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Improving the chemical profiling of complex natural extracts by joint ¹³C NMR and LC-HRMS² analysis and the querying of in silico generated chemical databases.

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The chemical profiling of complex natural mixtures emerges as a pivotal avenue of investigation for the discovery of new bioactive compounds. It requires a dereplication step generally based either on liquid chromatography-high resolution tandem mass spectrometry (LC-HRMS²) or on nuclear magnetic resonance (NMR) to quickly identify known compounds. The high sensitivity of MS results in numerous but sometimes incorrect candidate compounds, whereas the greater universality of NMR leads to fewer but more accurate annotations. These two analytical techniques are rarely used in combination despite their complementarity. This study focuses on the chemical profiling of Larix decidua (Pinaceae) bark by joint LC-HRMS² and ¹³C NMR data analysis and by querying custom in silico-generated chemical databases. MS-based dereplication allowed the annotation of 135 MS² spectra with at least two different annotation tools. Twenty-five compounds were annotated in parallel by NMR spectra analysis, including two previously undescribed myrtenic acid derivatives. Sixteen of these compounds were already reported in the Pinaceae family. Twelve compounds were jointly annotated with a high confidence level by comparing LC-HRMS² and ¹³C NMR dereplication results, including compounds not reported to date in Larix decidua. Our results show the benefits brought by combining LC-HRMS² and ¹³C NMR data and by querying custom in silico chemical databases to enhance the confidence level of data annotation during the chemical profiling of complex natural extracts.

Introduction

Living organisms produce low molecular weight organic natural products, called metabolites, to meet their functional needs. Primary metabolites are involved in vital biochemical pathways, while secondary or specialised metabolites mediate the interactions of living organisms with their biotic and abiotic environment. The locution "natural product" (NP) generally refers to secondary or specialised metabolites and primary metabolites, excluding macromolecules such as proteins, polysaccharides, or nucleic acids. Phytochemistry aims to chemically characterise NPs from various living organisms and to link this chemical information with biological activity profiles to address research questions mainly in chemical ecology or new active substances discovery for different economic sectors such as pharmaceutical, cosmetic, and agrochemical industries.

Three major events had a considerable impact on NP research. The first, in 1992, is linked to the Rio Convention on Biological Diversity, which laid the foundations for the Nagoya Protocol on access to genetic resources and the fair and equitable sharing of benefits arising from their use. This new regulation has been perceived as a constraint by many pharmaceutical companies and has contributed to a slowdown in research in the field. Nevertheless, an inventory of NPs introduced as drugs between 1981 and 2019 has been carried out by Newman and Cragg. Of the 1,881 molecules brought to market during these 39 years, 71 are NPs stricto sensu, 356 are derivatives of NPs, often obtained by semi-synthesis, and 272 are obtained by organic synthesis while having a pharmacophore inspired by NPs. This demonstrates, if proof was needed, that NPs are still an essential source of inspiration for Western medicine. They are also playing an increasingly important role in the cosmetics industry.

The second event concerns the advent of the bioeconomy and biobased chemistry concepts, which aims to add value to biomasses, mainly agricultural, algal or forestry, without competing with food resources. Therefore, while biodiversity hotspots have historically attracted the most significant research interest, the chemical profiling of dedicated biomasses or co-products from existing industries has become an essential challenge for achieving carbon neutrality by 2050 via the circular economy concept. This aspect of NP research is reinforced by recent results demonstrating unambiguously that the planetary boundary related to the introduction of new chemical entities (i.e., derived from organic synthesis) has widely been exceeded.

The last event concerns the increasing number of phytochemists who recently have actively engaged in addressing the principles of open science and promoting the production of FAIR data. Significant initiatives have been launched in this context, enabling our research activities to benefit from essential open-access tools in spectral data processing and databases.

Although the potential for NPs to reach high added value sectors and the interest in studying them for fundamental research questions are significant, the phytochemistry work often remains complex, time-consuming, tedious, and costly. Together with the entry into force of the Nagoya Protocol, this is one of the reasons for which industrial research programs on natural products have slowed down or even stopped since the beginning of the 2000s. Nevertheless, the massive arrival over the last 15 years of new dereplication approaches enabling known compounds to be chemically characterised as far upstream as possible in the process (1) has given new impetus to phytochemical research, both at academic and industrial levels.(2) Nevertheless, identifying unambiguous structures by describing their unequivocal molecular topology and their geometry, among mixtures, remains challenging. Hence, "annotation" is commonly used when structural information is incomplete.

The two main spectroscopic methods used to study complex mixtures of NPs are nuclear magnetic resonance (NMR) and mass spectrometry (MS), often hyphenated with HPLC for the latter.(3) Their sensitivity and universality performance affects the number and the relevance of annotated compounds.

As stated by Sumner *et al.*, exploiting the advantages of at least two unrelated techniques (*e.g.*, NMR and MS) should improve the annotation reliability.(4) Various strategies (online *versus* offline combinations) were reported by Letertre *et al.*(3) and Marshall *et al.*(5) and are shown in Fig. S1.

When examined under the same conditions, a compound originating from two distinct biological sources is expected to display identical spectral features. Consequently, rapidly identifying well-known compounds, achieved by matching experimental data with established reference spectra, is generally straightforward, assuming the necessary spectral data is accessible. However, this process becomes considerably more intricate when dealing with complex mixtures.

Presently, while structures within natural product structural databases are relatively easy to retrieve (*e.g.*, LOTUS, COCO-NUT-DB, NP-MRD, DNP, NPAtlas, ChEBI, KNApSAcK, *etc.*), the main challenge lies in locating databases where each structure is associated with its corresponding spectral data (*e.g.*, GNPS, MarinLit). Access to comprehensive and easily searchable experimental spectral data remains crucial to ensure a high confidence level in annotations. (6) Nevertheless, natural product databases encounter limitations, such as incompleteness, inconsistency, availability loss, and/or restricted query flexibility. Additionally, data from NMR and MS are scattered throughout scientific publications in formats that do not facilitate automated data extraction for constructing

databases. As a result, even when such data exist in the literature, they are not uniformly incorporated into databases.

Due to these limitations, querying a database for known natural products is hindered by barriers that significantly slow down the overall data mining process. As a solution, predicting spectral data has emerged as a way to improve current spectral databases. In this context, existing spectral databases serve a broader purpose than expediting dereplication (i.e., identifying known compounds). They also serve as the cornerstone for developing algorithms capable of predicting spectral data (e.g., MetFrag, CFM-ID, QCxMS). These tools can then enrich current experimental spectral databases by incorporating simulated in silico spectral data (e.g., UNPD-ISDB, FooDB, AntiBase, Mona, etc.), subsequently streamlining the dereplication process (e.g., Network Annotation propagation) or aiding to develop robust systems for automatic de novo elucidation (e.g., SIRIUS, Sherlock (7)). However, the DB size and content impact annotation performance, as a more extensive database increases the probability of returning irrelevant candidates, whereas a database that is too restricted a priori may lead to the omission of annotations.(8) DB content should be *a priori* restricted by taxonomy, either on biological and/or chemical grounds, to reduce the number of irrelevant matching spectra.(9)

The build of such DB is nowadays an obstacle course. Large chemical libraries must be first surveyed for compounds, according to criteria defined by the purposes of the study. The molecular structures of the returned compounds must then be gathered into a structural DB. This implies all the issues concerning the stereochemistry description and the drawing of the molecular structures. Finally, these structural DBs should be able to be used as inputs for spectral prediction software. Not all software is user-friendly, and the general workflow, including all the required unit steps, is not automated and available under a single software package.

This work aims to present a comprehensive workflow encompassing the creation of a relevant structural/spectral database and the combined exploitation of MS² and ¹³C NMR data for the chemical profiling of a bio-based extract, with a specific focus on an ethyl acetate solid-liquid extract from *Larix decidua* bark.

Results

The dereplication workflow described hereafter and applied to the chemical profiling of *Larix decidua* (Pinaceae) bark ethyl acetate crude extract is based on the combination of LC-HRMS² and ¹³C NMR data to annotate as many NPs as possible with the highest confidence level. First, barks were ground to powder and then macerated with EtOAc to obtain a crude extract with a mass yield of 6.2 %. A 3.81 g portion of this extract was submitted to centrifugal partition chromatography (CPC) fractionation, leading to 12 chemically simplified fractions.

These fractions were submitted to LC-HRMS² and to ¹³C NMR analysis. The resulting data were subjected to processing and visualisation procedures. The HRMS² data were presented as molecular networks, using Ion Identity Molecular Network (IMN), Feature-Based Molecular Network (FBMN), and Propagated Annotation Network (NAP) workflows.(10–12) ¹³C

NMR data were analysed by hierarchical cluster analysis after chemical shift alignment, and the resulting chemical shift clusters were visualised as a heatmap.(13) The HRMS² and NMR data were then annotated using spectral comparison with structures contained in a custom DB created by selecting structures from the LOTUS DB according to taxonomic criteria and with the help of *in silico*, experimental and *de novo* elucidation tools.(14) The results from both workflows were compiled, then ranked and finally a confidence score was assigned to each annotation using a script developed for this purpose.

Structural and *in silico* spectral database of compounds from Pinaceae

Creating a database containing NPs associated with their spectral properties requires collecting the corresponding chemical structures. At first, in the framework of this study, a DB containing the NPs isolated from plants belonging to the Pinaceae family was created. The corresponding chemical structures were downloaded from the LOTUS database using the VersaDB GUI. VersaDB is a recently developed Pythonbased graphical user interface that integrates the opensource database and webserver LOTUS, CFM-ID, and nmrshiftdb2 in a unique dereplication workflow.(14) It has been designed to create local custom databases of NPs selected according to biological and chemical taxonomic criteria from the LOTUS DB. The resulting structural DB and the corresponding MS and ¹³C NMR predicted spectral DBs can be further used to perform dereplication of complex mixtures through dedicated platforms and software such as SIRIUS, NAP (MetFrag algorithm will use the structural database to predict and compare spectra to experimental data), GNPS, MetGem and ACD/Labs DB.(14)

Structure inquiry was initiated targeting the Pinaceae plant family. As a result, 2,790 non-redundant structures were obtained. The 5 most represented chemical superclasses were distributed as follow: diterpenoids (469, 16.8%), triterpenoids (316, 11.3%), sesquiterpenoids (306, 11.0%), flavonoids (287, 10.3%), and lignans (173, 6.2%). Structures which could not be classified by NPClassifier were labelled as "nan". (Fig. S2B)

The structural DB was then used to build the MS^2 and the ^{13}C NMR spectral DBs using VersaDB program. The default prediction settings were applied for both prediction tools within the VersaDB application (CFM-ID4.2.6 and NMRShiftDB), and MS^2 spectra were predicted for three collision energies in $[M+H]^+$ mode. The prediction was operated in 1 h 15 min and 26 sec with a PRECISION 3650 computer, equipped with an Intel(R) Core(TM) i9-10900K CPU @ 3.70GHz 3.70 GHz processor and with 64.0 Go RAM.

The predicted ¹³C NMR chemical shift values were then imported to the database management software ACD/NMR Workbook to check and correct the predicted chemical shift of each structure. Among inspected structures, 258 contained suspicious chemical shifts and thus were corrected using the ACD/NMR Workbook Suite. This database was then used for spectral matching with each chemical shift clusters resulting from the HCA of experimental spectra.

Concerning MS² spectra, CFM-ID failed to predict spectra from 3 compounds since these were molecules for which the SMILES retrieved from LOTUS contained more than one structure. Consequently, MS² predicted spectral DB gathered 8,361 MS² spectra (3 collision energies x 2,787 compounds). To benefit from the NAP workflow as a complement to the forthcoming spectral matching with experimental databases, the structural database has been enriched, using the http://dorresteinappshub.ucsd.edu/, with the following information: "MonoisotopicMass InChI SMILES Identifier InChIKey2 InChIKey1 MolecularFormula kingdom_name superclass_name class_name subclass_name". (15)

Compound identification from $^{\rm 13}{\rm C}$ NMR data by the Cara-Mel workflow

The CaraMel chemical profiling workflow (13) is based on the unsupervised clustering of experimental ¹³C NMR chemical shift values using the peak intensity profiles across centrifugal partition chromatography (CPC) fractions as discriminating features, as explained in the Method section (¹³C NMR dereplication: CaraMel workflow). The resulting heatmap highlighted 25 chemical shift clusters (Fig. 1) from the 403 chemical shift values picked in the spectra of 12 CPC fractions of the EtOAc *Larix decidua* bark extract.

The chemical shift values in each cluster were compared to those from the custom Pinaceae 13 C NMR predicted database (2,790 structures associated with their predicted 13 C NMR chemical shifts).

