: Static headspace sampling gas chromatography mass spectrometry  
UPLC: Ultra-performance liquid chromatography  
Q-Tof-MS: Quadrupole time-of-flight mass spectrometry  
HRMS: High-resolution mass spectrometry  
PCA: Principal components analysis  
OPLS-DA: Orthogonal partial least squares discriminant analysis  
BCD: Burning-chisel-drilling  
Agar-Wit: Whole-tree agarwood-inducing technique  
PC: Principal component

PECs: 2-(2-phenylethyl)chromones  
THPECs: 5,6,7,8-tetrahydro-2-(2-phenylethyl)-chromones  
DEPECs: 5,6,7,8-diepoxy-tetrahydro-2-(2-phenylethyl)chromones  
EPECs: Epoxy-tetrahydro-2-(2-phenylethyl)chromones  
FTPECs: Flindersia type 2-(2-phenethyl)chromones.

**Data Availability**

All data included in this study are available upon request to the corresponding author.

**Conflicts of Interest**

The authors declare no conflicts of interest.

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**Supplementary Materials**

Tables S1–S2 and Figures S1–S3 in Supplementary Materials provide the details on comprehensive image analysis.

( Supplementary Materials)

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