

NMR metabolite profiling for a fast characterization of Vessalico garlic ecotype and bioactivity against *Xanthomonas campestris* pv. *campestris*

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1 Supplementary Figures and Tables

1.1 Supplementary Figures

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1.2 Supplementary Tables

Table S1. List of garlic accessions used in the study.

Table S2. ¹H NMR chemical shifts (δ) and coupling constants (Hz) of Chenomx 600 MHz library (CL) and custom library (CCL) metabolites.

1.1 Supplementary Figures

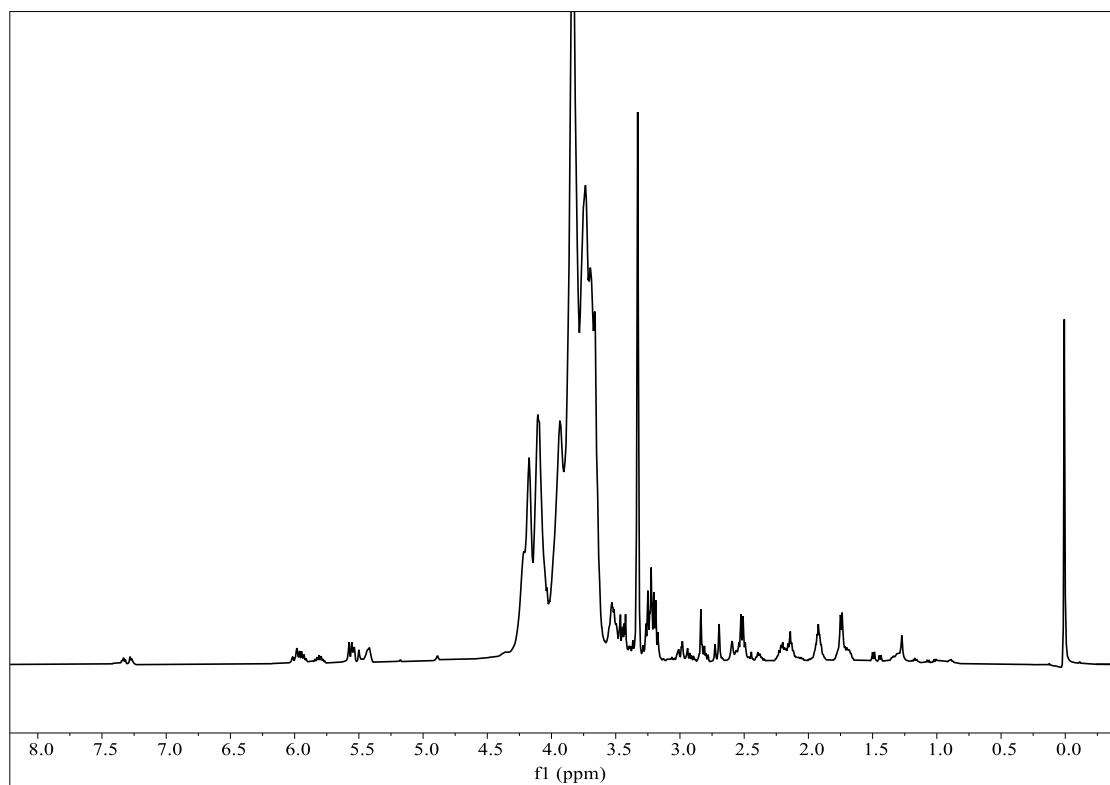


Figure S1. Representative ^1H -NMR spectrum of Vessalico garlic.
Accession 12 (Table S2) clove extract in $\text{CD}_3\text{OD-KH}_2\text{PO}_4$ in D_2O at pH 6.0, 600 MHz.

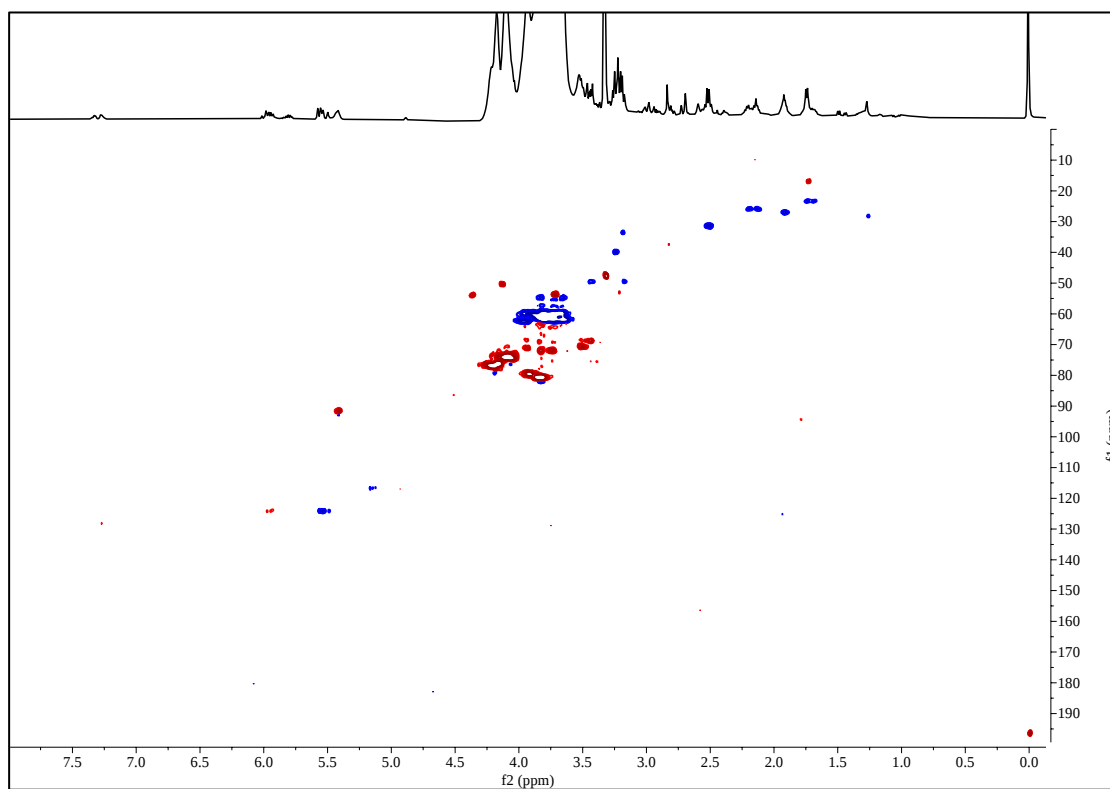


Figure S2. Representative HSQC spectrum of Vessalico garlic.
Accession 12 (Table S2) clove extract in $\text{CD}_3\text{OD-KH}_2\text{PO}_4$ in D_2O at pH 6.0, 600 MHz.