As a result, 12 compounds were directly annotated, as shown in Fig. 1, by submitting the ¹³C chemical shifts of the corresponding cluster from HCA to the $^{\rm 13}{\rm C}$ NMR Pinaceae database, using the structure search engine included in the ACD/NMR Workbook database management tool. If relevant, the annotations of these compounds contained in the EtOAc bark extract and their relative configurations were confirmed by the complementary analysis of ¹H and 2D NMR data (COSY, HSQC and HMBC spectra). These compounds, marked by a blue star symbol in Fig. 1, are catechin (cluster 1), epicatechin (cluster 2), quercetin-3-rhamnoside (cluster 3), trans-astringin (cluster 6), lavandoside (cluster 9), larixyl acetate (cluster 11), ferulic acid (cluster 13), larixinol isomers (cluster 14), 13epimanool (cluster 20), isopimaric acid (cluster 21), dehydroabietic acid (cluster 22) and 7-oxo-dehydroabietic acid (cluster 23).

Thirteen other structures were annotated (Fig.1) based on the initial proposals returned by ACD/NMR workbook (although erroneous, they often provide very useful hints for "manual" annotation) and from 2D NMR data of chemically simplified fractions: acetic acid (cluster 4), piceatannol-3'-oglucoside(trans) (cluster 5), glucosyl frambinone (cluster 7), glycerol monoacetate (cluster 8), glucosyl-trans-paracoumaric acid (cluster 10), 2-[2,4dihydroxybenzoyl)oxyphenyl]acetic acid (cluster 12), dianthoside (cluster 15), oleic acid (cluster 18), linoleic acid (cluster 19), butanol (cluster 1'), tannins (catechic units) (cluster 2'). Three of these NPs were not directly annotated even though they were present in the VersaDB Pinaceae database generated for this study because many of their chemical shifts have been clustered with those of other compounds

(e.g., trans-astringin (cluster 6) and trans-piceatannol-3'-oglucoside (cluster 5), or lavandoside (cluster 9) and glucosyl trans-para-coumaric acid (cluster 10). Moreover, the structures reported in Table S8 of two previously non-reported compounds: rhamnosyl-(1->6)-glucosyl-myrtenic acid and arabinosyl-(1->6)-glucosyl-myrtenic acid (respectively cluster 17 and 16), were elucidated based on 1D and 2D NMR data of CPC fraction 3. HMBC correlations were observed between protons H-1' and carbon C1, between proton H-1" and carbon C-6', and between protons H-6' and carbon C-1', thus confirming the structure of arabinosyl- $(1\rightarrow 6)$ -glucosyl-myrtenic acid. In addition, the correlation between carbons C-4" and C-5" and proton H-6" confirmed the rhamnosyl- $(1\rightarrow 6)$ -glucosylmyrtenic acid structure. These two compounds were also detected while analysing CPC fraction 3 by mass spectrometry, and the corresponding features have been annotated with their molecular formula (respectively C₂₁H₃₂O₁₁: feature 248, 483.18 m/z, [M+Na]⁺, feature 294, 499.16 m/z,[M+K]⁺; and C22H34O11: feature 306, 497.20 m/z, [M+Na]⁺). Their structures are close to that of sacranoside A, but with myrtenic acid as aglycon moiety instead of myrtenol. Interestingly, the monoterpene myrtenol and oxidised derivatives have also been previously identified from the bark of Picea abies (Pinaceae).(16)

As a result, 16 of the 25 compounds identified by the NMR workflow were previously reported in the Pinaceae and are marked with the PNC symbol in Fig. 1. The chemical diversity of the 25 identified compounds was studied according to NPClassifier output, including five diterpenoids (larixyl acetate, 13-epimanool, isopimaric acid, dehydroabietic acid, 7oxo-dehydroabietic acid), five flavonoids (catechin, epicatechin, suercetin-3-rhamnoside, larixinol isomers, tannins (catechic units)), three phenylpropanoids (lavandoside, glucosyltrans-para-coumaric acid, ferulic acid), three fatty acids (oleic acid, linoleic acid, butanol), two stilbenoids (trans-astringin, piceatannol-3-o-glucoside), two monoterpenoids arrabinosylglucosyl-myrtenic acid, rhamnosyl-glucoside-myrtenic acid), one aromatic polyketide (2-[2,4dihydroxybenzoyl)oxyphenyl]acetic acid), one cyclic polyketide (dianthoside), two undefined compounds (glucosylframbinone, glycerol monoacetate), and the erroneous annotation from NPClassifier for acetic acid as small peptide. Fig. S2B shows the global chemical distribution for these compounds.

Table 1 and Fig. 1 show the distribution of the identified compounds within the 12 fractions. The ¹³C and ¹H chemical shifts of the annotated compounds are reported in Table S8.

$\operatorname{LC-HRMS^2}$ data annotations through molecular network workflows

Dereplication applied to CPC fractions was performed using LC-HRMS², and experimental spectral data were subjected to annotation using specific algorithms, including IIMN (17), FBMN (18), NAP (10), and SIRIUS (19). The experimental data were also compared (18) to the simulated MS data of the custom Pinaceae structural and spectral DB using the open-source software MetGem (17). However, these results are not reported due to the low annotation rate, attributed to the poor quality of the predicted spectra as shown in Fig. S4.

The raw data processing resulted in the detection of 1,958 sets of signals corresponding to a chemical entity, also called features. MS² spectra similarity-based molecular network was generated using the Feature Based Molecular Network (FBMN) workflow (see GNPS platform (18)). The chromatographic peak shape correlation analysis was integrated into the molecular network by connecting and collapsing the different ion species of the same molecule via the lon Identity Molecular Networking (IIMN), which enhances annotations within the fragmentation pattern similarity molecular network. Annotations of MS² spectra through spectral matching with reference spectra (FBMN) were improved through the use of an *in silico* approach known as Network Annotation Propagation (NAP), as well as employing a *de novo* strategy provided by the SIRIUS software

IIMN + FBMN

The FBMN workflow returned 83 matches with the GNPS speclibs collection, 54 unique spectra (46 unique compounds): seven candidates were validated by the IIMN, encompassing six distinct spectra associated with unique compounds: epicatechin, quercetin, and curcumin, as well as norethindrone, gestoden and a piperazine-derivative antihistamine: cyclizine. The annotation of these three latter compounds within the natural substance mixture is indeed surprising, considering that they are of synthetic origin. However, norethindrone and gestoden, two synthetic steroids, could be considered as natural product-like compounds when calculating their natural product likeness scores (NaPLeS = 1.8 and 2, respectively), in comparison to cyclizine (NaPLeS = -0.6). The more positive the score, the higher the NPlikeness.(19) The most representative annotated chemical class were steroid (29%), diterpenoïd (23%), and flavonoid (17%), as shown in Fig. S2. B.

NAP

Fifty-three of the FBMN matches were also annotated with NAP. Thirty-seven of these annotations result from at least one structure candidate of the *in silico* fragmentation search with the MetFrag algorithm (20), including 25 unique spectra. NAP allowed the annotation of 156 not previously annotated features, using only *in silico* fragmentation search with Met-Frag. Hence, the *in silico* fragmentation search allowed the annotation of 39.8% of all the considered features by the NAP algorithm versus 4.2% of all data set using only experimental spectral database search. In total, NAP led to the annotation of 9.9% of all the detected features.

SIRIUS

The *de novo* strategy facilitated the annotation of 1,786 features with at least one molecular formula and corresponding adduct type. Among these, 896 features garnered multiple proposals, while 890 features were associated with a single proposal, resulting in a cumulative count of 2,847 proposals. Following the subsequent re-ranking step, 1,300 distinct molecular formulas were attributed to the 1,786 features.

Integration of the Pinaceae structural database into the annotation workflow enabled the annotation of 498 features, each associated with at least one potential structural candidate retrieved from the Pinaceae structural database, encompassing the assignment of 196 distinct structures as primary candidates. The five most frequently assigned structures the following: LTS0139452 ($C_{19}H_{28}O_2$, 22 times), LTS0056945 ($C_{20}H_{28}O_4$, 16 times), LTS0251392 ($C_{20}H_{28}O_3$, 15 times), LTS0084143 ($C_{20}H_{30}O_3$, 13 times) and LTS0241153 ($C_{20}H_{32}O_2$, 11 times). Ultimately, 1,728 features have been annotated up to the chemical class level using the webservice CANOPUS integrated in SIRIUS. These annotations are categorised across seven biosynthesis pathways: terpenoids (663 instances), fatty acids (307 instances), polyketides (234 instances), shikimates and phenylpropanoids (160 instances), alkaloids (116 instances), amino acids and peptides (106 instances), and finally carbohydrates (41 instances).

Combination of ¹³C NMR and LC-HRMS² data annotations

The experimental spectra were annotated through three strategies: experimental spectral databases matching, in silico spectral databases matching, and de novo annotation. The first one relies on the use of GNPS workflows and FBMN. The in silico annotation workflow relies on NAP and CaraMel. The manual analysis of NMR data and the application of the SIRI-US software, both employed in this study, are integral components of the third strategy. The latter allowed for the annotation of experimental spectra without depending on spectral databases, even in cases undisclosed chemical structures were involved. A new workflow was created to combine the annotations from all the GNPS workflows (i.e., FBMN, NAP), SIRIUS and the ¹³C NMR annotation workflow called CaraMel. This new workflow named « CATHEDRAL » for « Combining THE DiffeRent Annotation tooLs. » is an in-house script written in Python. This script allows the assignment of a confidence level score using a custom confidence level system, determined based on the outcomes of the comparative analysis.

First, FBMN and NAP results are merged to retrieve the spectral matches *versus* experimental spectral libraries from GNPS and Pinaceae *in a silico* database simulated by MetFrag.

Secondly, the resulting metadata is exported as a "recap.csv" file. A PRED column is added to the table, and features are "scored" as follows: 1 for the FBMN annotated features, 2 for the features without any FBMN annotation but with at least one NAP – MetFrag candidate, and 3 for all the remaining features.

The comparison process starts with the SMILES from the FBMN workflow and the MetFragSMILES from the NAP workflow. The corresponding InChIKeys are then generated *via* functions from the RDKit Python library.

The canonical SMILES from ¹³C NMR annotations structures, obtained from ChemDraw 20.1.1., and gathered into a file called «smile_nmr.txt», follow the same process as above.

The resulting InChiKeys for each feature and CaraMel annotations are thus compared to the corresponding SIRIUS annotations InChiKeys.

CATHEDRAL.py can be executed through a detailed stepby-step jupyter notebook (CATHEDRAL.ipynb) under an RDKit anaconda environment. The recap.csv file is first opened, and features corresponding to the PRED score = 1 are selected. FBMN annotations are compared to MetFrag candidates for each feature. In the second place, FBMN annotations are compared to the corresponding SIRIUS candidates. In the third place, the FBMN, NAP (MetFrag), and SIRIUS annotations are compared to each other to highlight those with the same candidate through the three annotation tools. The greater the number of tools giving the same annotation, the better the score and priority accorded to annotation tools based on their effectiveness in providing a relevant structural candidate (c.f. CASMI). Thus, a spectral match with a reference experimental spectrum (i.e., FBMN) will be prioritised over a predicted spectrum (i.e. MetFrag). However, de novo annotations (*i.e.*, SIRIUS) will be ranked higher than the latter. Furthermore, the scores are better when a candidate is corroborated by NMR, unlike cases where the annotation solely relies on MS/MS tools. Consequently, the features with consistent annotation along the three mass annotation tools are scored with a Confidence_Level = 9. The features with consistent annotations between FBMN and SIRIUS are scored 10, the features with consistent annotations between FBMN and MetFrag are scored 12, and features with consistent annotations between MetFrag and SIRIUS are scored 11+. The corresponding annotation for each scored feature is then compared to the CaraMel candidates. If the MS² candidate is part of CaraMel annotations, the header « Is_NMR_Annotated_SU » = 1, and the corresponding molecular name is given to the Molecular_Name_SU header « » else « Is NMR Annotated SU » = 0, where SU means Sum-Up. Consequently, features previously scored (9, 10, 11, 11+, 12), also annotated by CaraMel, are respectively re-scored 1,2, 3+ or 4. Then, only the data with a PRED score = 2 are selected. The features with consistent annotations between MetFrag and SIRIUS are scored 11. The annotation is then compared to the CaraMel annotations, as previously explained. If these features are also annotated with CaraMel, the score is raised to 3.

If an annotation is common to at least two MS² annotation tools, the third tool annotation is equally compared to the CaraMel annotations. If it matches, the corresponding molecular name is reported in the header « 3rd_Tool_SU » of the sum-up file.

