

Date : 2024-04-26

CERTIFICATE OF ANALYSIS - GC PROFILING

SAMPLE IDENTIFICATION

Internal code : 24D12-PTH08

Customer Identification : Ho Wood - China - H80109R

Type : Essential Oil

Source : *Cinnamomum camphora* ct. Linalool

Customer : Plant Therapy

Checked and approved by:

Alexis St-Gelais, Ph. D., Chimiste 2013-174

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GAS CHROMATOGRAPHIC ANALYSIS

Method : PC-MAT-014 - Analysis of the composition of an essential oil or other volatile liquide by FAST GC-FID

*ISO

Results : See analysis summary (next page)

Analyst : Sylvain Mercier, M. Sc., Chimiste 2014-005

Date : 2024-04-22

PHYSICOCHEMICAL DATA

Refractive index : 1.4622 ± 0.0003 (20 °C)

Method : PC-MAT-016 - Measure of the refractive index of a liquid.

Analyst : Cindy Caron B. Sc.

Date : 2024-04-12

CONCLUSION

No adulterant, contaminant or diluent has been detected using this method.

ANALYSIS SUMMARY - CONSOLIDATED CONTENTS

New readers of similar reports are encouraged to read table footnotes at least once.

Identification	%	Class
Sabinene	tr	Monoterpene
β-Pinene	tr	Monoterpene
Myrcene	0.03	Monoterpene
α-Phellandrene	0.01	Monoterpene
α-Thujene	tr	Monoterpene
α-Terpinene	0.01	Monoterpene
para-Cymene	0.01	Monoterpene
Limonene	0.03	Monoterpene
β-Phellandrene	0.03	Monoterpene
1,8-Cineole	0.01	Monoterpenic ether
(Z)-β-Ocimene	0.02	Monoterpene
(E)-β-Ocimene	0.02	Monoterpene
γ-Terpinene	0.01	Monoterpene
cis-Sabinene hydrate	0.08	Monoterpenic alcohol
cis-Linalool oxide (fur.)	0.13	Monoterpenic alcohol
Geranic oxide	tr	Monoterpenic ether
Fenchone	0.01	Monoterpenic ketone
trans-Linalool oxide (fur.)	0.18	Monoterpenic alcohol
Terpinolene	0.02	Monoterpene
Linalool	99.21	Monoterpenic alcohol
cis-para-Menth-2-en-1-ol	0.03	Monoterpenic alcohol
α-Campholenal	0.01	Monoterpenic aldehyde
Camphor	0.06	Monoterpenic ketone
Hodiendiol (2,6-dimethylocta-3,7-diene-2,6-diol)	0.01	Monoterpenic alcohol
Unknown	0.02	Unknown
Consolidated total	99.96	