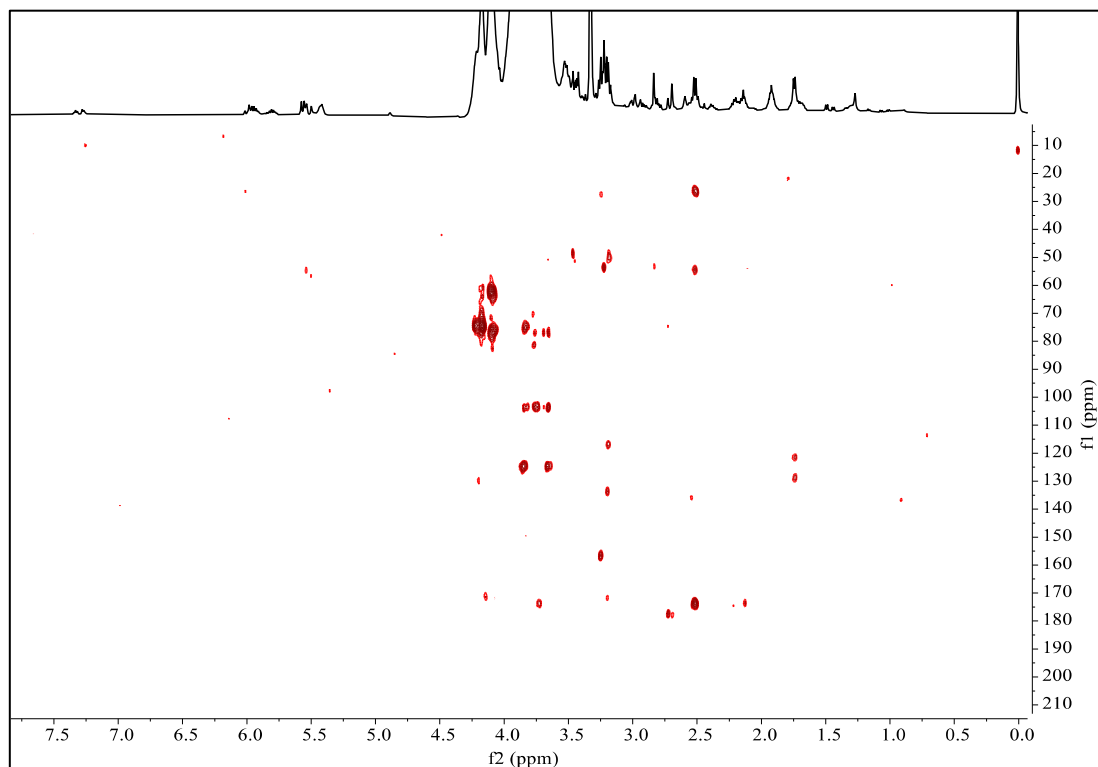


Figure S3. Representative HMBC spectrum of Vessalico garlic.
Accession 12 (Table S2) clove extract in CD₃OD-KH₂PO₄ in D₂O at pH 6.0, 600 MHz.

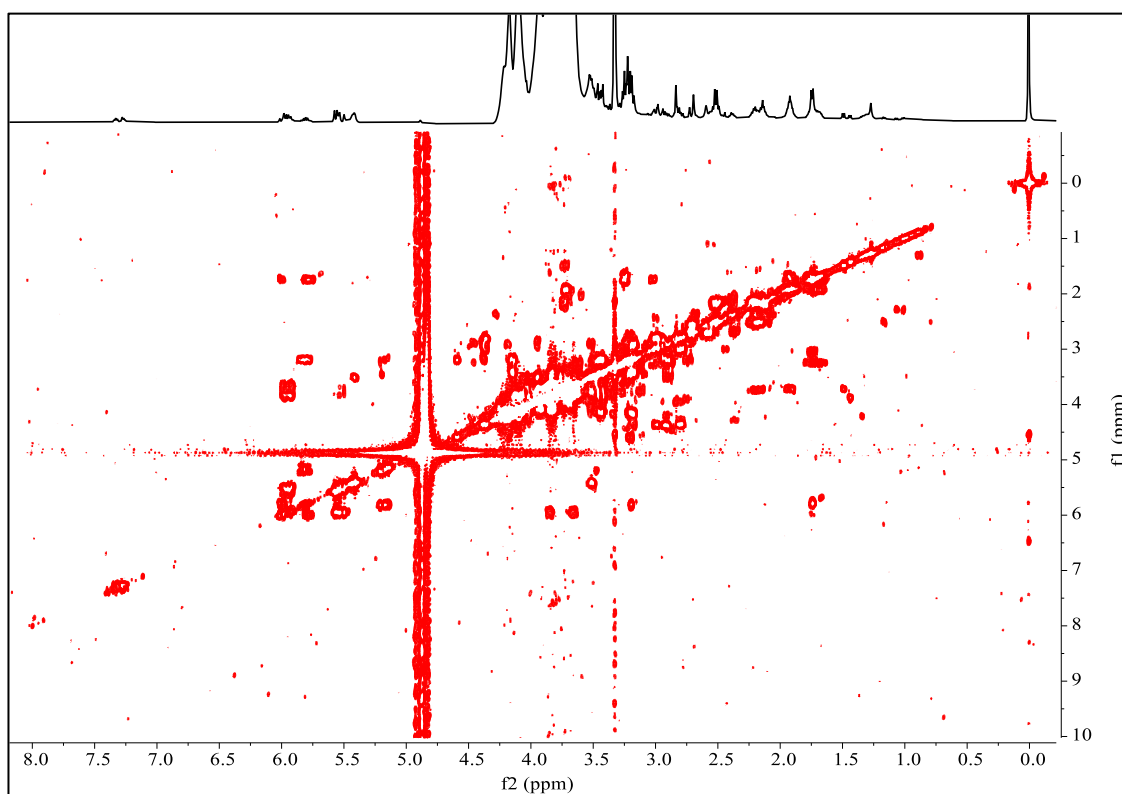


Figure S4. Representative COSY spectrum of Vessalico garlic.
Accession 12 (Table S2) clove extract in CD₃OD-KH₂PO₄ in D₂O at pH 6.0, 600 MHz.

