



Article New Hybrid Tomato Cultivars: An NMR-Based Chemical Characterization

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Featured Application: The knowledge of the hybrid tomatoes metabolite profile represents an important tool for fresh market, industries, and customers, driving them to a more targeted and valorized use of the fruits.

Abstract: Bamano, King Creole, Sugarland, and DulceMiel hybrid tomato cultivars have been recently introduced in the Lazio area (Central Italy) to expand and valorize the regional/national market. Tomatoes from these cultivars, together with tomatoes from the native Fiaschetta cultivar, were sampled at the proper ripening time for the fresh market and characterized to obtain and compare their metabolite profiles. The Bligh–Dyer extraction protocol was carried out, and the resulting organic and hydroalcoholic fractions were analyzed by high-field Nuclear Magnetic Resonance (NMR) spectroscopy. NMR data relative to quantified metabolites (sugars, amino acids, organic acids, sterols, and fatty acids) allowed to point out similarities and differences among cultivars. DulceMiel hybrid and Fiaschetta native cultivars showed some common aspects having the highest levels of the most abundant amino acids as well as comparable amounts of organic acids, amino acids, stigmasterol, and linoleic and linolenic acids. However, DulceMiel turned out to have higher levels of glucose, fructose, and galactose with respect to Fiaschetta, reflecting the particular taste of the DulceMiel product. King Creole, Bamano, and Sugarland hybrid cultivars were generally characterized by the lowest content of amino acids and organic acids. King Creole showed the highest content of malic acid, whereas Bamano was characterized by the highest levels of glucose and fructose.

Keywords: hybrid tomato; ¹H-NMR; metabolic profile

1. Introduction

Tomatoes (*Lycopersicum esculentum*) represent the most consumed vegetables worldwide grown for both fresh consumption and processing (mainly to make tomato paste), thus identifying two different industry segments. According to statistics [1], 182 million tons of tomatoes were globally grown in 2017, 25 million tons being produced in Europe. Forty percent of the European tomatoes are sold on the fresh market. Today, Italy represents the first major European producer (followed by Spain), capable of providing 6 million tons of product per year.

In the last two decades, breeding programs and hybrid seed production have been addressed to enhance plant tolerance to abiotic and biotic stresses, fruit nutritional value, and fruit quality [2].

Duvick [3] has reported that in the USA, as well as in most European, Asian, and Australian countries, the majority of tomato production (100% of fresh market and 80% of processing tomatoes) is obtained with hybrids [2]. Hybrid tomato cultivars are more frequently used than the open pollinated lines thanks to advantages such as high productivity, early maturation, high fruit uniformity, high fruit quality, and disease resistance. Tomato hybrid production often gives plants with desirable characteristics superior with respect to the native ones but, at the same time, the production of hybrid seeds is generally not easy and laborious [2].

Tomato features such as taste and fragrance as well as healthful properties depend on the fruit chemical composition. For instance, sugars are responsible for different sweetness levels; organic acids are correlated with sourness; amino acids such as glutamate, glutamine, and aspartate enhance the bitterness [4,5]; whereas glutamic acid seems to be correlated to umami taste [6].

Among different analytical methods, high-resolution Nuclear Magnetic Resonance (NMR) spectroscopy is one of the most suitable approaches to monitor the composition of foodstuffs, owing to its ability to identify and quantify major and minor components without physical separation in the same experiment [7,8]. The NMR metabolomic approach has been largely used to characterize tomatoes fruits and derived products. Besides the identification of principal components such as sugars, organic acids, amino acids, and minor components [9–12], several studies have been carried out to determine carotenoid composition [13], degree of ripeness [14–17], geographical origin [18], and sensory features [4,5,19]; to compare organic and conventional tomatoes [20]; and to evaluate genetic transformations [21,22].