All the other features that do not have consistent annotations between the different MS² tools are equally compared to the CaraMel annotations. If the annotation from FBMN matches, Confidence_Level corresponds to 5; if one annotation candidate matches for at least the NAP workflow, Confidence Level is equal to 7. Finally, if only a candidate annotated from SIRI-US matches with a CaraMel annotation, Confidence_Level will be 6. CaraMel annotations that do not correspond to any mass annotation are scored 8. All the corresponding confidence levels are reported in Table. 2.

The comparison outcomes are then written as a table in a sum-up file called « df_resume_confidence.tsv » starting with the following table column headers: « 3rd_Tool_SU, Confidence_Level, Feature_SU, GNPS_SU, Is_NMR_Annotated_SU, Molecular_Name, NAP_SU, Not_Matching_Tool_Annotation_SU, Rt_SU, SIRIUS_SU, m/z ». This sum-up file is then imported to the previously created

network collection in Cytoscape software, and metadata are overlayed on the graph *via* the in-house style, as exemplified in Fig. 2.

CATHEDRAL workflow highlights up to 41 unique compounds, annotated with at least two indiscriminately MS/MS or NMR dereplication tools, including 7 compounds in Fig. 3. The outcomes are reported in Tables. S1-S7. The best confidence level annotated structures correspond to those with the same candidates for all annotation tools. They are all classified as flavonoid derivatives and FBMN / SIRIUS jointly annotated features. In addition, 21 among the 27 structures jointly annotated by NAP / SIRIUS correspond to diterpenoids (abietane and derivatives) and triterpenoids. Most features annotated jointly by at least two different tools are [M+H]⁺ (37 among the 50 iterations of annotated compounds). The structure candidates highlighted by NAP / SIRIUS and SIRIUS / CaraMel comparisons also include other ion species and insource modifications, respectively [M+Na]⁺ and [M+K]⁺ adducts or in-source modifications.

As a result, the strategy for matching experimental spectral databases appears to prioritize annotating features associated with flavonoid chemical families and [M+H]⁺ adducts over other chemical families and ion species. Moreover, the chemical diversity is more prominent when considering the chemical classes related to annotated compounds from *in silico* and *de novo* strategies.

Finally, the *de novo* strategy for mass data annotation allowed the annotation of nine of the 25 structures found by ¹³C NMR data annotation. Five compounds among them were neither jointly listed by experimental nor predicted mass spectral database matching.

Conclusion & Discussion Summary

Untargeted metabolomics produces vast amounts of analytical data, yet the laborious analysis does not always result in a high annotation confidence level.

Due to the complementarity of ¹³C NMR and LC-HRMS², their combined use can potentially increase the number of detected compounds and significantly enhance the confidence level in their identification. Consequently, various strategies have been developed recently, such as online (*e.g.*, LC-MS-SPE-NMR) and offline approaches (*e.g.* multiblock).(3) However, the former requires bulky and expensive equipment (*i.e.*, ensuring compatibility of experimental conditions), while the latter enables in-depth data exploration but requires advanced expertise in chemometrics.

Hence, our strategy revolves around the cross-comparison of annotations from both ¹³C NMR and LC-HRMS² spectral datasets. Despite the high simplicity of our cross-comparison process, two major challenges persist. On one hand, automation should make conventional manual procedures more efficient. On the other hand, the expected gain in confidence is noticeable only for compounds detected by both techniques.

Consequently, compound annotation by a single technique remains ambiguous and requires more traditional strategies, such as exploiting multidimensional NMR data. Additionally, the quality of the annotation process, which heavily relies on spectral comparison, depends on the availability of relevant high-quality spectral or structural databases.

The initial step of the developed workflow involved the automatic compilation of tailored structural and predicted spectral databases, including the biological origin of the sample. This approach grants the crucial role of contextualization in NP research by narrowing down the pool of candidate structures whose affiliation with a specific plant species, genus, or family has been previously documented. It is also imperative to clarify that this step relies on queries from the LOTUS natural product database, built directly from Wikidata. The data are organized in triples, containing referenced structure-organism pairs that establish relationships between distinct molecular structures and the living organisms from which they were identified. Furthermore, the information comprised in LOTUS is associated to research findings provided by the scientific community, and as a result, it can suffer from the voluntary or involuntary omission of certain compounds in the chemical description of a given species, genus, or family. Nevertheless, to our knowledge, LOTUS represents the most extensive structural database available, including numerous metadata and is well-suited for automated online queries.

Consequently, integrating the Pinaceae family structural database with MS annotation tools resulted in a notable increase in annotated features. Interestingly, the Pinaceae contextualization of databases employed in NAP and SIRIUS processes enabled us to achieve 25% and 10% annotation rates, respectively, surpassing the 5% annotation rate achieved using the GNPS spectral library. However, the in silico annotation strategy is still limited by the accuracy of spectral prediction algorithms (e.g., the MS² predicted database generated by CFM-ID 4.0). According to LOTUS, this trend was also observed with NMR, where 16 of the 25 identified compounds were associated with the Pinaceae family. The remaining nine compounds were manually verified not to be genuinely related to any Pinaceae species: only dianthoside was an exception.(21) Furthermore, the last eight molecules correspond to substances that can be considered handling pollutants, such as butanol. Another concerns tannin, for which the complete molecular structure has not been elucidated. Others are not previously documented compounds, such as the myrtenic acid glycosides derivatives. The last molecules, piceatannol-3'-o-glucoside (i.e., a positional isomer of trans-astringin previously associated with Pinaceae), 2-[2,4-dihydroxy-6-(4-hydroxybenzoyl)oxyphenyl]acetic acid glycerol acetate and its hydrolysis product acetic acid, are likely the only compounds in the set that have never been described in Pinaceae, despite previous documentation. Given its meticulous compilation work, these results affirm our choice of considering LOTUS as the optimal starting point for contextualizing our databases.

The second step of this workflow involved efficiently scoring candidates through the CATHEDRAL script, using a custom confidence level system. Hence, a confidence level was assigned to 52 compounds, annotated with at least two annotation tools. This original scoring strategy markedly enhances the compound annotation reliability and reinforces the structure-organism pairs. This approach also reduces the requirement for tedious manual analysis, including successive purification and structural elucidation steps to confirm the presence of compounds in the studied matrix. As a result, it accelerates chemical profiling through dereplication.

Recent research has ultimately shown that expanding analytical dimensions for multidimensional metabolite annotation (*e.g.* Rt, m/z, chiroptical data, *etc.*) offers a significant boost, enhancing the accuracy and coverage of metabolites. More specially, ion mobility (IM) MS has emerged as a powerful technology that allows for the metabolite measurement of collision cross-section values (CSS). Consequently, the CCS aids in isomer characterization by providing an additional separation dimension to mass, increasing the identification confidence. (22–25) Similar to the four-dimensional untargeted metabolomic concept introduced by Cai *et al.*(26), including multidimensional data and/or metadata (*i.e.*, chemical ontology, scoring strategies, *etc.*) would significantly benefit our assessment methodology and, consequently, the consistency of identifications.

In turn, data from new identified compounds will participate to the completion of existing databases, thereby improving our understanding of the diversity of specialized metabolites biosynthetic pathways. This knowledge will contribute to optimizing the production of high added value compounds in the pharmaceutical industry, as seen in the case of artemisinin production in *Artemisia annua* through the up-regulation of amorphadiene synthase (multiplying yield by 2.3).(27)

Finally, the past and present habits in natural product chemistry research and publishing have led to issues, such as poorly defined chemical composition of plants. Insufficient description and standardization in the natural product practices, coupled with the irresponsible and inefficient management of experimental reference data, can render further research unreliable due to inherent inconsistencies in the substance reporting. This has contributed to the relative neglect of pharmacological studies on natural substances.(28) Additionally, it is essential to enhance the user-friendliness of tools to encourage phytochemists to persevere with better practices.

Despite the need for basic programming skills, this userfriendly approach contributes to global standardization in natural product research, following the ChemBioPrint[®] framework. Its consistency in providing high confidence levels for identified components in complex mixtures of natural products eases the selection of authentic molecules for pharmacological studies, such as reverse docking, thereby reducing compound to drug development time.

References

 Beutler JA, Alvarado AB, Schaufelberger DE, Andrews P, McCloud TG. Dereplication of phorbol bioactives: Lyngbya majuscula and croton cuneatus. J Nat Prod [Internet]. 1990 Jul [cited 2019 Nov 25];53(4):867–74. Available from: https://pubs.acs.org/doi/abs/10.1021/np50070a014

- Hubert J, Nuzillard JM, Renault JH. Dereplication strategies in natural product research: How many tools and methodologies behind the same concept? Phytochemistry Reviews. 2017;16(1):55–95.
- Letertre MPM, Dervilly G, Giraudeau P. Combined Nuclear Magnetic Resonance Spectroscopy and Mass Spectrometry Approaches for Metabolomics. Vol. 93, Analytical Chemistry. American Chemical Society; 2021. p. 500–18.
- Sumner LW, Amberg A, Barrett D, Beale MH, Beger R, Daykin CA, et al. Proposed minimum reporting standards for chemical analysis: Chemical Analysis Working Group (CAWG) Metabolomics Standards Initiative (MSI). Metabolomics. 2007;3(3):211–21.
- Marshall DD, Powers R. Beyond the paradigm: Combining mass spectrometry and nuclear magnetic resonance for metabolomics. Vol. 100, Progress in Nuclear Magnetic Resonance Spectroscopy. Elsevier B.V.; 2017. p. 1–16.
- Schymanski EL, Jeon J, Gulde R, Fenner K, Ruff M, Singer HP, et al. Identifying Small Molecules via High Resolution Mass Spectrometry: Communicating Confidence. Environ Sci Technol. 2014 Feb 18;48(4):2097–8.
- Wenk M, Nuzillard JM, Steinbeck C. Sherlock—A Free and Open-Source System for the Computer-Assisted Structure Elucidation of Organic Compounds from NMR Data. Molecules. 2023 Feb 2;28(3):1448.
- Bergold AN, Heaton P. Does filler database size influence identification accuracy? Law Hum Behav. 2018 Jun 1;42(3):227–43.
- Nuzillard JM. Taxonomy-Focused Natural Product Databases for Carbon-13 NMR-Based Dereplication. Analytica. 2021 Jun 28;2(3):50–6.
- Schmid R, Petras D, Nothias LF, Wang M, Aron AT, Jagels A, et al. Ion identity molecular networking for mass spectrometry-based metabolomics in the GNPS environment. Nat Commun. 2021 Dec 1;12(1).
- Nothias LF, Petras D, Schmid R, Dührkop K, Rainer J, Sarvepalli A, et al. Feature-based Molecular Networking in the GNPS Analysis Environment. bioRxiv. 2019 Oct;812404.
- da Silva RR, Wang M, lix Nothias LF, J van der Hooft JJ, Mauricio Caraballo-Rodríguez A, Fox E, et al. Propagating annotations of molecular networks using in silico fragmentation. 2018;
- Hubert J, Nuzillard JM, Purson S, Hamzaoui M, Borie N, Reynaud R, et al. Identification of natural metabolites in mixture: A pattern recognition strategy based on 13C NMR. Anal Chem [Internet]. 2014 Mar 18 [cited 2021 Jun 17];86(6):2955–62. Available from: https://pubmed.ncbi.nlm.nih.gov/24555703/
- Cordonnier J, Remy S, Renault JH, Nuzillard JM. Versa DB: Assisting 13 C NMR and MS/MS Joint Data Annotation Through On-Demand Databases. 2023; Available from: https://doi.org/10.1002/cmtd.202300020
- 15. Network Annotation Propagation (NAP) GNPS Documentation [Internet]. [cited 2023 Jan 13]. Available from: https://ccmsucsd.github.io/GNPSDocumentation/nap/#structure-database
- Heemann V, Francke W. Gaschromatographisch– Massenspektrometrische Untersuchungen der Flüchtigen Rindeninhaltsstoffe von Picea abies (l.) Karst. Planta Med. 1977 Dec 13;32(08):342–6.
- Olivon F, Elie N, Grelier G, Roussi F, Litaudon M, Touboul D. MetGem Software for the Generation of Molecular Networks Based on the t-SNE Algorithm. Anal Chem. 2018 Dec 4;90(23):13900–8.
- Wang M, Carver JJ, Phelan V V., Sanchez LM;, Garg N;, Peng Y;, et al. Sharing and community curation of mass spectrometry data with GNPS. Nat Biotechnol. 2017;34(8):828–37.
- Vanii Jayaseelan K, Moreno P, Truszkowski A, Ertl P, Steinbeck C. Natural product-likeness score revisited: an open-source, opendata implementation. BMC Bioinformatics. 2012 Dec 20;13(1):106.
- Wolf S, Schmidt S, Müller-Hannemann M, Neumann S. In silico fragmentation for computer assisted identification of metabolite mass spectra [Internet]. 2010. Available from: http://www.biomedcentral.com/1471-2105/11/148
- PLOUVIER V. Recherche du dianthoside chez les Pinacées et quelques autres groupes botaniques, et de l'érigéroside chez les Composées-Astérées. Comptes Rendus de l'Académie des Sciences, Série III. 1984;298:749–52.
- 22. Paglia G, Smith AJ, Astarita G. Ion mobility mass spectrometry in the omics era: Challenges and opportunities for metabolomics and

lipidomics. Mass Spectrom Rev [Internet]. 2022 Sep;41(5):722–65. Available from: https://analyticalsciencejournals.onlinelibrary.wiley.com/doi/10.10