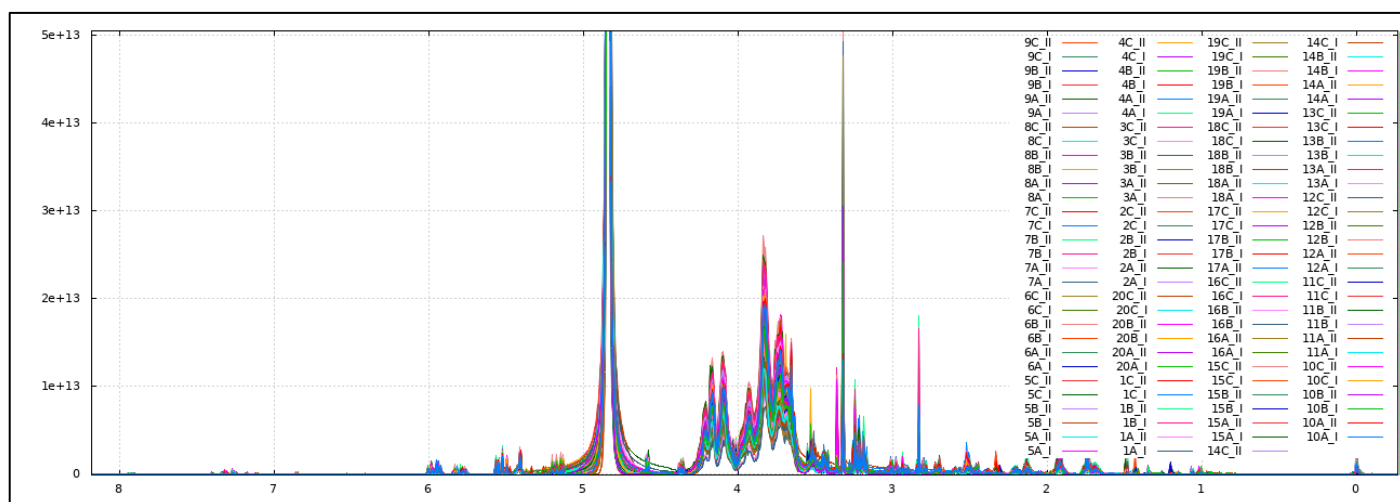


Figure S5. ^1H -NMR spectra of garlic accessions.

Spectra of the garlic extracts using the open access software NMRProcFlow v1.4.14. Spectra of the extracts are presented in different colours.

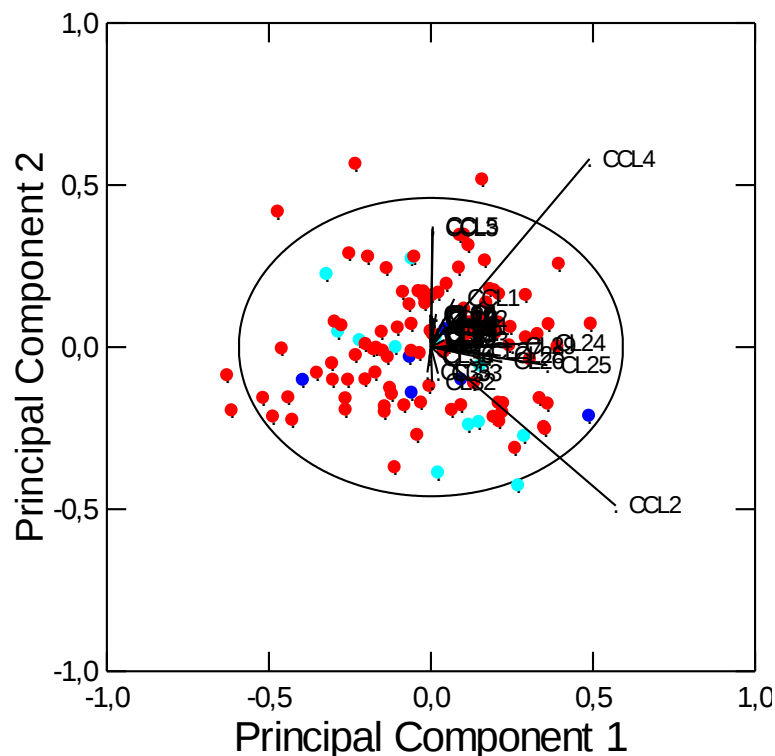


Figure S6. Results of PCA. CL and CCL variables.

The biplot shows the scores of the 120 spectra and the loadings of the 41 variables (Chenomx 600 MHz 36 library and 5 custom library metabolites) on Principal Components 1 and 2 (explaining 46.2% and 27.8% of the total variance, respectively): ● Vessalico; ● Messidor; ● Messidrôme). The ellipse represents the 95% confidence interval.

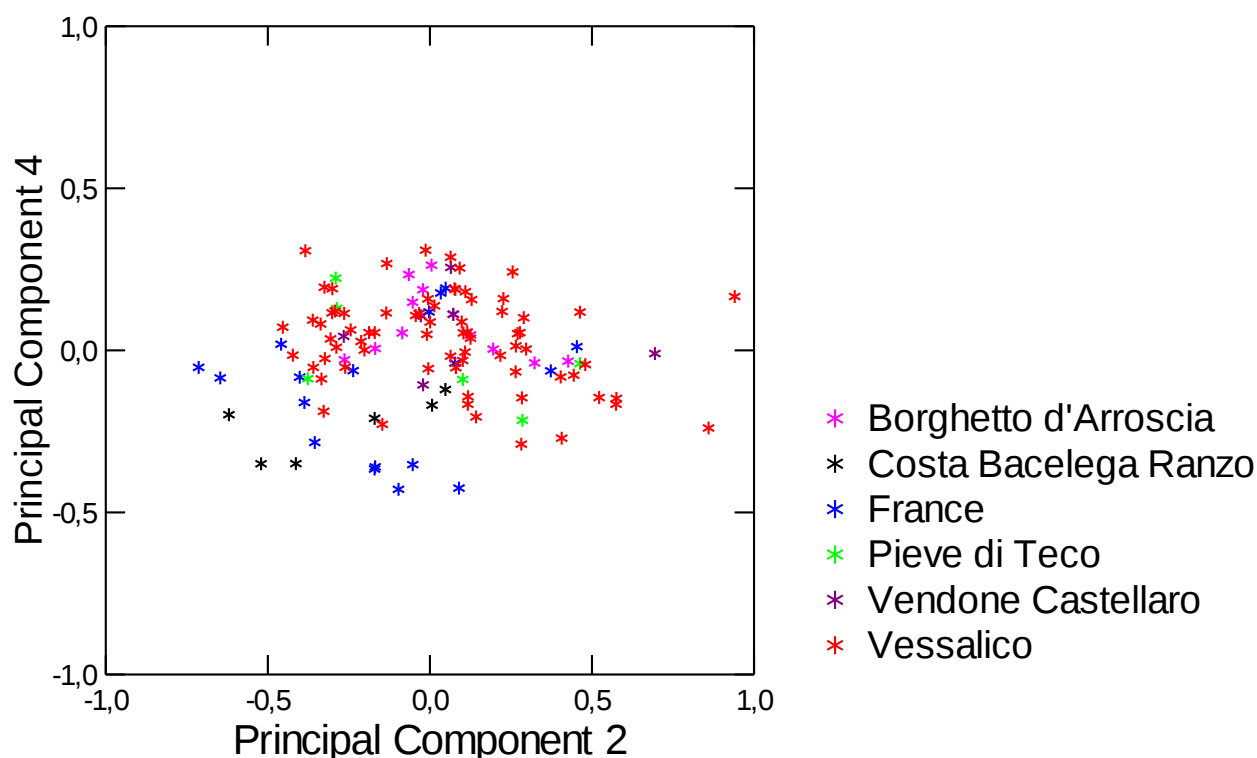


Figure S7. Results of PCA applied to garlic accessions from different locations and farms.