Within the regional project titled "eALIERB: an OPEN LAB to characterize and valorize foodstuffs and botanicals of Lazio region", the chemical composition of hybrid tomatoes namely Bamano, DulceMiel, Sugarland, and King Creole cultivars together with the native Italian Fiaschetta di Fondi cultivar was investigated for the first time. These hybrid cultivars have been recently introduced in the Lazio region (Central Italy) aiming at expanding the regional agronomic fresh market with products characterized by peculiar sensorial and nutritional features useful for industries and customers. In this paper, the NMR-based characterization of Bamano, DulceMiel, Sugarland, and King Creole hybrid cultivars together with the native Italian Fiaschetta di Fondi cultivar was carried out to obtain and compare their metabolite profiles.

2. Materials and Methods

2.1. Sampling

Tomatoes from four hybrid cultivars (Figure 1) namely Bamano, DulceMiel, Sugarland RZ F1, and King Creole and one native Fiaschetta di Fondi cultivar were cultivated in the southern area of the Lazio region (Fondi, LT) and collected by the "Terra del Sole" local farm. Tomatoes were cultivated within a greenhouse, without climate control systems, and were collected by hand during July according to their ripening degree proper for the market. Tomato seeds were provided by Syngenta, Basel, Switzerland (Bamano, DulceMiel and King Creole), Rijk Zwaan, De Lier, Netherlands (Sugarland RZ F1), and Terra del Sole, Fondi, Latina, Italy (Fiaschetta di Fondi) farms.



Figure 1. Tomato cultivars from Fondi area (Lazio region): (**A**) Fiaschetta di Fondi; (**B**) Bamano; (**C**) King Creole; (**D**) DulceMiel; (**E**) Sugarland.

2.2. Tomatoe Cultivars

Bamano tomatoes (Figure 1B) are uniform orange fruits (weight 10–12 g) with a consistent shape. King Creole fruits (Figure 1C, weight 30–40 g) are green in color, turning to dark red-brown when mature, and round in shape. Sugarland RZ F1 (Figure 1E) represents a cherry cultivar with small fruits, deep shiny red in color. DulceMiel (Figure 1D) is a sweet and aromatic cultivar characterized by green color, with honey shades when it is mature, oval in shape (weight 14–16 g). Fiaschetta di Fondi (Figure 1A) is a native tomato officially recognized in 2017 as a traditional agricultural product ("TAP") by the Italian Ministry of Agriculture. Fiaschetta belongs to *Lycopersicon esculentum* (L.) species, local ecotype "Fiaschetta di Fondi". It is characterized by a flask shape, deep and uniform red color (weight 40 g). The peel is consistent, and the pulp has a unique sweet-sour taste with a Brix grade of 5.

2.3. Extraction Procedure

Fifteen fresh tomatoes for each cultivar were frozen and ground in liquid nitrogen to obtain a homogeneous pool subjected to the Bligh–Dyer procedure. In detail, sequentially, 3 mL of methanol/chloroform (2:1 v/v) mixture, 1 mL of chloroform, and 1.2 mL of distilled water was added to about 1.0 g of homogenized tomatoes from each cultivar. After each addition, the sample was carefully shaken. The emulsion was preserved at 4 °C for 40 min. The sample was then centrifuged (4200 g for 15 min at 4 °C), and the hydroalcoholic and organic phases were carefully separated. The pellets were re-extracted using half of the solvent volumes (in the same conditions described above), and the separated fractions were pooled. Both fractions were dried under a gentle N₂ flow at room temperature until the solvent was completely evaporated. The dried phases were stored at -20 °C until further analyses [23].