02/mas.21686

- Luo MD, Zhou ZW, Zhu ZJ. The Application of Ion Mobility-Mass Spectrometry in Untargeted Metabolomics: from Separation to Identification. J Anal Test [Internet]. 2020 Jul 25;4(3):163–74. Available from: https://link.springer.com/10.1007/s41664-020-00133-0
- Zheng X, Smith RD, Baker ES. Recent advances in lipid separations and structural elucidation using mass spectrometry combined with ion mobility spectrometry, ion-molecule reactions and fragmentation approaches. Curr Opin Chem Biol [Internet]. 2018 Feb;42:111– 8. Available from: https://linkinghub.elsevier.com/retrieve/pii/S1367593117301540
- Picache JA, Rose BS, Balinski A, Leaptrot KL, Sherrod SD, May JC, et al. Collision cross section compendium to annotate and predict multi-omic compound identities. Chem Sci [Internet]. 2019;10(4):983–93. Available from: http://xlink.rsc.org/?DOI=C8SC04396E
- Cai Y, Zhou Z, Zhu ZJ. Advanced analytical and informatic strategies for metabolite annotation in untargeted metabolomics. Vol. 158, TrAC - Trends in Analytical Chemistry. Elsevier B.V.; 2023.
- 27. Wen W, Yu R. Artemisinin biosynthesis and its regulatory enzymes: Progress and perspective. Pharmacogn Rev [Internet]. [cited 2023 Nov 11];(10). Available from: www.phcogrev.com
- Lai CT, Shan JJ, Rodgers K, Sutherland SK. Challenges in Natural Health Product Research: The Importance of Standardization [Internet]. Vol. 50, Proc. West. Pharmacol. Soc. 2007. Available from: https://www.researchgate.net/publication/5246233
- Darme P, Spalenka J, Hubert J, Escotte-Binet S, Debelle L, Villena I, et al. Investigation of Antiparasitic Activity of 10 European Tree Bark Extracts on Toxoplasma gondii and Bioguided Identification of Triterpenes in Alnus glutinosa Barks [Internet]. 2022. Available from: https://journals.asm.org/journal/aac
- Pluskal T, Castillo S, Villar-Briones A, Orešič M. MZmine 2: Modular framework for processing, visualizing, and analyzing mass spectrometry-based molecular profile data [Internet]. 2010. Available from: http://www.biomedcentral.com/1471-2105/11/395
- Dührkop K, Fleischauer M, Ludwig M, Aksenov AA, Melnik A V., Meusel M, et al. SIRIUS 4: a rapid tool for turning tandem mass spectra into metabolite structure information. Nat Methods [Internet]. 2019 Apr 1 [cited 2021 Jun 17];16(4):299–302. Available from: https://www.nature.com/articles/s41592-019-0344-8
- 32. Stein SE, Scott DR. Optimization and Testing of Mass Spectral Library Search Algorithms for Compound Identification.

Supporting information

The relevant source code of CATHEDRAL script for the assignment of a confidence level score determined based on the outcomes of the comparative analysis comparison of annotations, can be found on GitHub at https://github.com/Jcrdnr/CATHEDRAL.

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Competing interest statement

The authors have no conflicts of interest to declare.

Keywords

Dereplication – Mass spectrometry – Nuclear Magnetic Resonance – Natural products

Materials and Methods

The dereplication workflows are summarised in Fig. 4. Plant material

The Nagoya Protocol was adopted by France in 2010 and entered into force in 2014. The French "Office National des Forêts" (ONF) granted the authors permission to collect samples and to use bark extracts at the University of

Reims Champagne-Ardenne. *Larix decidua* bark was collected on a standing tree by M. Verdeaux in Signy-L'Abbaye (Ardennes, France), route forestière Roban, 70th plot on the 11th of May 2017. The sample was dried for three days at 30°C before being finely powdered by a hammer mill (VEM Motors GmbH, Germany). The botanical identification of trees was made following the phenotypic characteristics, such as the shape, arrangement, and contours of leaves.(29)

Sample preparation

Collected Larix decidua barks were cleaned to remove unwanted materials such as moss and soil particles, and the crude extract was prepared by maceration of 70 g bark powder in 1 l ethyl acetate (EtOAc) at room temperature for 24 under magnetic stirring. After filtration under vacuum on a 40 μ m sintered glass, the solvent was evaporated under vacuum to leave a dry EtOAc extract. The extraction of Larix decidua bark by EtOAc yielded 4.34 g (6.2 %) of crude extract.

Sample fractionation

The crude extract was fractionated by Centrifugal Partition Chromatography (CPC). It is based on liquid–liquid compound partitioning between two nonmiscible liquid phases that stay close to their thermodynamic equilibrium. The absence of solid chromatographic support facilitates the fractionation of complex samples, eliminates the risk of stationary phase overloading or clogging, and avoids the potential deterioration of chemical compounds, thus leading to mass recovery rates close to 100 %.

CPC apparatus

CPC experiments were performed on an FCPC200 [®] apparatus (Rousselet Robatel Kromaton) equipped with a rotor containing 800 cells (260 ml total column capacity) and connected to Smartline Preparative Pump 1800 (Knauer). The eluent was collected with a LABOCOL Vario-4000 fraction collector (Knauer) in fractions of 20 ml.

Solvents system

The CPC fractionation of the extract was performed in two consecutive steps (CPC1 and CP2) to obtain the broadest possible diversity of fraction polarity. **Gradient elution system n-heptane/ethyl acetate/methanol/water:**

The initial mobile phase (lower phase of system 1: n-heptane/ethyl acetate/methanol/water (5:5:5:5, v/v)), the final mobile phase (lower phase of system 3: n-heptane/ethyl acetate/methanol/water (7:3:7:3, v/v)) and the organic stationary phase (upper phase of system 2: n-heptane/ethyl acetate/methanol/water (6:4:6:4, v/v)) were prepared separately.

Isocratic elution system ethyl acetate/acetonitrile/water:

The system composed of ethyl acetate/acetonitrile/water (3:3:4) (system 3) was used in the normal-phase mode (upper phase mobile).

Injection and CPC operating procedure

<u>CPC 1</u>

The column was filled with the stationary phase of the gradient elution system and equilibrated with the initial mobile phase at 10 ml/min and 1600 rpm. The crude extract (3.81 g) was dissolved in 3 ml of lower phase + 3 ml of upper phase and injected into the CPC column through a 20 ml loop. As described in Table, the mobile phase was pumped in descending mode for 110 minutes. 3. The column was extruded by switching the mode selection valve for 15 minutes at 20 ml/min. Fractions of 20 ml were collected over the whole experiment (elution and extrusion) and combined according to their thin layer chromatography profiles (TLC).

<u>CPC 2</u>

The column was filled with the stationary phase (upper phase) of the isocratic system and equilibrated with the lower phase at 10 ml/min and 1800 rpm. Then the CPC1 first fraction (252 mg) was dissolved in 3 ml of lower phase + 3 ml of upper phase and injected into the CPC column by a 20 ml loop. The mobile phase was pumped in ascending mode for 90 minutes. The column was extruded by switching the mode selection valve for 20 minutes at 20 ml/min. Fractions of 20 ml were collected over the elution step and combined according to their TLC profiles.

TLC of CPC fractions

The 20 ml fractions of the two CPC experiments (elution and extrusion) were characterised by TLC. TLC was performed on pre-coated silica gel 60 F254 Merck plates, with the migration solvent system consisting of ethyl acetate/toluene/acetic acid/formic acid (4/6/1/1, v/v). Compound migration was visualised under UV light at 254 nm and 366 nm and revealed by spraying consecutively the dried plates with 50% (v/v) H₂SO₄ acid and vanillin (10 g / l) ethanolic solution followed by heating. As a result, six fractions were obtained from CPC1. The first fraction (consisting in the seven first elution minutes (t0 of the experiment)) was used for CPC2. The other fractions were

noted F08, F09, F10, F11 and F12 (Figure S2). The fractions from CPC2 were arranged in reverse order of collection (because CPC1 was performed in descending mode while CPC2 was performed in the ascending mode) to obtain a chemical profile continuity between the two fractionation steps. The fractionation of the bark extract resulted in twelve chemically simplified fractions of polarity ranging from the highest (F01) to the lowest (F12) (Fig. S2), ten produced by elution and two by subsequent column extrusion.

The fractions were then analysed in parallel by LC-HRMS² and ¹³C NMR. Experimental spectral data were then processed and annotated while taking advantage of purposely created databases. A comparison of the resulting candidate structures from both workflows was carried out to enhance confidence in the final profiling outcomes.

Generation of custom structural and spectral databases: VersaDB

The VersaDB GUI was employed for constructing customised structural and spectral databases. The methodology previously outlined was subsequently applied. (14) Briefly, the process involved an automated structural inquiry utilising the 'for all selected categories' function of VersaDB. This function allowed the transmission of HTTP requests *via* the LOTUS API, applying a singular chosen criterion: 'T: All_Taxonomy_DB: family: Pinaceae'. This approach entailed a comprehensive search for structures documented within botanical species affiliated with the Pinaceae family. This was executed by leveraging the entirety of available taxonomic databases originating from the LOTUS Natural Product Occurrence Database.

The resultant compilation of structures, each accompanied by its respective LOTUS ID, was archived in two analogous files: cfmidinput.txt and structuraldb.txt. These files served as input for the MS² and ¹³C NMR spectra prediction or were employed as the structural database within the SIRIUS and NAP platforms. However, it requires formatting adjustments before it can be utilised in the NAP workflow. To achieve this, the file was subjected to processing using the http://dorresteinappshub.ucsd.edu/ web server. The original data, initially presented in the 'SMILES - CompoundID' format, undergo enhancement and transformation into a novel arrangement: "MonoisotopicMass InChI SMILES Identifier InChIKey2 InChIKey1 MolecularFormula kingdom_name superclass_name class_name subclass_name.". Additional metadata from LOTUS, encompassing chemical and physical attributes, taxonomic classification, and chemontology information according to the NPClassifier ontology, were consolidated and stored within a file named "cfmid input.tsv.".

The "predict both properties" functionality within VersaDB was subsequently employed to predict both $\rm MS^2$ and $\rm ^{13}C$ NMR spectral properties.

Concerning ¹³C chemical shifts prediction, the VersaDB system incorporated an adaptation of Kuhn and Nuzillard's approach, leveraging the nmrshiftdb2 packages. The outcome was the "13CNMRDatabase.sdf" file containing the structures, their ¹³C nmrshiftdb2-predicted chemical shifts in the style of ACD/Labs CNMR Predictor, and all compound-related metadata gathered from LOTUS, encompassing biological, chemical taxonomy, and physicochemical properties. This "13CNMRDatabase.sdf" file was then imported into ACD/NMR Workbook Suite 2012 (ACD/Labs, Ontario, Canada), where the predicted ¹³C chemical shifts underwent validation using the "check chemical shifts" option.

The prediction of MS² spectra was conducted using the CFM-ID 4.2.6.0 docker image. MS² spectra were predicted across three collision energies, employing the pre-trained CFM-ID models. This process culminated in the creation of the definitive custom *in silico* mass spectral database file, denoted as "MSMSspectraDatabase.mgf", along with the "annotationGNPSformat.tsv" file containing the compound's metadata. These outputs were fashioned to meet the requirements for publishing the database on the GNPS platform. ¹³C NMR dereplication: CaraMel workflow

Data acquisition

All samples were analysed using the same acquisition and processing parameters. Fractions were dried under vacuum, and aliquots (up to about 20 mg when possible) of CPC fractions were dissolved in 600 μ l DMSO-*d*₆ and analysed by nuclear magnetic resonance (¹H, ¹³C, HSQC, HMBC, and COSY) at 298 K on a Bruker Avance AVIII-600 spectrometer (Karlsruhe, Germany) equipped with a TCI cryoprobe. ¹³C NMR spectra were acquired at 150.91 MHz using a standard udeft pulse sequence with an FID acquisition time of 0.36 s, a relaxation delay of 3.00 s, and the accumulation of 1,024 scans.