Plot of the scores of the samples described by Chenomx 600 MHz 36 library and 5 custom library metabolites (CL and CCL, Table S2) on Principal Components 2 and 4.

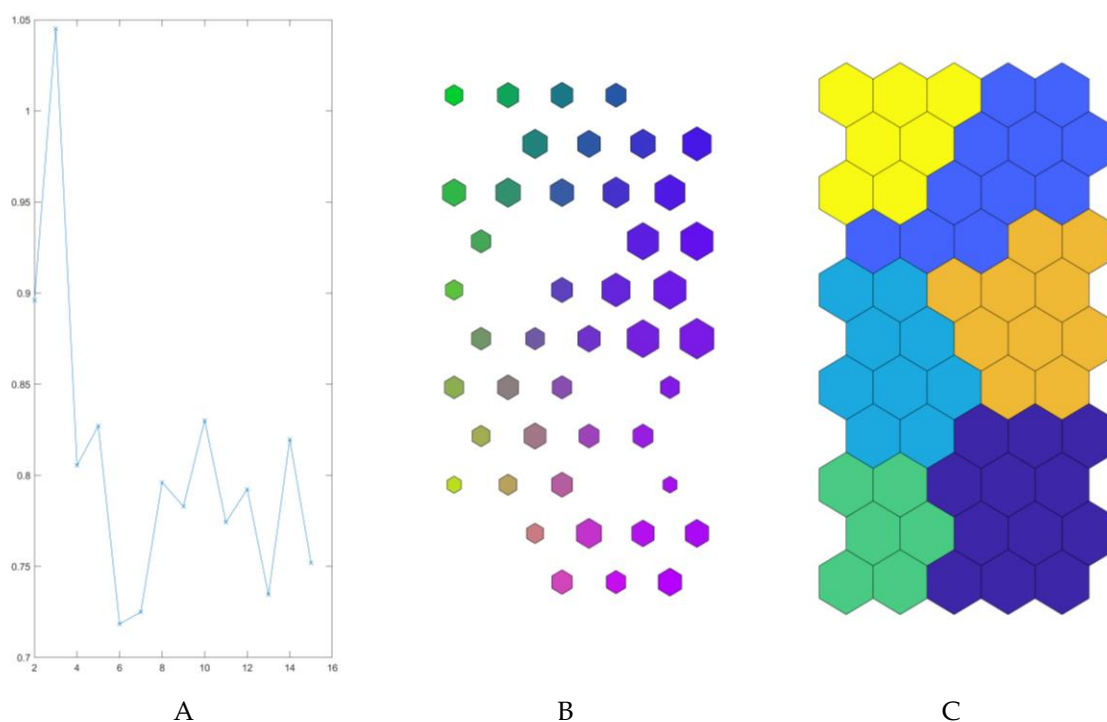


Figure S8. Results of SOMs. Map clusterization for CL and CCL variables.

Chenomx 600 MHz 36 library and 5 custom library metabolites (Table S2). A. Davies-Bouldine index progression: minimum value fits the best number of clusters. B. SOM output map with colour code association. Similar colours have similar characteristics. C. 6 clusters.

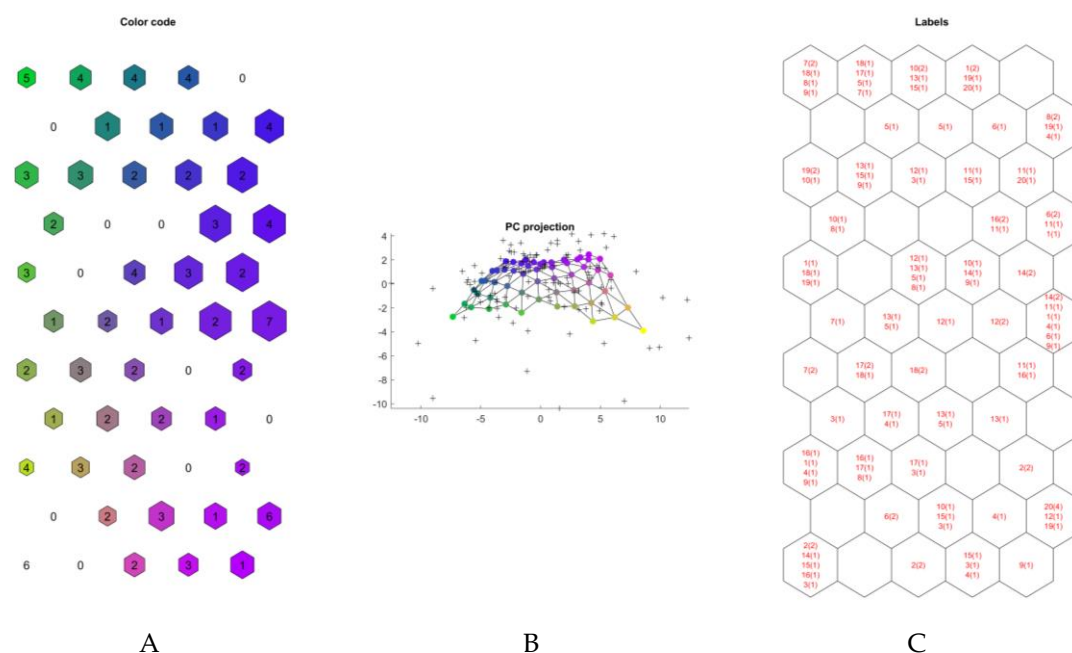


Figure S9. Results of SOMs. Graphical representation of map for CL and CCL variables.

Chenomx 600 MHz 36 library and 5 custom library metabolites (Table S2). A. SOM output map with colour code association. Similar colours have similar characteristics, numbers correspond to hit numbers. Dimensions of hexagons are related to the distance between neurons (biggest indicates greater distance); B. Principal Component projection of the map; C. labelled SOM output map, for each neuron the corresponding accession number (Table S1) and number of replicates (in parentheses) are shown.