2.4. NMR Analyses

The dried organic fraction of each sample was dissolved in 0.7 mL of a CDCl₃/CD₃OD mixture (2:1 v/v) and then placed into a 5 mm NMR tube. Finally, the NMR tube was flame sealed. Conversely, the dried hydroalcoholic phase of each sample was solubilized in 0.7 mL 400 mM phosphate buffer/D₂O containing 1 mM solution of TSP as internal standard and then transferred into a 5 mm NMR tube. NMR spectra of all hydroalcoholic and organic extracts were recorded at 27 °C on a Bruker AVANCE 600 spectrometer operating at the proton frequency of 600.13 MHz and equipped with a Bruker multinuclear z-gradient 5 mm probe head. ¹H spectra were referenced to methyl group signals of TSP $(\delta = 0.00 \text{ ppm})$ in D₂O and to the residual CHD₂ signal of methanol (set to 3.31 ppm) in CD₃OD/CDCl₃ mixture. ¹H spectra of hydroalcoholic extracts were acquired with 256 transients with a recycle delay of 5 s. The residual HDO signal was suppressed using a pre-saturation. The experiment was carried out by using 45° pulse of 6.5–7.5 μ s, 32 K data points. ¹H spectra of extracts in CD₃OD/CDCl₃ were acquired with 256 transients, recycle delay of 5 s, and 90° pulse of 9–11 µs, 32 K data points. The two-dimensional (2D) NMR experiments, such as ¹H-¹H TOCSY, ¹H-¹³C HSQC, and ¹H-¹³C HMBC, were carried out under the same experimental conditions previously reported [24]. The integrals of 25 selected signals in hydroalcoholic extract (Table 1) were measured using the Bruker TOPSPIN software and normalized with respect to the resonance at 0.00 ppm, due to methyl group signal of TSP, normalized to 100. Results have been expressed in mg/100 g FW. The integrals of 6 selected signals in organic extract (Table 2) were also measured using the Bruker TOPSPIN software and normalized with respect to the resonance at 2.30 ppm, due to signal of total fatty acids, set to 100. The molar % values of fatty acids and sterols were calculated using the following equations:

$$\%_{\beta-\text{SIT}} = 100 \times 0.66 I_{\beta-\text{SIT}} / I_{\text{tot}}$$
⁽¹⁾

$$\%_{\rm STIG} = 100 \times 0.66 I_{\rm STIG} / I_{\rm tot}$$
⁽²⁾

$$%_{\text{TRI}} = 100 \times 0.5 I_{\text{TRI}} / I_{\text{tot}}$$
 (3)

$$\%_{\rm DI} = 100 \times I_{\rm DI}/I_{\rm tot} \tag{4}$$

$$M_{MONO} = 100 \times (I_{UNS} - 2I_{Di} - 1.5I_{TRI})/I_{tot}$$
 (5)

$$%_{SAT} = 100 \times (I_{FA} - I_{DI} - 0.5I_{TRI} - I_{MONO})/I_{tot}$$
 (6)

where $%_{\beta-SIT}$, $%_{STIG}$, $%_{TRI}$, $%_{DI}$, $%_{MONO}$, $%_{SAT}$ are molar % of β -sitosterol, stigmasterol, tri-unsaturated fatty acids, di-unsaturated fatty acids, mono-unsaturated fatty acids, and saturated fatty acids, respectively. I_{β -SIT}, I_{STIG}, I_{TRI}, I_{DI}, I_{UNS}, I_{FA} are integrals (see Table 2), and I_{tot} is calculated according to the Equation (7)

$$I_{tot} = I_{FA} + 0.66I_{\beta-SIT} + 0.66I_{STIG}$$
(7)

Table 1. Compounds and relative signals (ppm) selected for quantitative analysis in the hydroalcoholic extracts.

ppm	Compounds	ppm	Compounds
0.96	Leu	3.21	Choline
0.99	Val	3.25	β-Glucose
1.01	Ile	4.04	Fructose
1.33	Lactic acid	4.31	Malic Acid
1.34	Thr	4.59	β-Galactose
1.49	Ala	5.25	α-Glucose
2.30	GABA	6.91	Tyr
2.35	Glu	7.34	Phe
2.46	Gln	7.74	Trp
2.55	Citric Acid	8.46	Formic Acid
2.81	Asp	9.13	Trigonelline
2.90	Asn		-

	ppm	Group	Compounds
$I_{\beta-SIT}$	0.66	CH ₃	β-Sitosterol
I _{STIG}	0.68	CH_3	Stigmasterol
I _{FA}	2.30	α -CH ₂	Total fatty acids
I _{DI}	2.73	CH ₂	Di-unsaturated fatty acids
I _{TRI}	2.77	CH ₂	Tri-unsaturated fatty acids
I _{UNS}	5.31	CH=CH	Total unsaturated fatty acids

Table 2. Compounds and relative signals (ppm) selected for quantitative analysis in the organic extracts.