Data processing

The absolute intensities of all ^{13}C NMR signals detected in all spectra were collected by automatic peak picking, after spectra processing (manually phased, baseline corrected, and referenced by setting the central resonance of DMSO-d_6 at δ 39.80 ppm) using the TOPSPIN 4.1.3 software (Bruker, Rheinstetten, Germany). Each peak list was converted into a text file contain-

ing peak positions and absolute peak intensities. Peaks positions from the whole set of spectra were then aligned by an algorithm written in the Python language. The principle was to divide the ¹³C spectral width (from 0 to 240 ppm) into regular bins of 0.2 ppm width and to place the absolute intensity of each ¹³C peak into the corresponding bin. The bins without any signal, regardless of fraction, were removed from the bin list. The resulting global table contains 12 columns, each corresponding a CPC fraction, and 403 rows, corresponding to the NMR spectral buckets for which at least one ¹³C NMR peak was detected in at least one spectrum. This table was imported into the PermutMatrix software (version 1.9.3, LIRMM, Montpellier, France) for hierarchical clustering analysis (HCA). This operation reordered the rows of the global peak table so that similar rows, corresponding to similar chromatographic emergence profiles, were grouped together and lead to define clusters of chemical shifts values, with ideally one cluster defined per compound contained in the EtOAc extract of Larix decidua bark. The similarity between table rows was measured by the Euclidian distance, and data agglomeration was performed with the Ward's method. The resulting clusters of ¹³C NMR chemical shifts were visualised as dendrograms on a heat map.

Each ¹³C NMR chemical shift cluster obtained from HCA was manually submitted to the VersaDB-generated database containing the structures and corresponding predicted NMR chemical shifts values (nmrshiftdb2) of 2790 natural metabolites found in the Pinaceae family (April 2022), via the structure search engine from ACD/NMR Workbook Suite 2012 (ACD/Labs, Ontario, Canada). This dereplication procedure was described in a previous article.(13) A ¹³C NMR chemical shift difference between the predicted and experimental spectra was tolerated between 2 and 3 ppm, and the minimum number of ¹³C query shift values to match was set at about 80% of the number of chemical shifts in the cluster. Finally, each structure proposal provided by the database query and its associated relative configurations was confirmed by interpretation of 1D and 2D NMR data (¹H, ¹³C, HSQC, HMBC, COSY). LC-HRMS² dereplication

Data acquisition

LC analyses were performed with Waters QSM Acquity, equipped with an UPTISPHERE Strategy C18 column ($2.2\mu m \times 150 mm \times 2.1 \mu m$, Interchim). The eluent consisted of H₂O + 0.1% formic acid (A) and MeCN (B), following a gradient 5-30 % B in 4 min, then 30-80 % B in 14 min, then 80-100 % in 0.50 min, then maintaining 100 % B for 9.5 min at a flow rate of 0.5 ml/min. The wavelength range of the UV detector was set from 210 to 400 nm.

Mass data were acquired with a Waters SYNAPT G2-Si (QTof) mass spectrometer. The electrospray ionisation source was set as follow: positive mode, source temperature 100°C, capillary 3 kV, desolvation temperature 450°C, nebuliser gas flow 5 Bar, desolvation gas flow 700 l/h. MS scans were performed in full-scan mode from m/z 100 to 1200 (scan time 0.1 sec) with a resolution of 40 000 (FWHM). An MS¹ scan was followed by MS² scans of the three most intense ions above a threshold of 3000 counts (exclusion window 3 sec). The selected parent ions were fragmented according to the following energy ramp: low mass start: 35 eV, low mass end: 55 eV, high mass start: 70 eV high mass end: 130 eV. Leucine-enkephalin (1 ng/µL) was used as a reference mass *via* a lock spray interface at a flow rate of 10 µl/min for positive ion mode monitoring ([M + H]⁺ = 556.277).

MZMine 3 data processing

The 12 MS^2 raw files were converted from the .raw (Waters) standard data format to .mzml format using the MSConvert software, part of the ProteoWizard package (version 3.0.21349, Palo Alto, CA). All .mzml were processed by MZMine 3 v0.21 beta(30) in batch mode.

The mass detection was realised by keeping the noise level at 50. The ADAP chromatogram builder was used with a minimum group size of scans of 5, a group intensity threshold of 5000, a minimum highest intensity of 7500, and m/z tolerance of 0.005 (or 20 ppm). The ADAP feature resolver was used for the deconvolution step and the wavelets deconvolution algorithm was applied with the following standard settings: S/N threshold = 5, minimum feature height = 7500, coefficient/area threshold = 20, peak duration range 0.01-1 min, RT wavelet range 0.01-0.12 min. MS² scans were paired using an m/z tolerance range of 0.05 Da and RT tolerance range of 0.5 min. Isotopologues were grouped using ¹³C isotope filter (isotopic peak grouper) algorithm with an m/z tolerance of 0.005 (or 20 ppm) and a retention time (RT) tolerance of 0.2 min and a maximum charge of 3. The peak alignment was performed using the join aligner module [m/z tolerance = 0.004 (or 10 ppm), weight for m/z = 2, weight for RT = 1, absolute RT tolerance 0.5 min]. The peak list was gap-filled with the same RT and m/z range gap filler module [m/z tolerance of 0.004 (or 10 ppm)]. The feature list was filtered using the feature filter algorithm as follow: Duration 0-3 min, data points 3-10000. Features were then filtered using the row filter algorithm as minimum features in an isotope pattern set to 2.

Finally, the metaCorrelate algorithm was applied to the dataset with the following parameters, RT tolerance 0.1 min, feature height correlation and intensity correlation threshold were set to 0. Min sample in all 1, Min % intensity overlap 40%, Exclude estimated features (gap-filled) turn on. Concerning correlation grouping parameter, min datapoints 5, min data points on edge 2, measure Pearson, min feature shape correlation 85%. Concerning the feature height correlation parameters, min datapoints 3, measure Pearson, min correlation 60%. For the next step, the Ion Identity Networking algorithm was used as follow: m/z tolerance 0.001 (10 ppm), check all features, min-height 0. MS mode positive, maximum charge 2, maximum molecules/cluster 2, adducts: [M+H]+, [M+Na]⁺, [M+NH4]⁺, [M+2H]2⁺, [M-H+2Na]⁺, [M+H+Na]2⁺, modifications: [M-H₂O], [M-2H₂O]. Less common adducts were then added: [M+K]+, [M+Ca]2⁺, [M+Fe]2⁺, [M+H+NH₄]2⁺, [M+H+K]2⁺, [M+Ca-H]⁺, [M+Fe-H]⁺. Finally, adducts that tend to form clusters/in-source fragments were added: [M+H]⁺, [M+NH₄]⁺, [M+2H]2⁺, and modifications: [M+HFA], [M+ACN], [M-H₂O], [M-2H₂O], [M-3H₂O], [M-4H₂O]. Ion identity network refinement was applied using the following parameters: minimum size 3, delete smaller networks: link threshold 5. All ion identities were checked with m/z tolerance (MS^2) 0.002 (15 ppm), with checking for multimers and neutral losses ($MS^1 \rightarrow MS^2$).

Eventually, the .mgf preclustered spectral data file and its corresponding .csv metadata files (for RT, areas, and formulas integration and edges MS¹ annotation) were exported using the dedicated "Export for GNPS" built-in options. Processed spectral data were exported to SIRIUS with m/z tolerance of 0.002 (10 ppm) without merging MS/MS.

SIRIUS annotation

SIRIUS 4 is considered a state-of-the-art metabolite annotation solution, which combines molecular formula calculation and the prediction of a molecular fingerprint of a query compound from its fragmentation tree and spectrum.(31) Although the SIRIUS algorithm is integrated into GNPS, it was executed locally with Sirius 4.8.2 GUI, as it allows to use an in-house database for the CSI FingerID contrarily to GNPS. The parameters used to proceed to the spectral data analysis were the following for SIRIUS molecular formula calculation: possible ionisation [M+H]⁺, [M+K]⁺, [M+Na]⁺, instrument: Q-TOF, ppm tolerance 10 ppm, Top molecular formula candidates: 5, filter: formulas from all available DBs, Elements allowed in Molecular Formula: H, C, O, N (max: 3). Zodiac algorithm was used for re-ranking predicted formula as follow: [candidates 300m/z 10, candidates 800 m/z 50, use 2-step approach. Edge Threshold 0.95, min local connections 10, iterations 20 000, burn-in 2000, separates runs 10]. For the CSI: FingerID step, the parameters were the following: possible adducts: [M+H]⁺, [M+K]⁺, [M+Na]⁺, [M-H₂O+H]⁺, filter: compounds present in the in-house PINACEAE DB, the maximal number of returned candidate structures: unlimited. Eventually, the CANOPUS algorithm was applied to predict the chemical class of compounds based on NPClassifier chemical ontology.

Ion Identity Molecular Network

The molecular networks are based on matching spectral patterns between paired mass spectra, accomplished by calculating cosine similarity. This metric measures the cosine of the angle between the vectors representing non-zero spectral data points in a multidimensional hyperspace, where the dimensions correspond to the considered mass-to-charge ratio (m/z) variables. The intensity of each data point corresponds to the coordinate value along the respective mass axis within this hyperspace. (26,32)

The molecular networks were created using the online workflow at Global Natural Products Social molecular networking (http://gnps.ucsd.edu). An Ion Identity Molecular Network (IIMN) was created (iob: 25b1448341ab454c9002c1767fba98e1) where edges were filtered to have a cosine score above 0.4 and at least 4 matching peaks. Further edges between two nodes were kept in the network if and only if each of the nodes appeared in each other's respective top 10 most similar nodes. All matches kept between the network spectra and the library spectra were required to have a score above 0.7 and at least 4 matched peaks. The default speclibs from the GNPS platform were used for the spectral library search. A Feature Based Molecular Network was realised (iob: Of64e360d2ba4227beb6a2e0a03a5335) to decrease the library search cosine-score threshold from 0.7 to 0.4, with at least 4 matching peaks. The default speclibs from the GNPS platform were used for the spectral library search. The FBMN results were used for Network Annotation Propagation (NAP) step (iob: 35984a76a5794491acd88cb694e30843). The parameters used to proceed with analysis were the following: N first candidates for consensus score 10, Accuracy for exact mass candidate search (ppm) 20, acquisition mode Positive, Multiple adduct types $[M+H]^{\ast},\ [M+Na]^{\ast},\ [M+2O+H]^{\ast},\ user-provided database: NAP formatted in house Pinaceae structural database. The maximum number of candidate structures in the graph was set to 3.$

Merging and comparing SIRIUS annotations with IIMN, NAP

IIMN, FBMN, NAP, and MolNetEnhancer results were downloaded from the GNPS platform as archive folders. Metadata from the different GNPS annotation workflows were merged via Cytoscape 3.8.2 software (https://cytoscape.org/), following the method explained in Fig. S3. The IIMN graph was kept visualising the molecular network. Quantification table and the "df_resume_confidence.tsv" file resulting from the CATHEDRAL comparison data were overlayed on the molecular network using custom style.



Figure 1: Annotated heatmap, built with ¹³C NMR chemical shifts of the twelve fractions from EtOAc *L. decidua* extract. (M) Compound equally annotated through MS2 data, orange compounds: previously undescribed compounds, (PNC) Compound structure described in the ¹³C NRM predicted Pinaceae database, (Blue star) Compound directly annotated by Caramel workflow



Figure 2: IIMN visualization with CATHEDRAL comparison outcomes overlayed



Figure 3: Comparison of MS/MS clusters A to C to the annotated CaraMel heatmap.



