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Nutrient, fibre, sorbitol and chlorogenic acid content of prunes 1 (Prunus domestica): an updated analysis and comparison of 2 different countries of origin and database values 3 4 5 Samantha K Gill*, Ellen Lever*, Peter W Emery, Kevin Whelan King's College London, Department of Nutritional Sciences, London, United Kingdom 6 7 Corresponding author: Professor Kevin Whelan, King's College London, Department of 8 Nutritional Sciences, 150 Stamford Street, London, SE1 9NH, United Kingdom, + 44 20 78 48 9 38 58, kevin.whelan@kcl.ac.uk 10 11 12 Short running head: Prune composition 13 Key words: dried fruit, dried plums, prunes, gastrointestinal health, fibre, polyphenols 14 *Joint first authors 15 16 Disclosure of interest: This study was funded by the California Dried Plum Board. The funders 17 played no role in study conception, design, analysis, interpretation or writing of the 18 manuscript. KW has served as a consultant for Danone, has received speaker fees from Alpro 19 20 and the National Dried Fruit Trade Association UK Ltd and has received research funding from Almond Board of California, California Dried Plum Board, Clasado Biosciences, Nestec Ltd, 21 International Nut and Dried Fruit Council, and receives royalties from FoodMaestro. The 22

23 remaining authors report no conflicts of interest.

24 Abstract

Current prune composition data is outdated and requires a comprehensive and comparative 25 26 re-analysis. This novel study aimed to: (i) analyse and compare prune composition from major 27 countries of origin; and (ii) provide a comprehensive compositional analysis of prunes of USA origin and compare this with UK and USA database data. Prune samples were analysed for 28 major nutrients and bioactive compounds and compared between countries of origin. Total 29 fibre was higher in prunes from the USA (12.0 g/100g) and Chile (11.5 g/100g) compared with 30 France (8.4 g/100g) and Argentina (8.9 g/100g), while prunes from all countries contained 31 high levels of sorbitol (11.2-15.5 g/100g). Differences of energy and starch values compared 32 33 with national databases reflected different approaches to sampling and analysis. In 34 conclusion, prunes contain high levels of fibre and other bioactive compounds. Variations 35 between country of origin and database values highlight the importance of transparency in 36 documenting sampling and analysis methods.

37 Introduction

Studies have highlighted the potential benefits of dried fruits on a variety of health outcomes 38 39 (Chang et al. 2016). In particular their high fibre content has led to investigation of the role of 40 dried fruit in the maintenance and promotion of gastrointestinal health (Lever et al. 2015) 41 which is considered of major public health importance (DuBois 2004; Wald et al. 2007). The impact of dietary fibre on health is affected by variations in its chemical composition (e.g. 42 distribution of different fibre fractions) and physical structure (e.g. degree of polymerisation, 43 molecular weight and linkages) that alter its solubility, viscosity and fermentability. Given that 44 45 dried fruits are nutritionally comparable to whole fresh fruits, only provided in a smaller and 46 more concentrated form, they may be a convenient and versatile option for increasing fruit 47 consumption across population groups (Sadler et al. 2019).

48

Plums are taxonomically diverse stone fruits of Prunus domestica L. and are commonly 49 consumed in their dried form, termed prunes. Data from various sources, including the United 50 Kingdom (Finglas 2015) and the United States of America (USA) (US Department of Agriculture 51 2018), indicate that prunes are naturally high in a variety of poorly-fermented and readily-52 53 fermented dietary fibres (>6 g/100g including hemicellulose, pectin, cellulose). In addition, 54 prunes contain other bioactive compounds such as polyphenols, which may stimulate colonic proliferation of microorganisms such as Bifidobacteria and Lactobacilli. Furthermore, prunes 55 contain high amounts of sorbitol (~12 g/100g) which is known to have laxative effects (Yao et 56 al. 2014). Indeed, a systematic review concluded that prunes may play a role in 57 58 gastrointestinal health by increasing stool frequency and improving stool consistency (Lever et al. 2014). 59

60

Data on prune composition require updating for several reasons. Firstly, existing databases, such as McCance and Widdowson (UK) and USDA (USA) databases, do not report a wide range of components relevant to gut health (e.g. different fibre fractions and sorbitol content), and secondly the current data were compiled between 1980 and 2001 and therefore may no longer accurately reflect present-day prune composition. Thirdly, the USDA data calculates total carbohydrate 'by difference', which does not account for the lower energy contributions from unavailable carbohydrates. Fourthly, the composition of prunes may vary depending on 68 a variety of factors including growing and harvesting conditions and post-harvest processes (e.g. drying, dehydration and rehydration, storage conditions). Given that the vast majority of 69 70 global supply of prunes originates from four countries: the USA (largely California, 43%), Chile 71 (24%), France (16%) and Argentina (15%) (Buncher 2012), currently-available prune 72 composition data may be confounded by variations in origin. For example, standard yellow 73 plums have been shown to contain higher vitamin and phenolic compound content than 74 organically grown plums (Lombardi-Boccia et al. 2004), while prunes from Australia have been 75 shown to contain higher iron and folate contents than prunes from USA and Chile (Bennett et 76 al. 2011). Finally, variations in the nutrient composition of prunes of different countries of 77 origin will impact the database values in each country. For example, databases in the USA 78 (USDA) and France (CIQUAL), both of whom are large producers of prunes, will reflect the 79 composition of prunes from those countries, whereas the database in the UK (McCance and 80 Widdowson), which does not grow large supplies of prunes, will reflect the composite of 81 prunes from different countries of origin.