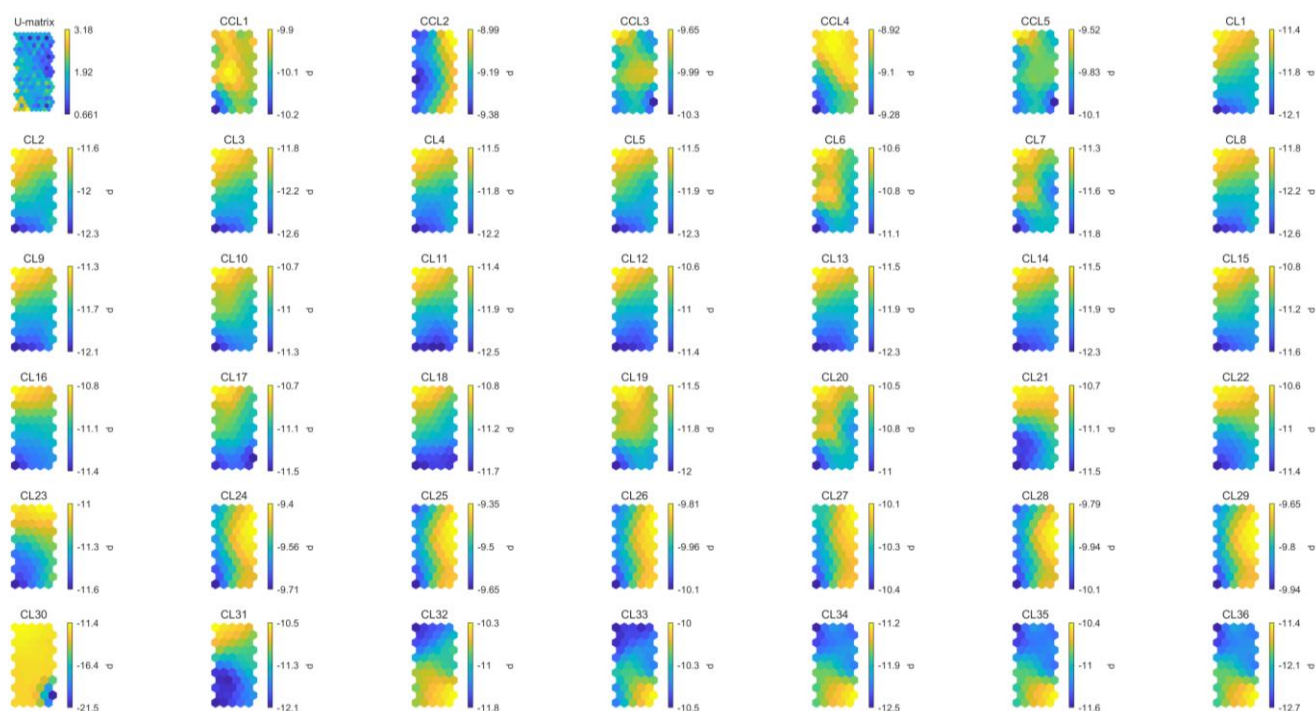


Figure S10. Results of SOMs. U-matrix and maps for each CL and CCL variables.

Chenomx 600 MHz 36 library and 5 custom library metabolites (Table S2). Similar color gradations indicate highly correlated variables.

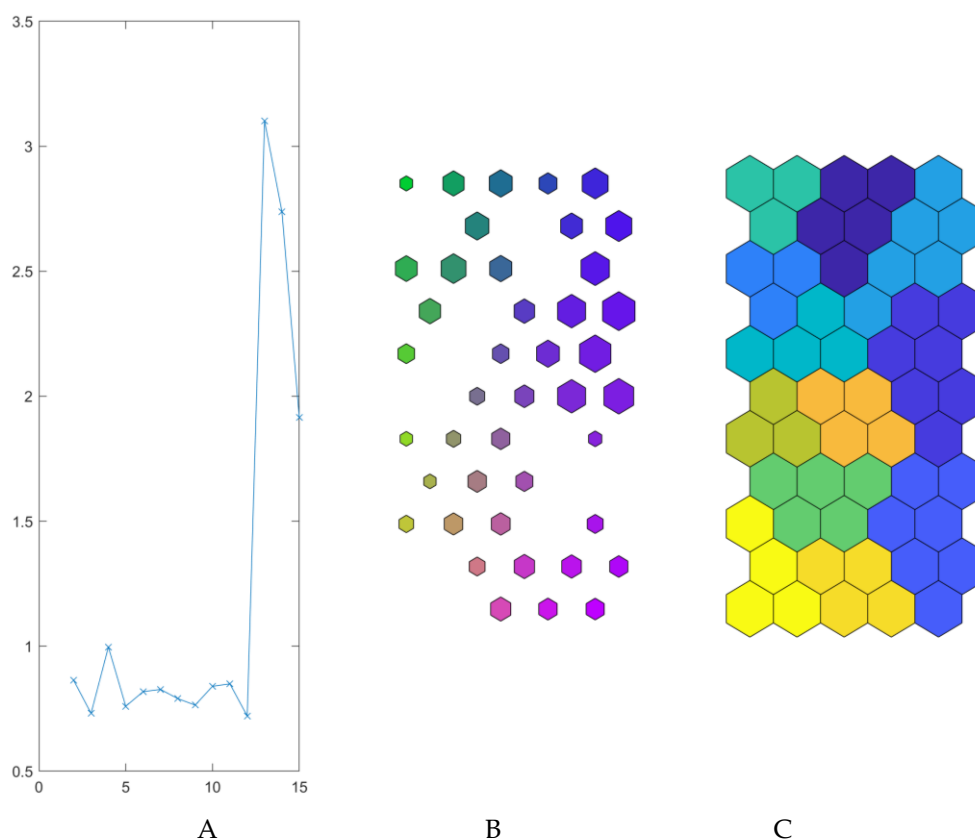


Figure S11. Results of SOMs. Map clusterization for CL variables.

Chenomx 600 MHz 36 library metabolites (Table S2). A. Davies-Bouldine index progression: minimum value fits the best number of clusters. B. SOM output map with colour code association. Similar colours have similar characteristics. C. 12 clusters.