Figure 4: Dereplication workflows based on ¹³C NMR (CaraMel) and LC-HRMS² of *Larix decidua* EtOAc crude extract by Centrifugal Partition Chromatography

CPC fractions	Mass (mg)	% Crude extract	Composition
01 – elution	42	10.8 %	dianthoside (Maj – cluster 15)
02 – elution	12	3.1 %	glucosyl-frambinone (Min – cluster 7); glycerol- monoacetate (Maj – cluster 8); lavandoside (Min – cluster 9); glucosyl <i>trans-para</i> -coumaric acid (Min – cluster 10); rhamnosyl-(1->6)-glucosyl- Myrtenic acid (Min – cluster 17)
03 – elution	18	4.7 %	glucosyl-frambinone (Med – cluster 7); arabino- syl-(1->6)-glucosyl-myrtenic acid (Maj – cluster 16); rhamnosyl-(1->6)-glucosyl-myrtenic acid (Maj – cluster 17)
04 – elution	85	22.0 %	acetic acid (Min – cluster 4); astringin (trans) (Maj – cluster 6); piceatannol-3'-O-glucoside (Maj – cluster 5); tannins (catechin unit) (Med – cluster 2')
05 – elution	29	7.5 %	catechin (Maj – cluster 1); epicatechin (Min – cluster 2); quercetin-3-rhamnoside (Med – clus- ter 3); acetic acid (Min – cluster 4); oleic acid (Min – cluster 18); linoleic acid (Min – cluster 19)
06 – elution	26	6.7 %	epicatechin (Min – cluster 2); acetic acid (Min – cluster 4); 2-[2,4-dihydroxy-6-(4- hydroxybenzoyl)oxyphenyl]acetic acid (Maj - cluster 12); ferulic acid (Med – cluster 13); lar- ixinol isomer 1 (Min – cluster 14); quercetin-3- rhamnoside (Min – cluster 3); oleic acid (Min – cluster 18); linoleic acid (Min – cluster 19)
07 – elution	38	9.8 %	larixinol isomer 1 (Maj - cluster 14); larixinol isomer 2 (Med – cluster 14); acetic acid (Min – cluster 4); oleic acid (Min – cluster 18); linoleic acid (Min – cluster 19)
08 – elution	16	4.1 %	catechin (Med – cluster 1); astringin (<i>trans</i>) (Min – cluster 6); piceatannol-3'-O-glucoside (Min – cluster 5); glycerol-monoacetate (Min – cluster 8); ferulic acid (Med – cluster 13); oleic acid (Min – cluster 18); linoleic acid (Min – cluster 19)
09 – elution	8	2.1 %	oleic acid (Med – cluster 18); linoleic acid (Min – cluster 19); 7-oxodehydroabietic acid (Min – cluster 23)
10 – elution	17	4.4 %	glycerol-monoacetate (Min – cluster 8); larixyl acetate (Maj – cluster 11); oleic acid (Min – clus- ter 18); linoleic acid (Min – cluster 19); 7- oxodehydroabietic acid (Min – cluster 23)
11 – extrusion	83	21.4 %	larixyl acetate (Min – cluster 11); oleic acid (Med – cluster 18); linoleic acid (Med – cluster 19); dehydroabietic acid (Min – cluster 22)
12 – extrusion	13	3.4 %	glycerol-monoacetate (Min – cluster 8); oleic acid (Med – cluster 18); linoleic acid (Min – clus- ter 19); 13-epimanool (Maj – cluster 20); isopi- maric acid (Maj – cluster 21); dehydroabietic acid (Med – cluster 22)

Table 1: Mass and global composition of the CPC fractions. (Maj=major; Med=medium; Min=minor). Compounds are related to their corresponding cluster on Fig.1.

Feature jointly annotated with	<u>Confidence</u> <u>level</u>		
FBMN, NAP, SIRIUS, CaraMel	1		
FBMN, SIRIUS, CaraMel	2		
NAP, SIRIUS, CaraMel	2+		
(+ other FBMN annotation)	37		
NAP, SIRIUS, CaraMel	2		
(no other FBMN annotation)	5		
FBMN, NAP, CaraMel	4		
FBMN, CaraMel	5		
SIRIUS, CaraMel	6		
NAP (one structure candidate of the <i>in silico</i> fragmentation search with MetFrag), CaraMel	7		
CaraMel	8		
FBMN, NAP, SIRIUS	9		
FBMN, SIRIUS	10		
NAP, SIRIUS	11+		
(+ other FBMN annotation)	11+		
NAP, SIRIUS	11		
(no other FBMN annotation)			
FBMN, NAP	12		

Table 2: Custom confidence levels established according to the tools for which the feature was jointly annotated with.

Time (min)	Flow rate (ml / min)	% Lower phase system 1	% Lower phase sytem2
0	1	100 %	0 %
3	10	100 %	0 %
15	10	100 %	0 %
90	10	0 %	100 %
110	10	0 %	100 %
125	Extrusion (20 ml / min)	Extrusion	Extrusion

Table 3: Mobile phase composition during CPC 1 fractionation.

ChemRxiv[®]

Improving the chemical profiling of complex natural extracts by joint ¹³C NMR and LC-HRMS² analysis and the querying of *in silico* generated chemical

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Table S3: Summary of compounds jointly annotated by NAP, SIRIUS and eventually CaraMel, with another FBMN candidate (*c.f. Excel file Table_S3.xlsx*).

Table S4: Summary of compounds jointly annotated by NAP, SIRIUS and eventually CaraMel, without any FBMN candidate (c.f. Excel file Table_S4.xlsx).

Table S5: Summary of compounds jointly annotated by SIRIUS and CaraMel (c.f. Excel file Table_S5.xlsx).

Table S6: Summary of compounds jointly annotated by NAP and CaraMel (*c.f. Excel file Table_S6.xlsx*).

Table S7: Summary of compounds jointly annotated by FBMN and SIRIUS (c.f. Excel file Table_S7.xlsx).

Figure S1: The reported strategies for the joint use of MS and NMR analytical data, with their benefits and constraints

Figure S2: Chemical distributions

Figure S3: Merging the annotations produced by the MS workflows, leading to the recap.csv file

Figure S4: Comparison of experimental spectra of dehydroabietic acid from GNPS library to predicted spectra of dehydroabietic acid from LOTUS structure (LTS0252977) from CFM-ID 4.0 algorithm.

ted	52	~	S3	S4	~	7		SS			S6				-			S2		22	ß	ÿ	7
Rela	Table.	IN	Table.	Table.	Nat	Na		Table.			Table.				Table			Table.		Table.	Table.	Table	Nat
<u>Graphical</u> shape	Yellow diamond	NaN	Yellow rectangle	Yellow rectangle	NaN	NaN		V shape			Triangle				Yellow higlighted			Diamond		Hexagon	Rectangle	Rectangle	NaN
Cumulative nb compounds	3	3	4	9	6	6		12			12				23			25		28	31	52	52
Compound(s) name	catechin / epicatechin querectani-Ahammoside catechir chai	AN Concentration of the concen	dehydroabietic acid	dehydroabietic acid Ianrihoil komens - sociadrihotelacia acid	AN A	NaN	catechin / epicatechin limoleic acid dehydroableitic acid	isomerator Isophanic acid	7-oxo-denyarioabiette acid astringion	13-septimation	astringin 7-oxodehydroabietic add dehydroabietic add	acetic acid piceatamol-3:-o-glucoside	glucosyl frambinone oliverol monoacetate	jayanda ku	glucosyl trans para coumaric acid 2-12.4 digydroxy-6-(4-tiydroxybenzoyl)oxyphenyl]acet ic acid	dianthoside arabinosy/1-5-65-glucosy/ myrtenic acid	rhamnosyl-(1->6)-glucosyl myrtenic acid oleic acid	quercetin kaempferol	odanielic odd	nour querceth-3-galactoside morreceth-3-galactoside	1,44-dimethyl/7-lprop-1-en-2-ylh-1,2,3,4,8,9,10,10 a-octahydrophenanthrene-1-carboxylic acid 19-nor-4-hydroxyabieta-8,11,13-trien-7-one 18-nor-4,15-dihydroxyabieta-8,11,13-trien-7-one	picetational picetational picetatione a productione A provider Statistic and method and a still 3 prodocamatrien 18 oic add method a still 3 treem 18 oic add methyd febydroubietate methyd febydroubietate (9 tes 23) 2 3 (4 treem 18 oic add methyd 2 3 (2 shydroxyposan 2 yr) / 4 (11,13 stream methyd 2 3 2 2 2 0 2 8.06.8 0.12.17) foctadecame 15,2 'oxoland) 6 yllpropanoate methyd 19,2 3 (4 stream methyd 2 3 2 15,16,17 decahydro-1H-cyclopental/a) phenanthren 17 - ylpentanoic add (9 tes 23,24 at) (13,14 spentamethyd 2 3,56,7,81 2 15,16,17 decahydro-1H-cyclopental/a) phenanthren 17 - ylpentanoic add (9 strathory 44,10,13,14 spentamethyd 2 3,56,7,81 2 15,16,17 decahydro-1H-cyclopental/a) phenanthren 17 - ylpentanoic add (9 strathory 44,10,13,14 spentamethyd 2 3,56,7,81 2 15,16,17 decahydro-1H-cyclopental/a) phenanthren 17 - ylpentanoic add (9 strathory 44,10,13,14 spentamethyd 2 3,11,11,13 strathar methyd 3 5,16,181,141,179,2 strathydrone 4,4,10,13,1 (1 2 - 6,16) strathar 11,2	NaN
Nb unique compound	е	NaN	-	e	NaN	Nan		6			3				E			2		s	5 features with a unique MetFrag candidate 23 unique compounds	41 features with andrugue MeFFag candidate & 2 1 unique compounds	NaN
Nb Features	4	NaN	-	9	NaN	NaN		42			4				NaN			4		12	15	103	NaN
CararMel	×	×	×	×	×	×		×			×			;	×								
SIRIUS	×	×	×	×				×										×		×	×	×	
AN	×		×	×	×						×							×			×	×	×
FBMN	×	×			×	×												×		×			×
<u>Confidence</u> Level	-	2	3+	e	4	5		é			7				80			6		10	+11+	=	12

Table S1: Summary of compounds annotated during chemical profiling of *Larix decidua bark* crude extract, after the comparison of the candidate coming from each annotation tool (*c.f. Excel file Table_S1.xlsx*).

स						Carr
CaraM	2	YIS	g	YES	XIS .	Classifier
NPClassifier Superclass	Flavonoids	Flavoneids	Flavoncids	Flavoncids	Flavonoids	M
SiRiUS annotation	ALTMATREDAJIN CENADOZIN + NI+	игдеккоминен сонакордин + н-	Interenticopia + H+	Prizvesco-Poten Costeolie + H-	orgucurrowward careboal ((M + H)+	notation and Stiffild
MSMS spectrum						MS MS spectrum
NAP - MelFrag candidate (LOTUS ID)	(150004651	LT5006122	179155822	1150265245	862386238	NAP - MetFrag
FBMN matched mirror spectra	itter ühesi and stij förstedate tradi jähnskolli 1406/03/13/13/60.060/05/15/16/comfer aft 1506/060/13/13/16/06/16/15/15/16/comfer aft	tten üzen soal eki finensiste irozi in hendoli Holikonal 20 ekisionalosi Sistemenen al	ther/Jimes und eleftretedd efred acherdade 19560aut20e60ac6030336erreter af 19560aut20e00ac60ac603	tttor üpen sod ett freedde freed a fraidold Holdonal David sod an 2018 freeder af L	the all planes and end interface (a decade and a decade	FBMM matched mirror spectra
GMPS spectrum ID	CONFUNDODOS739139	CONSUMPORES N2589	CONSUM0005 NeSs	Taltesonophismoo	Contraction	GMPS spectrum ID
MS1 spectrum						MS1 spectrum
Mol. Weight	302.24	678.52	106.24	290.27	846.38	Mol. Weight
Name	guercelin	catechic tannin	ksempferol	catechia	querceto 3-thermocide	Name
Structure	Ho to to to		to Control of			Structure
Feature ID	120	15 2011	198 182	8	287	Q
						Featu

CaraMel	YES	2	2	2	
NPClassifier Superclass	Diterpenoid	Diterpenoid	Diterpenoid	Diterpenoid	
SIRIUS annotation	NFWOWWBERKOV CONSOZIA + HI-	NGBRPGLXCOJPU CZ046802(M + M)-	PTOFVCHKVSVCM C1942602[M + H]+	sochzekopfinist, crish2so2(M + Na)+	
MS MS spectrum					
candidate (LOTUS ID)	LT50034017	LT50068733	LTS0108097	LT50171116	
FBMN matched mirror spectra	Otee	Otee	Oter	Otex	
GNPS spectrum ID	Oher	Oher	Oher	Oher	
MS1 spectrum		4 4 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
Mol. Weight	300.44	298.42	286.41	302.41	
Name	dehydroubiele-acid	1,4a-dreethy,7,1prop.1-en- 2,4%,10,10a- 1,2,3,4,4,8,10,10a- cethoryfic acid cethoryfic acid	19-nor-4-hydroxysbiete- 8,11,13-then-7-one	18-nor-4,15- dihydroryabela- 8,11,13-blen-7-one	
Structure		- Here Level		The second secon	
Feature ID	1211	497 750 854	822	668	
-					

Table S2: Summary of compounds jointly annotated by all annotation tools (c.f. Excel file Table_S2: xlsx). Table S3: Summary of compounds jointly annotated by NAP, SIRIUS and eventually CaraMel, with another FBMN candidate (c.f. Excel file Table_S3.xlsx). file Table S3.xlsx).