tr: The compound has been detected below 0.005% of the total signal

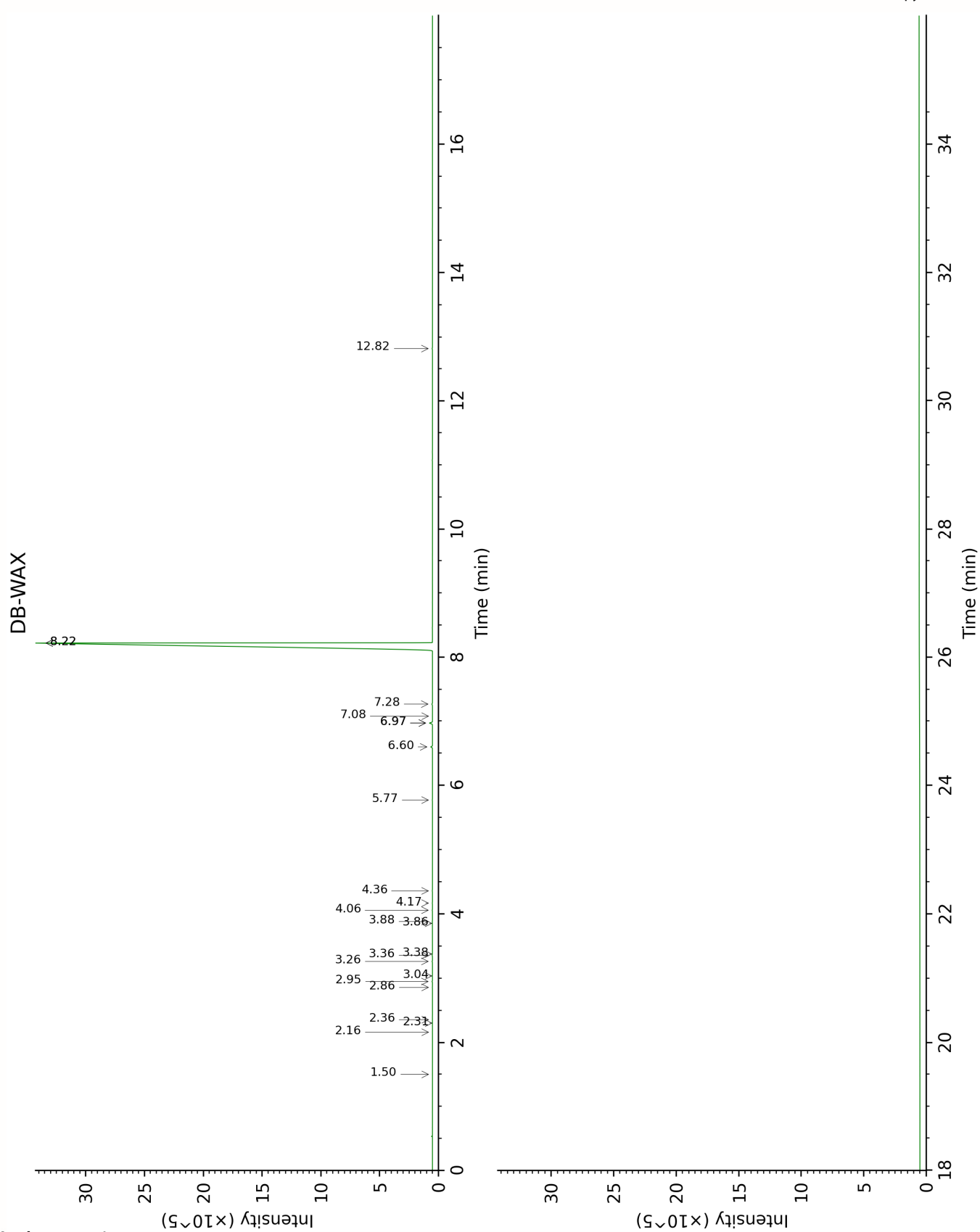
Note: no correction factor was applied

About "consolidated" data: The table above presents the breakdown of the sample volatile constituents after applying an algorithm to collapse data acquired from the multi-columns system of PhytoChemia into a single set of consolidated contents. In case of discrepancies between columns, the algorithm is set to prioritize data from the most standard DB-5 column, and smallest values so as to avoid overestimating individual content. This process is semi-automatic. Advanced users are invited to consult the "Full analysis data" table after the chromatograms in this report to access the full untreated data and perform their own calculations if needed.

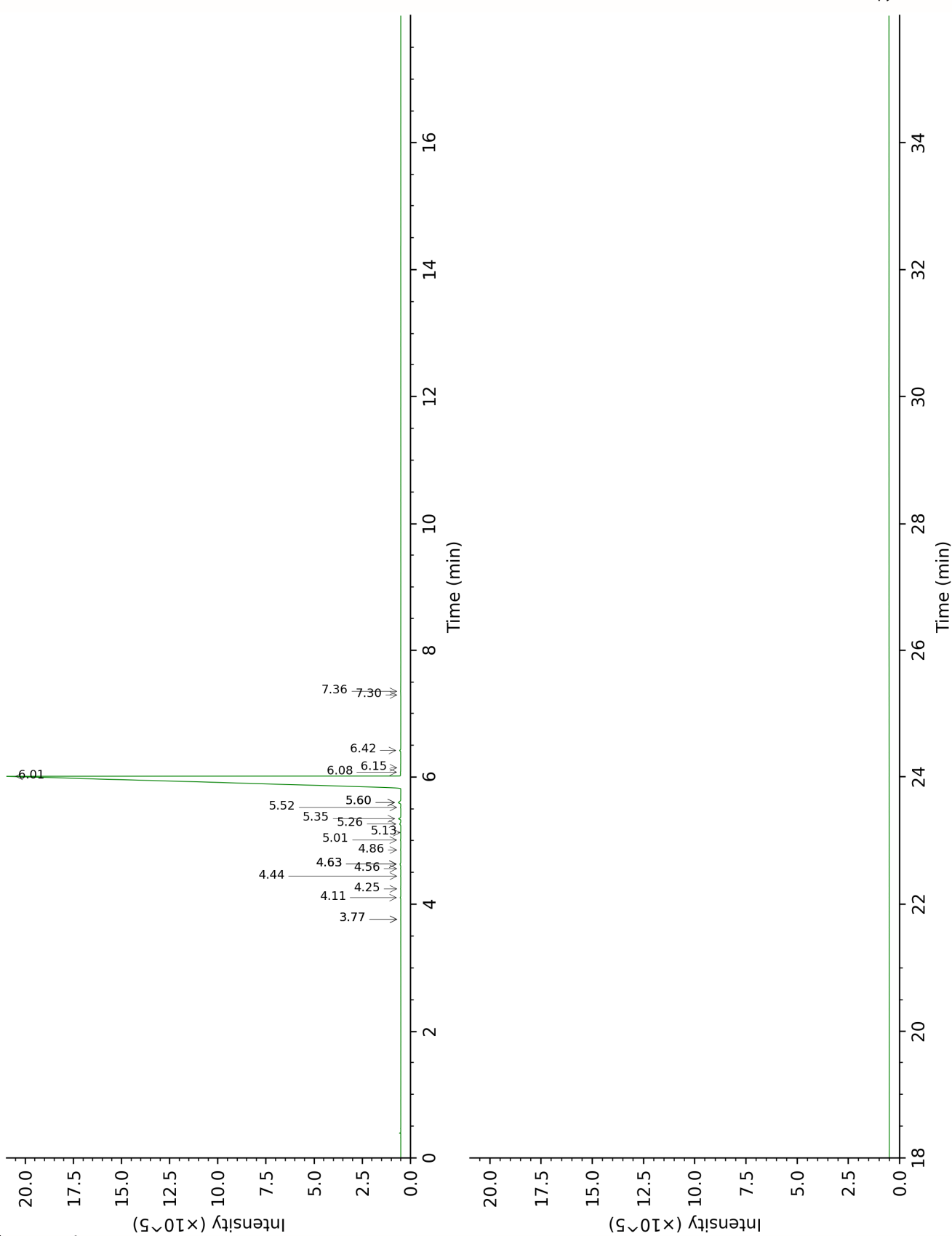
Unknowns: Unknown compounds' mass spectral data is presented in the "Full analysis data" table. The occurrence of unknown compounds is to be expected in many samples, and does not denote particular problems unless noted otherwise in the conclusion.