2.5. Tree Clustering Analysis

Statistical analysis of the NMR data was carried out using STATISTICA software (version 5.1). Before performing the Tree Clustering Analysis, all the selected variables were mean-centered, and each variable was divided by its standard deviation (autoscaling). For Tree Clustering Analysis [25] the Euclidean distance, Equation (8), was used as a measure of distance between samples. For the measure of distance between clusters the complete linkage method was used (when the distance between clusters is determined using the greatest distance between any two objects in the different clusters):

distance
$$(x, y) = [\Sigma_i (x_i - y_i)^2]^{1/2}$$
 (8)

3. Results

The results of the NMR analysis regarding the tomato hydroalcoholic and organic fractions will be reported separately.

3.1. Tomato Hydroalcoholic Fraction

The 1D NMR spectrum assignment of the hydroalcoholic extracts solubilized in D₂O phosphate buffer (data not reported) was obtained using literature data [11,16,17] and confirmed by 2D experiments (Figures S1–S3). Compounds and relative NMR signals selected for quantitative analysis are reported in Table 1.

The amino acid profiles (Figure 2A) of the five cultivars showed some common aspects: GABA, alanine, glutamine, and glutamic acid were always the most abundant amino acids [6,26–28]. In particular, glutamic acid was the most abundant amino acid in all cultivars. On the other hand, valine, isoleucine, tyrosine, tryptophan, and leucine were always present at lower levels with respect to other amino acids. Concerning the difference among the cultivars, glutamic acid showed significant variations reaching 447 mg/100 g in DulceMiel cultivar, at least double with respect to the other cultivars. DulceMiel cultivar was also the cultivar with the major content of leucine, aspartic acid, and alanine. Fiaschetta cultivar showed a higher level of isoleucine, tyrosine, asparagine, GABA, and glutamine with respect to the others. Valine, tryptophan, threonine, and phenylalanine were found to be more abundant in Sugarland. Bamano and King Creole cultivars were characterized by a lower content of amino acids compared to the other samples.

Citric, malic, lactic, and formic acids were detected and quantified in all five cultivars with citric acid as the most abundant organic acid [17,29,30] (Figure 2B). DulceMiel cultivar showed the highest content of citric acid (460 mg/100 g) whereas King Creole showed an almost double malic acid content with respect to the other tomato cultivars. Formic acid was always present in minor amount being lower than 1.1 mg/100 g. Fiaschetta cultivar was characterized by high lactic acid level (26.5 mg/100 g), at least 3.5-fold higher than in the other cultivars and more than doubled with respect to malic acid content (10.4 mg/100 g), which is known to be, together with citric acid, one of the most abundant organic acid in tomatoes [11,20,21]. Note that lactic acid has been rarely detected and not always quantified in tomato fruits [14,30]. The presence of higher amount of lactic acid in Fiachetta cultivar could be ascribed to the fermentation of tomato pulp by the bacteriocin-producing lactic acid bacteria [31,32].

Fructose was the most abundant sugar in all tomato fruits [5,10,12,17,21], followed by glucose and galactose. DulceMiel, Bamano, and Sugarland cultivars showed the highest levels of fructose,

galactose, and glucose, whereas both King Creole and Fiaschetta cultivars were characterized by the lowest levels, nearly halved with respect to the other cultivars (Figure 2C). These data suggest that Fiaschetta and King Creole cultivars could be useful in hypoglycemic diet.

Regarding the content of other metabolites (Figure 2D), the concentration of choline ranged from 6.9 mg/100 g in King Creole to 15.4 mg/100 g in DulceMiel cultivar, whereas trigonelline content was found to be lower than 2.5 mg/100 g, varying from 0.9 mg/100 in King Creole to 2.4 mg/100 g in Fiaschetta.