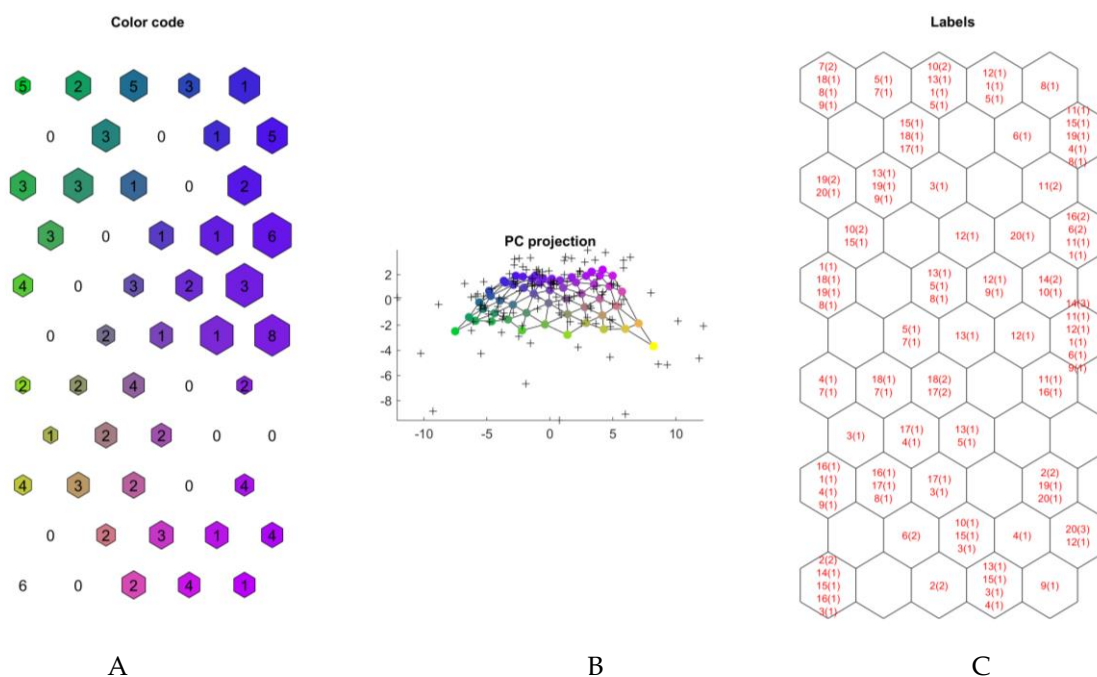


Figure S12. Results of SOMs. Graphical representation of map for CL variables.

Chenomx 600 MHz 36 library metabolites (Table S2). A. SOM output map with color code association. Similar colours have similar characteristics, numbers correspond to hit numbers. Dimensions of hexagons are related to the distance between neurons (biggest indicates greater distance); B. Principal Component projection of the map; C. labelled SOM output map, for each neuron the corresponding accession number (Table S1) and number of replicates (in parentheses) are shown.

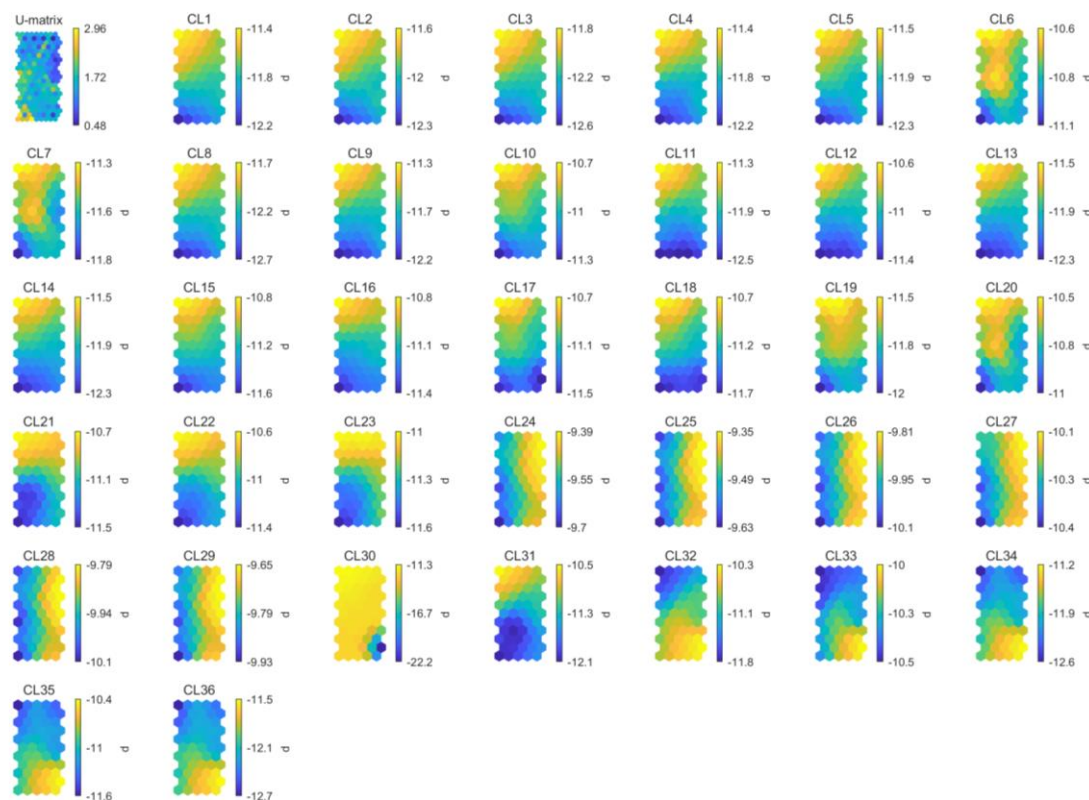


Figure S13. Results of SOMs. U-matrix and maps for each CL variable.

Chenomx 600 MHz 36library metabolites (Table S2). Similar color gradations indicate highly correlated variables.

1.2 Supplementary Tables

Table S1. List of garlic accessions used in the study.

Accession number	Ecotype or Cultivar name	Grower	Location	Breeding status (*)	Soil texture type	Yield (ton x ha ⁻¹)
1	Vessalico	A	Costa Bacelega Ranzo (Imperia, Italy) ^a	Ecotype	sandy clay loam soil	80 (**)
2	Vessalico	B	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
3	Vessalico	C	Borghetto d'Arroschia (Imperia, Italy) ^c	Ecotype	sandy clay loam soil	80 (**)
4	Vessalico	D	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
5	Vessalico	E	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
6	Vessalico	F	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
7	Vessalico	G	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
8	Vessalico	H	Vendone fraz. Castellaro (Imperia, Italy)	Ecotype	sandy clay loam soil	80 (**)
9	Vessalico	I	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
10	Vessalico	J	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
11	Vessalico	K	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
12	Vessalico	L	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
13	Vessalico	M	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
14	Vessalico	N	Borghetto d'Arroschia (Imperia, Italy) ^c	Ecotype	sandy clay loam soil	80 (**)
15	Vessalico	O	Pieve di Teco (Imperia, Italy) ^d	Ecotype	sandy clay loam soil	80 (**)
16	Vessalico	P	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
17	Vessalico	Q	Vessalico (Imperia, Italy) ^b	Ecotype	sandy clay loam soil	80 (**)
18	Messidor	R	France	Released variety	-	9-12 (***)
19	Messidrôme	S	France	Released variety	-	8-14 (***)
20	Messidor	T	France	Released variety	-	9-12 (***)

^a Coordinates of the downtown of the village: 44°04'51"N; 8°01'51"E; elevation 450 (m.a.s.l).