_									
CaraMel	YES	YES	YES	9	2	9	Ŷ	9	2
NPCIassifier Superclass	Diterpenoid	Fisconoid	Diterpanoid	Sitheroid	Triterpanoid	Diterpencid	Diterpenoid	Lignan	Phicroglucinol
SiRiUS annotation	иятикомонаясного соннароди + на-	Provisou Zycski canazoran - H-	MSW.JSD.NPCSSNW C20H2403[M + H]+	connuccencre crantoopa . My.	NFRISIOHEADFRO CJOHHROJM - M-	-М- Меоганер: ганысинанопп	DWHTYJMRWXJOJI, C17H22010 + Naj-	OTWLSOPCSOEBAY CIGHTBOTH = Nal-	ouwrrawnow.ec.comecogiw + H+
MS/MS spectrum									
NAP - MetFrag sandidate (LOTUS ID)	LTS0034017	LTS0269975	LTS0229171	LTS0094372	LTS0224467	LTS0079469	LTS0083733	LTS0044016	LTS0194549
EBMN matched mirror spectra	Ober	Others	Other	Other	Other	Other	Other	Ober	Oher
GNPS spectrum ID	Ober	Ober	Otes	Otex	Ober	Ober	Other	Otex	Obec
MS1 spectrum			المراجع المراجع المراجع المراجع ا	The second					
Mol. Weight	900 44	542 49	314.42	244.24	424.70	310.30	214.36	370.35	326.43
Name	dehydroebielic acid	britinol isomers	7-ciso-detrychoabietic acid	picestarred	cyclosecolations	picostactions A	13-hydroy-8,1113-podicarpation-	meridinol	ocalionatione
Structure	- Care	5 2 2 2 2 2 2 2 2 2 2 2 2 2		8-0-5 2-0-5	25, Far	-	E Contraction of the second se		
Feature ID	674 849 1272	276 336	1185	158	1267	888	2	230	629

 Table S4 (Part 1.):
 Summary of compounds jointly annotated by NAP, SIRIUS and eventually CaraMel, without any FBMN candidate (c.f.

 Excel file Table_S4.xlsx).





	-				
Ŷ	Ŷ	Q.	Ŷ	2	Ŷ
Diterpencid	Triterpanoid	Triarpenoid	Lignan	Diterpencid	Diterpencid
ouucreakeu.FS controjn - Hi-	KUMASYJANPIHK CITHEOUIM - H-	SoleEXWISIOUEZ CISH44C0(N + M-	FITARICNICYDO CI94240[4 + M-	socid28/convest, cri+recordin + Hj-	SOMLIGNOOH CI948IQM + Hj-
(TTTTTTTTTTT Hereine		12 12 10 10 10 10 10 10 10 10 10 10 10 10 10			(1)(1)(1)(1)(1)(1)(1)(1)(1)(1)(1)(1)(1)(
LTS0210076	LTS0041481	LTS0087433	LTS0155650	LTS017116	LTS0118921
Ces	Celer	Office	Celes	Office	Otes
Ohee	Oher	Oher	Oner	Oher	Ohee
			the state of the s	El Contraction of the second s	
270.45	406.73	424.06	368.39	302.41	272.43
dehydratiotherun	piceanold B	(26.5R.96.10R.13R.14R.17R)-2- ethological and the sentementy, 17.12R)- 4.4.10.131.4.andtamethy, 17.12R)- 4.6.5.0.11.251.5.6.17.costby490-1H cyclopenta(a)phenanthren-3-cos	lonage4.64(modg);7,1-(i)	18-nor-4, 15-dihydi oyyabista- 8,11, 13-krien-7-ona	(15.445) 1045)-1.44 dmmthyl-7. prostr2-34 2.34.5.10,05 headhydrophenanthree 1.41
-695	The second secon	HALL HALL	A A A A A A A A A A A A A A A A A A A		
090 028 904 1219 1425	1347	1366	665 961 1316	423 565 516	676

Table S4 (Part 3.): Summary of compounds jointly annotated by NAP, SIRIUS and eventually CaraMel, without any FBMN candidate (*c.f. Excel file Table_S4.xisx*).

CaraMel	YES	YES	YES	VES	YES	YES	YES	YES	YES
MCClassifier. Superclass	Favoreid	Fatty Acid	Diterpenoid	Phenyteropaneid (O6-C3)	Diterpenoid	Diterpenoid	Satemoid	Diterpenoid	Diterprecid
StillS. antotation	егаменорогими станаоцім. ноо + н +	OMIQUUCINUIS CHRISOZIM + Naj-	NPWOWWNERACY CONTROLM + H-	KSEBMYGRYZTDHS CIGHTOOUJM - H2O + Hj-	-In- com- micognocy karantariana	MSW.JSDLIPCSSW C20H2603[M + H]+	PERFIRILUZIOUN CZOHOZONIM + M-	CERTIFICAN CONHADJM - HOO + HJ+	исисионансциен сонносији - ноо - нј-
MG MG apoctican				t. totale the second					
MAP - MelFrag sandidate (LOTUS, ID)	NaN	NaN	NPN	New	NWN	NaN	NaN	NaN	NaN
EBMN matched mirror spectra	NaN	NaN	NaN	NaN	NaN	Nev	NaN	haN	New
GNPS associum D	New	NaN	NaN	New	NaN	Nev	New	NaN	New
MS1 aperium		t talentalah merinden en en en				1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
Mol. Weight	290.27	280.45	300.44	194.18	302.45	314.42	8C 90#	290.48	348 52
Marrie	catechin / epicatechin	Brokić add	dehydrodbiefic acid	fourfic acid	hophmark actd	7-oso-dehydroeleelis acid	ubute suc	13 eșinurool	laring acceler
Stucture	Ho Contraction of the second s	and a state of the		at the second se			HO H, O HO H	A A A A A A A A A A A A A A A A A A A	2 (2 (2)) 00 (2 (2 (2)) 00 (2 (2)) (2 (2)) (2 (2)) (2 (2)) (2 (2
Eesture ID	76 250	209 958 959 960	436 743 858 855 935 1134 11233 2014	57 196 700 1036 1194 1194 11328 1617 1910	063 876 8701 8701	396 400 400 461 461 461 461 583 678 678 1279 11279	159	1047	1182

Table S5: Summary of compounds jointly annotated by SIRIUS and CaraMel (c.f. Excel file Table_S5.xlsx).

CaraMel	YES	YES	YES	
NPCIassifier Superclass	Diterpenoid	Stibmoid	Diterpenoid	
SiffUS annotation	Nev	NaN	hav	
MS-MS spectrum				
NAP - MetFrag. sandidate (LOTUS ID)	LTS0251851	LT50251851	LT50229171	
FBMN matched mirror spectra	NAM	New	NRM	
GNPS spectrum ID	Mak	hav	hav	
MS1 spectrum	10 	Veranse and a second se		
Mol. Weight	300.44	90 OP	314.42	
Name	denyal cabieto acid	unganisas ansir	7-oue dehydroxbielic acid	
Structure				
Feature ID	942	124	420 816	

Table S6: Summary of compounds jointly annotated by NAP and CaraMel (c.f. Excel file Table_S6.xlsx).

CaraMel	Q	Q	Q	Q	Q.
NPClassifier Superclass	Diterpenoid	Fitworod	Filtyonoid	Fisvonoid	Flavonoid
SIRIUS annotation	ZOHUKYYELWEOK CZOH26O3M + H]+	IKGXIBGEBALURG C2740001694 + 4]+	Internet Control Control (14)	ovsorbikcevzwak cziłczorzyw + H+	REFANTPEDVLIN CISHIOO72M + HJ-
MS/MS spectrum					
NAP - MetFrag candidate (LOTUS ID)	NaN	NaN	NaN	NsN	Nez
FBIMN matched mirror spectra	Millor Targos usola edu Thrano CAF Anneu A Jap Taris - Of Hospicotom 422 robotinos concessos Henne 1964 al annotations . DB	htten: target usså eku Proteol Afrekreut jag frats 4014-60000-1422 holden annon sonstatsforkreur 1864 all annotations DB	Marchinges and edu Protocol Farment ing The Attraction of the Article State of the Attraction of the Attraction of Attraction of Attract	Mino. Tansa usat eki Printo A.F. enun la Drata Mino. Tansa usat ana Printo A.F. enun la Drata Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino. Mino.	Magacitypes useds each Pranto A Fahrends Jac Theorem 4000-4000 Concentration Concentration 2000-4000 Concentration Concentration 2000-4000 Concentration 2000-
GNPS spectrum ID	CCMSLIB00006570661 CCMSLIB00006570644	COMSUB000022062	CCMSUB000574966	CCMSLIB0005738276	OCMSUB000574873
MS1 spectrum	0 0 0 0 0 0 0 0 0 0 0 0 0 0				
Mot. Weight	316.44	610.52	286.24	464.38	302.24
Name	danieliic acid	ugn	kaampferol	quercelin-3-galactoside	ujecuarb
Structure			HO OH OH OH	OH HO HO HO HO HO HO HO HO HO HO HO HO H	HO HO HO
Feature ID	334 422 426 436 489 489 489 559 577 1366	162	530	280	345











Figure S2: Chemical distributions (A) HPTLC profile of the 12 CPC fractions of the *Larix decidua* EtOAc extract. The plate was sprayed with vanillin/ H_2SO_4 derivatization reagent, heated, and observed under white light (B) Sunburst charts representing the chemical distribution of the Pinaceae structural database, CaraMel annotations, FBMN annotations



6. Create PRED column in recap.csv and score features:

- 1: FBMN annotated features
- 2: NAP (\geq 1 MetFrag candidate) annotated features
- 3: all the remaining features



Figure S3: Merging the annotations produced by the MS workflows, leading to the recap.csv file



Figure S4: Comparison c 0 100 200 300 400 500 600 700 hydroabietic acid from LOTUS structure (LTS0252977) from CFM-ID 4.0 algorithm. (A) GNPS spectra vs. predicted spectra (10 eV). (B) GNPS spectra vs. predicted spectra (20 eV). (C) GNPS spectra vs. predicted spectra (40 eV). Red spectrum: CCMSLIB0000840371 Blue spectra: CFM-ID 4.0 predicted spectrum.

Table S8: Annotated structures by CaraMel workflow, linked to their ¹³C NMR and ¹H chemical shifts.



$C_{15}H_{14}O_6$	290.3 g/mol	CAS 154-23-4
Atom number	¹³ C (ppm)	¹ H (ppm)
2	81.5	4.48
3	66.6	3.81
4	28.3	2.66/2.35
4a	99.6	-
5	156.6	-
6	94.1	5.68
7	156.8	-
8	95.6	5.89
8a	155.7	-
1'	131.0	-
2'	114.9	6.72
3'	145.2	-
4'	145.3	-
5′	115.3	6.68
6′	118.8	6.59

epicatechin



$C_{15}H_{14}O_6$	290.3 g/mol	CAS 490-46-0
Atom number	¹³ C (ppm)	¹H (ppm)
2	78.3	4.73
3	65.3	4.00
4	28.7	2.67/2.47
4a	98.9	-
5	156.0	-
6	94.4	5.72
7	156.5	-
8	95.6	5.89
8a	156.1	-
1'	130.9	-
2'	118.3	6.65
3'	144.7	-
3'	144.7	-
4'	144.8	-
5'	115.2	6.66
6'	115.4	6.88



$C_{21}H_{20}O_{11}$	448.4 g/mol	CAS 522-12-3
Atom number	¹³ C (ppm)	¹ H (ppm)
2	156.8	-
3	134.5	-
4	178.1	-
4a	104.4	-
5	161.6	-
6	99.0	6.21
7	164.6	-
8	94.0	6.40
8a	157.7	-
1'	121.1	-
2'	121.5	7.25
3'	115.8	6.87
4'	148.8	-
5'	145.5	-
6'	116.0	7.29
1"	102.1	5.25
2"	70.4	3.98
3"	70.7	3.51
4"	71.5	3.15
5"	71.0	3.20
6"	17.9	0.81

	acetic acid O 2 OH	
$C_2H_4O_2$	60.5 g/mol	CAS 64-19-7
Atom number	¹³ C (ppm)	¹ H (ppm)
1	172.3	-
2	21.4	1.91

Astringin (trans)