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With this in mind, we aimed to investigate the energy, macronutrient, micronutrient, fibre, sorbitol and polyphenol composition of prunes by: (i) analysing and comparing the composition of prunes from major countries of origin (USA, Chile, France and Argentina); and (ii) undertaking an in-depth analysis of prunes of USA (Californian) origin and comparing this with data from food composition databases.

88

89 Materials and Methods

90 Sample collection

Prune samples grown by the four largest producers of prunes were collected in order that 91 92 composition could be both globally representative and compared between country of origin (USA, Chile, France and Argentina). Prune samples were purchased from major population 93 94 centres in five countries across Europe (France, Germany, Italy, Spain, United Kingdom) as 95 these are five major European markets for prunes and thus data would reflect the 96 composition of prunes available across Europe, as well as meeting the Food Information to 97 Consumers Legislation (European Commission 2011). Prune samples were purchased as sold 98 to the customer from major retail outlets including supermarkets, department stores and health food stores and including a range of brands (where available) to ensure purchase of 99

prunes representative of those most frequently consumed, i.e. with the highest volume of sales. Prune samples were purchased at the same time of year and within use-by-dates. Samples were required to be in unopened packets of ≥ 100 g with a remaining shelf life of ≥ 6 months. Prunes were stored unopened until analysis to minimise drying, water absorption and contamination. Prunes were purchased pitted (stone removed) or whole. If purchased whole, stones were removed prior to compositional analysis.

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In total, the goal was to purchase 10 to 12 samples from each of the four countries of origin,
with at least 3 samples from each sampling country. This number is recommended for nutrient
composition database data and based upon guidance from Greenfield and Southgate (2003),
though this depends on the variability of the nutrients being measured.

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Prune samples from each country of origin were pooled prior to analysis. Funding restrictions meant that the study could either: (i) individually analyse a number of prune samples from a single country, thus allowing measurement of within-country variation but not betweencountry variation; or (ii) analyse a pooled sample from a number of sampling countries, thus allowing measurement of between-country variation albeit without statistical comparison.

Given the wide geographic difference in countries of origin (USA, France, Chile, Argentina), it was felt that between-country variations, rather than within-country variations, were likely to be larger and therefore of greater nutritional relevance.

120

121 Sample preparation and analysis

Samples were pooled according to country of origin (**Table 1**). This pooled sample comprised an equal weight of 500 g (i.e. 100 g adjusted weight from each sampling country), of prunes from each of the five sampling countries. Pooled samples were homogenised using a hand mincer, divided into aliquots, stored frozen at -80°C and defrosted prior to analysis.

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Prune samples were analysed at Leatherhead Food Research, Surrey, UK. The pooled sample from each of the four countries of origin was analysed using standard methods for energy (calculated from macronutrient data), protein (total nitrogen), fat (Soxhlet), carbohydrate (calculated by difference), sugars and sorbitol (ion-exchange chromatography), a range of fibre classifications (AOAC methods, Englyst), and chlorogenic and neochlorogenic acid (ultraperformance liquid chromatography tandem mass spectrometry, UPLC with MS-MS) (**Table 2**). In addition, further in-depth analyses were performed on the pooled sample of prunes from the largest global producer (California, USA) including fatty acids (gas chromatography with flame ionisation detection), sugars (ion-exchange chromatography) and major micronutrients (inductively coupled plasma optical emission spectrometry ICP-OES, high performance liquid chromatography HPLC) (**Table 2**).

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In terms of the chromatographic methods, for sugar and sorbitol, extraction from the prune 139 140 samples was performed by sonication in hot water and treatment with Carrez reagents. The 141 filtered solution was then analysed using high-performance anion-exchange chromatography 142 coupled with pulsed electrochemical detection (HPAEC-PED) using a Dionex PA20 column 143 (Corradini et al, 2012). For chlorogenic and neochlorogenic acids, extraction from the prune 144 samples was performed in hot water and methanol and the solution analysed using UPLC with 145 MS-MS equipped with an ethylene bridged hybrid column (C18 2.1 x 50 mm, 1.7 μ m). For 146 fatty acids, transmethylation was undertaken to form methyl esters which were analysed using gas-liquid chromatography with a flame ionisation detection (Seppänen-Laakso, et al, 147 2002). For vitamin B analysis, extraction was performed using HCl and the solution analysed 148 149 using HPLC with fluorescence detection using a C18 conventional column (250 x 4.6 mm, 5 μm). 150

151

Duplicate analyses were carried out for analytes that were not routinely measured at the research centre. However, routine analyses were not performed in duplicate as these had criteria defining the limits of repeatability.