^b Coordinates of the downtown of the village: 44°02'46"N; 7°57'43"E; elevation 197 (m.a.s.l).

^c Coordinates of the downtown of the village: 44°03'26"N; 7°58'54"E; elevation 155 (m.a.s.l).

^d Coordinates of the downtown of the village: 44°02'52"N; 7°54'57"E; elevation 240 (m.a.s.l).

* Landrace / Farmers' variety / Breeding line / Released variety/ ecotype.

** Maximum production capacity; only the cultivated area (excluding passages and other cultivation tares); included the fresh aerial part of the plant (foliage).

*** From breeder catalogue (INRA). Excluded aerial part of the plant; included passages and cultivation tares.

Cultivars	Foliage Colour **	Foliage Attitude ***	Bulb skin color ****	Bulb (cm)	Bulb Weight(g)	Shape	Colour of tunic	No. of bulblets	Clove size (cm)	Clove Weight (g)	Garlic flavor (+)
Vessalico*	Dark green	Erect	White	3,5-8	50-100	Oval	White/cream	10-15	3-5	5-10	+
Messidor°	Dark green	Erect	White	8-10	40-60	Oval	White/cream	10-15	2-3	2-5	++
Messidrôme°	Dark green	Erect	White	12-15	>60	Oval	White/cream	10-15	2-3	3-5	++

* From PGI production specification.

° From breeder catalogue (INRA).

** Dark green / Light green.

*** Erect / Droop.

****Purple tinge / White.

The best climatic conditions for garlic cultivation in Liguria are found in the inland areas, away from the sea, but characterized by a Mediterranean climate: a limited day-night temperature range between 5 and 10 °C; relative humidity of the air between 50 and 75%; rainfall concentrated mainly in the spring period (February-April), before maturation, and in autumn (September-October), before bulb planting.

The soil, from a chemical-physical point of view, is characteristic of the maritime slopes of the Ligurian Apennines and Alps: calcareous, rich in skeletal material, with a high clay and sandy component, with a modest water supply, especially in the period approaching harvest.

All these conditions, combined with the adoption of cultivation techniques refined over time by generations of farmers, have converged in a mountainous area behind Imperia (Italy) and identifiable with the territory of Vessalico (IM) and other 10 municipalities of the Arroscia Valley (IM): Vessalico, Aquila d'Arroscia, Armo, Borghetto d'Arroscia, Cosio d'Arroscia, Mendatica, Montegrosso Pian Latte, Pornassio, Pieve di Teco, Ranzo, Rezzo.

The garlic cultivated in that area has always been recognized under the name of the municipality of Vessalico which, due to its strategic position at the crossroads of ancient trade routes, has always been the center of trade for this product. The cultivation technique of Vessalico garlic is detailed in the Vessalico Garlic Specification for Protected Geographical Indication (PGI) candidacy. Below is a summary.

Vessalico garlic is cultivated in a hilly environment characterized by small, terraced plots, typical of the Ligurian hinterland. The soil has a medium texture and a good presence of skeletal material. Crop rotation is essential to avoid phytosanitary problems, and it is recommended to alternate garlic with other crops for at least two cycles. Vessalico garlic can be cultivated using both conventional and organic or biodynamic methods, complying with current regulations. Soil preparation occurs in the summer through plowing and refinement, with the possible addition of organic matter. Planting is preferably done in autumn, by February, and can be done manually or with specific machinery. It is important to follow the indications for planting depth and spacing. After planting, one or two spring hoeing operations are necessary to eliminate weeds, preferably with eco-sustainable techniques. Emergency irrigation is allowed in case of drought, but it is advisable to limit water supply before harvesting to promote bulb maturation. Fertilization can be done with mature organic substances or mineral fertilizers, depending on crop needs. Harvesting takes place when garlic leaves begin to yellow and dry, and it can be done manually or mechanically. Plants are then selected based on size and subjected to a slow drying period to preserve quality. Vessalico garlic is then handcrafted into characteristic "reste" (braids) and stored in cool, dry environments until commercialization.



Table S2. ¹H NMR chemical shifts (δ) and coupling constants (Hz) of Chenomx 600 MHz library (CL) and custom library (CCL) metabolites.

NMR spectra of CCL metabolites were recorded in CD₃OD-KH₂PO₄ in D₂O at pH 6.0, 600 MHz.