Figure 2. Histograms comparing the level of metabolites present in tomato hydroalcoholic extracts from the five investigated cultivars: (**A**) Amino acids; (**B**) Organic acids; (**C**) Carbohydrates; (**D**) Miscellaneous (B: Bamano, KC: King Creole, S: Sugarland, DM: DulceMiel, F: Fiaschetta). Data shown are expressed as mg/100 g FW.

The ¹H and ¹³C NMR assignments of Bamano, King Creole, DulceMiel, Sugarland, and Fiaschetta tomatoes extracts in CDCl₃/CD₃OH was obtained by 2D experiments (Figures S4–S6). In Table 3, the metabolites identified in the tomatoes organic fraction are reported along with the assignment of ¹H and corresponding ¹³C resonances. Previously, the ¹H NMR spectra assignment of organic fraction from lyophilized cherry tomatoes (Naomi and Shiren cultivars) in CDCl₃ has been reported [9,14]. Taking into account different plant material, a different extraction procedure, and a different solvent (CDCl₃/CD₃OH instead of CDCl₃) used in the present study, a new ¹H NMR spectra assignment was required.

In the upfield spectral region, the CH₃-18 signals of β -sitosterol at 0.66 ppm (I_{β -SIT}), and stigmasterol at 0.68 ppm (I_{STIG}) were detected and quantified (Table 3 and Figures 3 and 4). The resonance centered at 2.30 ppm (I_{FA}) was ascribed to α -CH₂ of all fatty acid chains. Resonances at 2.73 (I_{DI}) and 2.77 ppm (I_{TRI}) were ascribed to diallylic CH₂ of linoleic and linolenic acids. The multiplet at 5.31 ppm (I_{UNS}) belongs to double bond CH groups of all unsaturated fatty acids.



Figure 3. ¹H NMR spectrum of organic extract from tomato fruits of "Fiaschetta di Fondi" in CDCl₃/CD₃OD 2:1 *v/v* mixture. Signals used for the quantitative analysis have been labelled: $I_{\beta-SIT}$, CH₃, β -sitosterol; I_{STIG} , CH₃, stigmasterol; I_{FA} , α -CH₂, all fatty acids; I_{DI} , diallylic CH₂, linoleic acid; I_{TRI} , diallylic CH₂, linolenic acid; I_{UNS} , CH=CH of all unsaturated fatty acids.

Table 3. Metabolites identified and quantified in the 600.13 MHz ¹H spectra (27 °C) of organic extracts from Bamano, DuceMiel, King Creole, Sugarland, and Fiaschetta tomato cultivars.

Compound	Assignment ^a	¹ H (ppm)	Multiplicity: J [Hz]	¹³ C (ppm)
Oleic fatty chain	COO			174.4
$(C18:1 \Delta^9)$	CH ₂ -2	2.30		34.6
, , , , , , , , , , , , , , , , , , ,	CH ₂ -3	1.58	m	25.3
	CH ₂ -4,7	1.30	m	29.5
	CH2-8	2.01	m	27.6
	CH=CH 9,10	5.31	m	130.4
	CH ₂ -11	2.01	m	27.6
	CH ₂ -12,15	1.33-1.30	m	29.4-30.2
	CH ₂ -16	1.28	m	31.9
	CH ₂ -17	1.26	m	22.9
	CH ₃ -18	0.84	t	14.3