$C_{20}H_{22}O_{9}$	406.4 g/mol	CAS: 29884-49-9
Atom number	¹³ C (ppm)	¹ H (ppm)
1	129.0	-
2	113.7	6.99
3	145.8	-
4	146.0	-
5	116.1	6.73
6	119.2	6.85
7	129.3	6.96
8	125.5	6.79
1'	139.8	-
2'	105.2	6.73
3'	159.3	-
4'	103.1	6.34
5′	158.7	-
6'	107.5	6.58
1"	101.0	4.81
2"	73.7	3.23
3"	77.1	3.28
4"	70.2	3.18
5″	77.5	3.37
6"	61.1	3.50/3.74

piceatannol-3'-O-glucoside (trans)



$C_{20}H_{22}O_{9}$	406.4 g/mol	CAS: 94356-26-0
Atom number	¹³ C (ppm)	¹ H (ppm)
1	139.6	-
2	104.8	6.40
3	158.9	-
4	102.3	6.14
5	158.9	-
6	104.8	6.40
7	126.8	6.85
8	128.2	6.91
1'	129.3	-
2'	114.6	7.45
3′	146.1	-
4'	147.0	-
5′	116.3	6.80
6′	122.4	7.06
1"	102.9	4.75
2"	73.8	3.34
3"	76.4	3.33
4"	70.5	3.18
5″	77.8	3.43
6"	61.3	3.50/3.79

glucosyl-frambinone



$C_{16}H_{22}O_7$	326.3 g/mol	CAS: 38963-94-9
Atom number	¹³ C (ppm)	¹ H (ppm)
1	30.2	2.09
2	208.2	-
3	44.7	2.72
4	28.7	2.72
1'	134.8	-
2'/6'	129.3	7.11
3′/5′	116.5	6.93
4'	156.0	-
1"	100.9	4.79
2"	73.5	3.22
3"	76.7	3.24
4"	70.0	3.15
5″	77.3	3.28
6"	60.9	3.47/3.68



$C_5H_{10}O_4$	134.1 g/mol	CAS 106-61-6
Atom number	¹³ C (ppm)	¹ H (ppm)
1	170.9	-
2	21.3	2.00
1'	63.1	3.33
2'	69.7	3.63
3'	66.0	3.88/4.02

glucosyl-trans-para-coumaric acid



$C_{15}H_8O_8$	326.3 g/mol	CAS 14364-05-7
Atom number	¹³ C (ppm)	¹ H (ppm)
1	168.3	-
2	117.4	6.41
3	144.1	7.59
1′	128.3	-
2'/6'	130.2	7.65
3'/5'	116.8	7.05
4'	159.3	-
1"	100.3	4.95
2"	73.6	3.26
3"	76.9	3.27
4"	69.9	3.16
5″	77.6	3.35
6"	61.1	3.46/3.68

$C_{16}H_{20}O_9$	356.3 g/mol	CAS: 177405-51-3
Atom number	¹³ C (ppm)	¹ H (ppm)
1	168.3	-
2	118.2	6.46
3	143.7	7.50
1'	128.5	-
2'	115.1	7.09
3′	122.3	7.17
4'	148.5	-
5′	149.3	-
6′	111.3	7.32
7'	55.9	3.82
1"	99.9	4.97
2"	73.4	3.27
3"	77.3	3.29
4"	70.0	3.16
5″	77.4	3.35
6"	60.9	3.45/3.66

larixyl acetate



$C_{22}H_{36}O_{3}$	348.5 g/mol	CAS: 4608-49-5
Atom number	¹³ C (ppm)	¹ H (ppm)
1	38.8	1.04/1.67
2	18.8	1.45/1.48
3	43.4	1.21/1.32
4	33.4	-
5	56.9	1.44
6	72.6	4.90
7	44.2	2.02/2.54
8	145.0	-
9	55.8	1.61
10	39.7	-
11	17.9	1.27/1.48
12	41.6	1.16/1.56
13	72.1	-
14	146.6	5.84
15	111.1	4.94/5.11
16	28.0	1.13
17	109.5	4.65/4.88
18	36.3	1.00
19	22.4	0.81
20	15.9	0.67
1'	169.8	-
2'	21.9	2.00

2-O-(4-Hydroxybenzoyl)-2,4,6-trihydroxyphenylacetic acid



$C_{15}H_{12}O_7$	304.3 g/mol	CAS: -
Atom number	¹³ C (ppm)	¹ H (ppm)
1	163.9	-
2	119.8	-
3/7	132.4	7.92
4/6	115.9	6.92
5	162.8	-
1'	172.7	-
2'	29.3	3.28
3'	106.1	-
4'	151.0	-
5'	101.0	6.09
6'	157.1	-
7'	100.0	6.24
8'	157.1	-

ferulic acid

$C_{10}H_{10}O_4$	194.2 g/mol	CAS: 1135-24-6
Atom number	¹³ C (ppm)	¹ H (ppm)
1	168.2	-
2	115.9	6.36
3	144.8	7.49
1'	126.0	-
2'	111.4	7.28
3′	148.2	-
4'	149.3	-
5'	115.9	6.8
6′	123.0	7.08
7'	56.0	3.82

larixinol isomer 1 + 2



$C_{30}H_{22}O_{10}$	542.5 g/mol	CAS: 101046-79-1
Atom number	¹³ C (ppm)	¹ H (ppm)
2	77.7 / 78.4	4.60 / 4.90
3	64.5 / <mark>63.7</mark>	4.02 / 4.23
4	28.6 / <mark>28.6</mark>	2.46/2.51 / 2.67/2.82
4a	103.4 / 103.8	-
5	156.7 / <mark>157.2</mark>	-
6	90.1 / <mark>90.7</mark>	6.08 / <mark>6.20</mark>
7	150.7 / <mark>151.8</mark>	-
8	104.2 / 104.4	-
8a	151.9 / <mark>151.7</mark>	-
9	129.6 / <mark>129.5</mark>	-
10/14	127.7 / <mark>127.9</mark>	6.93 / <mark>7.09</mark>
11/13	114.8 / <mark>114.8</mark>	6.69 / <mark>6.69</mark>
12	156.5 / <mark>156.7</mark>	-
2'	92.9 / <mark>88.9</mark>	5.72 / <mark>6.14</mark>
3′	60.1 / <mark>59.9</mark>	-
4'	178.7 / <mark>174.7</mark>	-
5′	154.1 / <mark>154.9</mark>	-
6'	96.1 / <mark>96.1</mark>	5.88 / <mark>5.79</mark>
7'	160.3 / <mark>160.2</mark>	-
8'	89.8 / <mark>89.4</mark>	5.90 / <mark>5.88</mark>
9'	162.1 / <mark>163.1</mark>	-
10'	104.9 / <mark>104.1</mark>	-
11'	126.2 / <mark>126.2</mark>	-
12'/16'	127.0 / <mark>127.3</mark>	6.90 / <mark>6.91</mark>
13'/15'	114.8 / <mark>115.2</mark>	6.52 / <mark>6.68</mark>
14'	157.2 / <mark>157.8</mark>	-

dianthoside



$C_{12}H_{16}O_8$	288.3 g/mol	CAS: 20847-13-6
Atom number	¹³ C (ppm)	¹ H (ppm)
2	161.8	-
3	142.1	-
4	174.6	-
5	116.5	6.44
6	156.0	8.14
7	15.6	2.37
1'	104.0	4.74
2'	74.3	3.15
3'	76.7	3.19
4'	69.9	3.10
5′	77.7	3.10
6′	31.3	3.44/3.64

arabinosyl-(1->6)-glucosyl-myrtenic acid



$C_{21}H_{32}O_{11}$	460.5 g/mol	CAS: -
Atom number	¹³ C (ppm)	¹ H (ppm)
1	164.1	-
2	138.9	-
3	138.6	6.91
4	32.1	2.41/2.48
5	40.0	2.11
6	37.5	-
7	40.9	2.70
8	31.1	1.03/2.46
9	25.9	1.31
10	21.2	0.76
1'	94.6	5.36
2'	72.8	3.16
3'	76.6	3.24
4'	70.1	3.08
5'	76.6	3.41
6'	67.2	3.38/3.88
1"	108.7	4.71
2"	82.3	3.79
3"	77.5	3.63
4"	84.0	3.70
5"	31.5	3.41/3.54

rhamnosyl-(1->6)-glucosyl-myrtenic acid



C ₂₂ H ₃₄ O ₁₁	474.5 g/mol	CAS: -
Atom number	¹³ C (ppm)	¹ H (ppm)
1	164.1	-
2	138.9	-
3	138.6	6.91
4	32.1	2.41/2.48
5	40.0	2.11
6	37.5	-
7	40.9	2.70
8	31.1	1.03/2.46
9	25.9	1.31
10	21.2	0.76
1'	94.7	5.37
2'	72.8	3.16
3'	76.6	3.24
4'	69.8	3.10
5'	76.7	3.36
6'	66.5	3.45/3.78
1"	100.8	4.53
2"	70.9	3.40
3"	70.7	3.59
4"	72.2	3.17
5"	68.7	3.42
6"	18.2	1.10

oleic acid

18



$C_{18}H_{34}O_2$	282.5 g/mol	CAS: 112-80-1
Atom number	¹³ C (ppm)	¹ H (ppm)
1	174.8	-
2	34.0	2.17
3	24.8	1.46
4-7	28.8-29.4	1.22-1.30
8	26.9	1.97
9	129.9	5.32
10	129.9	5.32
11	26.9	1.97
12-15	28.8-29.4	1.22-1.30
16	31.6	1.23
17	22.3	1.25
18	14.3	0.85



$C_{18}H_{32}O_2$	280.4 g/mol	CAS: 60-33-3
Atom number	¹³ C (ppm)	¹ H (ppm)
1	174.7	-
2	34.0	2.16
3	24.9	1.48
4	29-30	1.25
5	29-30	1.2-1.3
6	29-30	1.2-1.3
7	29-30	1.31
8	27.0	2.00
9	129.9	5.32
10	128.0	5.30
11	25.6	2.73
12	128.0	5.30
13	129.9	5.32
14	27.0	2.00
15	29-30	1.31
16	31.3	1.25
17	22.4	1.25
18	14.2	0.85



2

$C_{20}H_{34}O_1$	290.5 g/mol	CAS: 1438-62-6
Atom number	¹³ C (ppm)	¹ H (ppm)
1	38.7	1.05/1.80
2	19.2	1.44/1.50
3	42.0	1.14/1.36
4	33.5	-
5	55.1	1.08
6	24.2	1.24
7	38.0	1.81/2.32
8	148.5	-
9	56.9	1.49
10	39.6	-
11	17.9	1.48
12	41.6	1.12/1.56
13	72.0	-
14	146.6	5.83
15	111.1	4.93/5.11
16	27.9	1.12
17	106.9	4.52/4.79
18	21.8	0.77
19	33.7	0.85
20	14.6	0.62

isopimaric acid



$C_{20}H_{30}O_2$	302.5 g/mol	CAS: 5835-26-7
Atom number	¹³ C (ppm)	¹ H (ppm)
1	38.8	1.05/1.80
2	17.9	1.47
3	36.9	1.53/1.65
4	45.6	-
5	44.9	1.83
6	25.0	1.55/1.90
7	121.1	5.31
8	135.5	-
9	51.8	1.69
10	34.8	-
11	18.4	1.63/1.71
12	35.8	1.32/1.44
13	36.8	-
14	45.8	1.86/1.93
15	150.2	5.80
16	110.0	4.87/4.93
17	21.5	0.83
18	179.8	-
19	17.5	1.16
20	15.3	0.85

dehydroabietic acid



$C_{20}H_{28}O_2$	300.4 g/mol	CAS: 1740-19-8
Atom number	¹³ C (ppm)	¹ H (ppm)
1	38.1	1.31/2.30
2	18.4	1.63/1.72
3	36.6	1.56/1.68
4	46.7	-
5	45.0	2.02
6	21.4	1.41/1.75
7	29.8	2.75/2.82
8	134.4	-
9	147.1	-
10	36.7	-
11	124.3	7.15
12	124.0	6.97
13	145.3	-
14	126.7	6.84
15	33.2	2.77
16	24.3	1.16
17	24.3	1.16
18	179.7	-
19	16.7	1.16
20	25.1	1.13

7-oxodehydroabietic acid



$C_{20}H_{26}O_3$	314.4 g/mol	CAS: 18684-55-4
Atom number	¹³ C (ppm)	¹ H (ppm)
1	37.3	1.47/2.37
2	Not detected	Not detected
3	37.2	1.49/2.37
4	46.0	-
5	44.3	2.53
6	37.9	2.24/2.72
7	198.2	-
8	130.5	-
9	153.8	-
10	37.4	-
11	124.5	7.41
12	132.8	7.50
13	146.5	-
14	124.0	7.70
15	33.2	2.92
16	23.9	1.20
17	23.9	1.20
18	179.8	-
19	16.8	1.20
20	23.9	1.20