CCL/CL number	metabolite	selected characteristic NMR chemical shifts in ppm - cluster midpoint (multiplicity, J)	other chemical shifts in ppm - cluster midpoint (multiplicity, J)	MSI Status ^b	references
CCL1	S-methyl-L-cysteine	2.17 (3H, s)	2.97 ^a , 3.86 ^a	1	(Higuchi et al., 2003)
CCL2	methiin	2.83 (3H, s)	3.25 ^a , 3.48 ^a , 4.13 ^a	1	(Hibi et al., 2013; Ingallina et al., 2023)
CCL3	S-allyl-L-cysteine	5.17 (1H) ^a	2.92 ^a , 3.10 ^a , 3.23 ^a , 3.77 ^a , 5.24 ^a , 5.83 ^a	1	(Higuchi et al., 2003; Maldonado et al., 2011)
CCL4	L-alliin	5.50 (1H) ^a	3.21 ^a , 3.66 ^a , 3.82 ^a , 4.09 ^a , 5.53 ^a , 5.96 ^a	1	(Hibi et al., 2013)
CCL5	allicin	5.88 (1H) ^a	3.13 ^a , 3.32 ^a , 3.37 ^a , 3.60 ^a , 5.28 ^a	1	(Pacholczyk-Sienicka et al., 2024; Ritota et al., 2012)
CL1	leucine	0.95 (3H) ^a	0.94 ^a , 1.67 ^a , 1.71 ^a , 1.73 ^a , 3.73 ^a	2	(Le Mao et al., 2021; Liang et al., 2015; Pacholczyk-Sienicka et al., 2024; Verpoorte et al., 2007)
CL2	isoleucine	1.00 (3H, d, J = 7.0)	0.93 ^a , 1.25 ^a , 1.46 ^a , 1.97 ^a , 3.66 ^a	2	(Le Mao et al., 2021; Liang et al., 2015; Pacholczyk-Sienicka et al., 2024)
CL3	valine	1.03 (3H, d, J = 7.0)	0.98 ^a , 2.26 ^a , 3.60 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024; Verpoorte et al., 2007)
CL4	threonine	1.32 (3H, d, J = 6.6)	3.58 ^a , 4.25 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024; Verpoorte et al., 2007)
CL6	alanine	1.47 (3H, d, J = 7.2)	3.77 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024)
CL5	lysine	1.71 (2H) ^a	1.43 ^a , 1.50 ^a , 1.88 ^a , 1.91 ^a , 3.02 ^a , 3.75 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024)
CL7	acetic acid	1.91 (3H) ^a		2	(Fulmer et al., 2010)
CL8	acetamide	1.99 (3H) ^a		2	(Matsuoka et al., 2015)
CL9	homoserine	2.01 (2H) ^a	2.14 ^a , 3.77 ^a , 3.85 ^a	2	(Jamieson et al., 2009; Wang et al., 2008)
CL10	glutamine	2.11 (1H) ^a	2.14 ^a , 2.43 ^a , 2.46 ^a , 3.77 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024; Verpoorte et al., 2007)
CL11	succinylacetone	2.26 (3H) ^a	2.41 ^a , 2.81 ^a , 3.85 ^a	2	(Bal et al., 2009)
CL12	glutamic acid	2.33 (1H) ^a	2.04 ^a , 2.12 ^a , 2.36 ^a , 3.75 ^a	2	(Liang et al., 2015; Lucas-Torres et al., 2018; Sabino et al., 2019; Verpoorte et al., 2007)
CL13	pyruvic acid	2.36 (3H) ^a		2	(Bourafai-Aziez et al., 2022; Le Mao et al., 2021)
CL14	succinic acid	2.39 (4H) ^a		2	(Gaviglio and Doctorovich, 2008; Kostidis et al., 2017)
CL15	riboflavin	2.48 (3H) ^a	2.58 ^a , 3.73 ^a , 3.87 ^a , 3.93 ^a , 3.97 ^a , 4.44 ^a , 4.95 ^a , 5.14 ^a , 7.96 ^a , 7.98 ^a	2	(Jiménez-Nava et al., 2023)
CL16	citric acid	2.69 (2H) ^a	2.53 ^a	2	(Le Mao et al., 2021; Pacholczyk-Sienicka et al., 2024; Verpoorte et al., 2007)
CL17	aspartic acid	2.80 (1H) ^a	2.67 ^a , 3.89 ^a	2	(Liang et al., 2015; Pacholczyk-Sienicka et al., 2024)
CL18	asparagine	2.94 (1H) ^a	2.85 ^a , 4.00 ^a	2	(Liang et al., 2015; Lucas-Torres et al., 2018; Pacholczyk-Sienicka et al., 2024)
CL19	choline	3.19 (9H, s)	3.51 ^a , 4.06 ^a	2	(Le Mao et al., 2021; Liang et al., 2015)
CL20	arginine	3.24 (2H) ^a	1.64 ^a , 1.72 ^a , 1.89 ^a , 1.92 ^a , 3.76 ^a	2	(Nilsson et al., 2004)
CL21	cystine	3.38 (2H) ^a	3.18 ^a , 4.11 ^a	2	(Lucas-Torres et al., 2018)
CL22	proline	3.41 (1H) ^a	1.98 ^a , 2.02 ^a , 2.06 ^a , 2.34 ^a , 3.33 ^a , 4.12 ^a	2	(Le Mao et al., 2021; Liang et al., 2015; Verpoorte et al., 2007)
CL23	glycine	3.55 (2H, s)		2	(Kostidis et al., 2017)

					Supplementary Materials
CL24	gluconic acid	3.77 (1H) ^a	3.66 ^a , 3.75 ^a , 3.82 ^a , 4.03 ^a , 4.13 ^a	2	(Horton et al., 1983; Ramos et al., 2000)
CL25	methionine	3.85(1H) ^a	2.11 ^a , 2.13 ^a , 2.19 ^a , 2.63 ^a	2	(Kostidis et al., 2017)
CL26	serine	3.94 (1H)	3.84 ^a , 3.98 ^a	2	(Kostidis et al., 2017)
CL27	fructose	4.01(2H) ^a	3.56 ^a , 3.70 ^a , 3.78 ^a , 3.88 ^a , 3.99 ^a , 4.09 ^a	2	(Le Mao et al., 2021)
CL28	lactic acid	4.11 (1H) ^a	1.32 (3H) ^a	2	(Le Mao et al., 2021; Liang et al., 2015)
CL29	pyroglutamic acid	4.17 (1H) ^a	2.02 ^a , 2.38 ^a , 2.41 ^a , 2.50 ^a	2	(Kostidis et al., 2017; Liang et al., 2015)
CL30	trigonelline	4.43 (3H, s)	8.07 ^a , 8.82 ^a , 8.83 ^a , 9.11 ^a	2	(Le Mao et al., 2021; Ritota et al., 2012; Sabino et al., 2019)
CL31	xylose	4.57 (1H, d, J = 3.75)	3.22 ^a , 3.31 ^a , 3.43 ^a , 3.52 (1H) ^a , 3.60 ^a , 3.62 ^a , 3.66 ^a , 3.67 ^a , 3.69 ^a , 3.92 ^a , 5.19 (1H, d, J = 3.66)	2	(Le Mao et al., 2021; Moing et al., 2004)
CL32	α-glucose	5.23 (1H, d, J = 3.75)	3.23 ^a , 3.39 ^a , 3.45 ^a , 3.48 ^a , 3.52 ^a , 3.70 ^a , 3.72 ^a , 3.82 ^a , 3.89 ^a , 4.64 ^a	2	(Bourafai-Aziez et al., 2022; Le Mao et al., 2021)
CL33	sucrose	5.40 (1H, d, J = 3.85)	3.47 ^a , 3.55 ^a , 3.67 ^a , 3.76 ^a , 3.80 ^a , 3.82 ^a , 3.83 ^a , 3.88 (1H) ^a , 4.04 ^a , 4.21 ^a	2	(Sabino et al., 2019)
CL34	fumaric acid	6.51 (2H, s)		2	(Pacholczyk-Sienicka et al., 2024)
CL35	histidine	7.09 (1H, s)	3.14 ^a , 3.24 ^a , 3.98 ^a , 7.87 ^a	2	(Pacholczyk-Sienicka et al., 2024)
CL36	phenylalanine	7.42 (2H, m)	3.12 ^a , 3.28 ^a , 3.99 ^a , 7.32 ^a , 7.37 ^a	2	(Kostidis et al., 2017)

^a Overlapped signals

^b MSI level of identification according to Sumner *et al.* (Sumner et al., 2007).

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