Compound	Assignment ^a	¹ H (ppm)	Multiplicity: J [Hz]	¹³ C (ppm)
Di-unsaturated fatty				
acids (DUFA)	COO			174.4
Linoleic fatty chain		• • •		24.4
$(C18:2 \Delta^{9,12})$	CH ₂ -2	2.30		34.6
	CH ₂ -3	1.58	m	25.3
	CH ₂ -4,7	1.32-1.28	m	29.5
	CH ₂ -8	2.02	m	27.6
	CH-9	5.33	m	130.4
	CH-10	5.30	m	128.6
	CH ₂ -11	2.73	t [6.8]	26.0
	CH-12	5.30	m	128.6
	CH-13	5.33	m	130.4
	CH ₂ -14	2.02	m	27.6
	CH ₂ -15	1.27	m	29.4
	CH ₂ -16	1.27	m	31.6
	CH ₂ -17	1.23	m	22.9
	CH ₃ -18	0.85	t	14.3
Tri-unsaturated fatty				
acids (TUFA)	COO			174.4
Linolenic fatty chain				
$(C18:3 \Delta^{9,12,15})$	CH ₂ -2	2.30		34.6
	CH ₂ -3	1.58	m	25.3
	CH ₂ -4,7	1.30	m	29.5
	CH2-8	2.03	m	27.6
	CH-9	5.33	m	130.4
	CH-10	5.30	m	128.6
	CH ₂ 11	2.77	t [6.2]	26.0
	CH=CH 12,13	5.30	m	128.6
	CH ₂ -14	2.77	t [6.2]	26.0
	CH-15	5.27	m	127.4
	CH-16	5.35	m	132.2
	CH ₂ -17	2.04	m	20.9
	CH3-18	0.94	t [7.6]	14.4
Diacylglycerol moiety	CH ₂ - sn 1	4.34;4.16		62.6
	CH- sn 2	5.20	m	70.2
	CH ₂ - sn 3	4.07		64.7
Saturated fatty acids (SFA)	COO			174.4
(011)	CH ₂ -2	2.28		34.6
	CH2-3	1.58	m	25.3
	CH ₂	1.28-1.22	m	29.6-32.0
	CH ₂ n-1	1.26		22.9
	CH ₃ n	0.84	t	14.2
β-Sitosterol	CH ₃ -18	0.66	S	12.2
Stigmasterol	CH ₃ -18	0.68	S	12.2
Sugnasteror	CH=CH-22.23	5.12: 4.99		
	CH(OH)-3	3.50		71.7
Squalene	CH ₃ -a	1.56		16.2
	CH ₃ -b	1.64		25.8
	CH-c	5.07	m	124.7
	CH2-d	2.03	5.0.02	27.4
	CH ₂ -e	1.94		40.1
Phosphatidylcholine	N(CH ₃) ₃	3.21	S	54.5

Table 3. Cont.

^a For structure fragment numbering see [33].

The results of quantitative analysis, reported as histograms in Figure 4, pointed out some interesting differences between the investigated tomatoes cultivars. The amount of β -sitosterol changed from 3.1% (Fiaschetta) to 5.9% (Bamano), whereas stigmasterol ranged from 0.7% in King Creole to 1.8% in Bamano cultivar. The total concentration of unsaturated fatty acids (UFA) is very close in all the selected tomato fruits; it varies from 51.7% in Bamano to 65.8% in Fiaschetta. In terms of mono-unsaturated fatty acids (MUFA), Fiaschetta cultivar was found to be the richest one (24.7%), whereas Sugarland

and Bamano (15.9 and 16.5%, respectively) were characterized by the lowest contents. Di-unsaturated fatty acids (DUFA) level (linoleic acid) was comparable in the five cultivars ranging from 24.7% in King Creole to 30.7% in DulceMiel, whereas tri-unsaturated fatty acid content (TUFA, linolenic acid) was found to be nearly doubled in King Creole (15.96%) compared to Bamano (8.9%) which was characterized by the lowest content.



Figure 4. Histograms comparing the concentration (% molar) of metabolites present in tomato organic extracts from the investigated five cultivars (B: Bamano, KC: King Creole, S: Sugarland, DM: DulceMiel, F: Fiaschetta).

3.3. Hydroalcoholic and Organic Fractions: A Comparison Among Cultivars

NMR data including all the quantified metabolites from both hydroalcoholic and organic fractions were submitted to tree cluster analysis (TCA) to point out general similarities/differences in the metabolite profile of the investigated cultivars (Figure 5). The first level of the dendrogram showed two clusters: the first cluster is composed by Fiaschetta (native) and DulceMiel (hybrid) cultivars, and the second one includes the three remaining hybrid cultivars. Cutting the dendrogram at a lower level the previous second cluster is further divided in two groups: one consisting of King Creole cultivar and the other one consisting of the remaining two hybrid cultivars, Bamano and Sugarland.