Bracketed value ([xx]): A bracketed percent value indicate that two or more compound percentage could not be solved due to coelution.

This page was intentionally left blank. The following pages present the complete data of the analysis.



DB-5



FULL ANALYSIS DATA

Sabinene	Column DB-WAX			Column DB-5		
	2.36	1086.3	tr	3.77*	970.9	[0.01]
β-Pinene	2.16	1068.0	tr	3.77*	970.9	[0.01]
Myrcene	2.95	1135.3	0.03	4.11	993.4	0.03
α-Phellandrene	2.86	1128.4	0.01	4.24	1002.5	0.01
α-Thujene	1.50	1003.3	tr			
α-Terpinene	3.04	1141.9	0.01	4.44	1015.1	0.01
para-Cymene	4.17	1226.3	0.02	4.56	1022.5	0.01
Limonene	3.26	1159.0	0.03	4.63*	1027.0	[0.06]
β-Phellandrene	3.36	1166.1	0.03	4.63*	1027.0	[0.06]
1,8-Cineole	3.38	1167.7	0.01	4.63*	1027.0	[0.06]
(Z)-β-Ocimene	3.86	1203.8	0.02	4.86	1040.8	0.02
(E)-β-Ocimene	4.06	1218.5	0.02	5.01	1050.6	0.02
γ-Terpinene	3.88	1206.0	0.01	5.13	1058.0	0.01
cis-Sabinene hydrate	6.97*	1428.2	[0.27]	5.26	1066.4	0.08
cis-Linalool oxide (fur.)	6.60	1400.6	0.14	5.35	1071.7	0.13
Geranic oxide	2.30	1081.3	tr			
Fenchone	5.77	1340.9	0.02	5.52	1082.7	0.01
trans-Linalool oxide (fur.)	6.97*	1428.2	[0.27]	5.60*	1087.6	[0.20]
Terpinolene	4.36	1239.8	0.02	5.60*	1087.6	[0.20]
Linalool	8.22*	1523.2	[99.07]	6.01	1113.5	99.21
cis-para-Menth- 2-en-1-ol	8.22*	1523.2	[99.07]	6.08	1117.6	0.03
α-Campholenal	7.08	1436.2	0.01	6.15	1122.0	0.01
Camphor	7.28	1451.1	0.06	6.42	1139.4	0.06
Hodiendiol (2,6- dimethylocta- 3,7-diene-2,6- diol)	12.82	1905.0	0.04	7.30	1195.7	0.01
Unknown SASC VI [m/z 43, 71 (80), 67 (55), 59 (51), 68 (44), 41 (43)...]				7.36	1199.4	0.02
Total reported	99.82%			99.94%		

*: Two or more compounds are coeluting on this column

[xx]: Duplicate percentage due to coelutions, only the first one is taken into account in the consolidated total

†: Peaks apexes were resolved, but peaks overlapped and were summed for analysis

tr: The compound has been detected below 0.005% of total signal.

Essential Oil, *Cinnamomum camphora* ct. Linalool
Internal code: 24D12-PTH08

Ho Wood - China - H80109R

Report prepared for:
Plant Therapy

Note: no correction factor was applied
R.T.: Retention time (minutes)
R.I.: Retention index

Laboratoire
PhytoChemia

Plus que des analyses... des conseils

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