DulceMiel and Fiaschetta are clearly distinguished from the rest, having, as shown in the histograms of Figure 2, the highest levels of the most abundant amino acids namely glutamic acid, glutamine, alanine, GABA, and aspartic acid as well as citric and lactic acids. Moreover, both the cultivars have comparable amounts of citric and malic acids, aspartic acid, phenylalanine, threonine, valine, choline, stigmasterol, DUFA, and TUFA. A significant difference between these two cultivars is the carbohydrate profile: higher levels of glucose, fructose, and galactose were found in DulceMiel than in Fiaschetta, the latter being characterized by the lowest content of sugars among all cultivars.

King Creole, Bamano, and Sugarland belong to the second TCA cluster. They were characterized by the lowest content of many hydroalcoholic metabolites. However, it is important to note that King Creole showed the highest content of malic acid. Bamano and Sugarland showed a very similar chemical composition, being characterized by high levels of glucose, fructose, and galactose. In fact, the producers declare these two cultivars to be the sweetest compared to the others, likely due to a balance between acids and sugars in favor of the last ones. Bamano and Sugarland were similar also in terms of organic extracts composition: higher levels of sterols (β -sitosterol and stigmasterol) and SFA, and lower levels of UFA (in particular MUFA) were found in both tomato fruits.



Figure 5. Dendrogram (TCA) showing the clustering, based on NMR data (both hydroalcoholic and organic metabolites quantified) of F: Fiaschetta, B: Bamano, DM: DulceMiel, S: Sugarland, KC: King Creole. Two levels corresponding to a different number of groups are marked.

4. Conclusions

The NMR-based characterization allowed to obtain the metabolite profiling of hybrid tomato cultivars essential to define nutritional properties. This untargeted approach allowed to characterize each cultivar: King Creole showed the highest content of malic acid, whereas Bamano and Sugarland were characterized by high levels of glucose and fructose. Fiaschetta native cultivar showed the highest level of lactic acid and MUFA and the lowest levels of sugars, whereas DulceMiel cultivar was characterized by high levels of both sugars and acids. The content of water-soluble metabolites in analyzed tomatoes is generally consistent with the literature data obtained by NMR-based characterization of other tomato cultivars ("Ailsa Craig" [17], "Palmiro" and "Clotilde" [10], "Moneymaker" [12]). Nevertheless, a few metabolites showed either relatively lower (malic acid) or higher content (alanine, threonine) with respect to the literature data mentioned above.

The knowledge of the nutritional profile and chemical composition of hybrid tomatoes may represent an important tool for fresh market, industries and customers, driving them to a more targeted and optimized use of the fruits.

Supplementary Materials: The following are available online at http://www.mdpi.com/2076-3417/10/5/1887/s1, Figure S1: ¹H-¹H TOCSY spectrum of hydroalcoholic extract from "Fiaschetta di Fondi" tomato fruits in 400 mM phosphate buffer/D₂O 1 mM TSP, Figure S2: ¹H-¹³C HSQC spectrum of hydroalcoholic extract from "Fiaschetta di Fondi" tomato fruits in 400 mM phosphate buffer/D₂O 1 mM TSP, Figure S3: ¹H-¹³C HMBC spectrum of hydroalcoholic extract from "Fiaschetta di Fondi" tomato fruits in 400 mM phosphate buffer/D₂O 1 mM TSP, Figure S4: ¹H-¹H TOCSY spectrum of organic extract from "Fiaschetta di Fondi" tomato fruits in CDCl₃/CD₃OD 2:1 v/v mixture, Figure S5: ¹H-¹³C HSQC spectrum of organic extract from "Fiaschetta di Fondi" tomato fruits in CDCl₃/CD₃OD 2:1 v/v mixture, Figure S6: ¹H-¹³C HMBC spectrum of organic extract from "Fiaschetta di Fondi" tomato fruits in CDCl₃/CD₃OD 2:1 v/v mixture.

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