155

156 **Results**

157 Sample purchases

Sample purchases were made in France (Normandy), Germany (Bonn), Italy (Milan, Novara),
Spain (Madrid), and the UK (London) between March and June 2013. The pack sizes of the
purchased prune samples varied between 120 g and 1000 g. **Table 1** shows the number of
prune samples purchased and analysed from each sampling country and by country of origin.
One sample was excluded as it exceeded the use-by-date by the time of analysis and four

samples were excluded because the country of origin was unclear. Eighteen different samples were pooled and analysed for USA and French prunes, fifteen for Chilean prunes, but only five for Argentinian prunes (all purchased from Spain) due to their lack of availability in Europe at that time. The amount analysed from each individual sample was weighted so that an equal amount from each sampling country was included and pooled to make up a total of 500g from each country of origin (**Table 1**).

169

170 Composition Data

The composition of prunes from the pooled samples originating from USA, Chile, France and
Argentina are shown in **Table 2.** In general there were few major differences in nutrients and
fibre fractions between prunes of different countries of origin.

174

Differences in starch content were observed between countries, being lower in prunes of French origin (1.9 g/100 g) compared with others (5.7-6.6 g/100g). Total fibre (measured using AOAC 2011.25) was higher in prunes from the USA (12.0g/100g) and Chile (11.5g/100g) compared with those from France (8.4g/100g) and Argentina (8.9g/100g).

179

Prunes contained high levels of sorbitol (11.2-15.5g/100g) with broadly similar values across the countries of origin. High levels of the phenolic compounds chlorogenic acid and neochlorogenic acid were also found in prunes, however, in general these were higher in prunes from the USA (3.6 and 89.3 g/100g) and France (3.9 and 92.0 g/100g) compared with prunes from Chile (1.3 and 39.8 g/100g) and Argentina (1.3 and 40.3 g/100g) (**Table 2**).

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The composition of prunes from the USA pooled sample compared with data from the USDA
nutrient database and McCance and Widdowson's the composition of foods is shown in Table
3. Concentrations per wet weight are presented in order to be consistent with these
databases. Energy and starch values (230 kcals/100g and 6.6g/100g, respectively) were closer
to values published by the USDA database (240 kcals/100g and 5.1 g/100g, respectively), than
McCance and Widdowson (141 kcals/100g and 0.0 g/100g, respectively).

193 Discussion

The current study aimed to investigate the energy, macronutrient, micronutrient, fibre, sorbitol and polyphenol composition of prunes by: (i) analysing and comparing the composition of prunes from major countries of origin (USA, Chile, France and Argentina); and (ii) undertaking an in-depth analysis of prunes of USA (Californian) origin and comparing this with data from food composition databases.

199

In regards to the measured composition of prunes from different countries of origin, while the pooling of samples precluded statistical comparisons, at face value there were few differences in energy and macronutrient content between them, apart from starch which was lower in prunes of French origin (**Table 2**). Given that the same methods of analysis were used for prunes of all countries of origin and analysis occurred at the same time and in the same run, these differences likely reflect true compositional differences in prune samples between countries.

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There were no major differences in dietary fibre content when measured using the AOAC 208 209 991.43 method (which excludes low molecular weight fibres and most types of resistant starch) nor using the Englyst method (non-starch polysaccharides). However, when measured 210 211 using the AOAC 2011.25 method, fibre was higher in prunes from USA and Chile compared 212 with France and Argentina. The AOAC 2011.25 method includes all categories of dietary fibre, high and low molecular weight fibres and all types of resistant starch. Taken together, this 213 suggests that US and Chilean prunes likely contain greater low molecular weight fibres and 214 215 resistant starch than French and Argentinian prunes. Low molecular weight fibres are soluble, explaining the higher soluble fibre content in USA (7.6 g/100 g) and Chilean (6.3 g/100 g) 216 prunes compared with others (4.4-4.6 g/100 g). In addition, French prunes contained less 217 sorbitol, while Chilean and Argentinian prunes had lower chlorogenic and neochlorogenic acid 218 219 content than the prunes of other origins. As previously mentioned, these differences may be due to variations in soil management, plum ripeness or storage conditions (Donen 1939; Piga 220 221 et al. 2003), although the reasons for any differences was not investigated here.

In regards to prunes of USA (Californian) origin (Table 3), there were minor differences 223 between the current analytical data and those published by USDA, though these were small 224 and likely negligible from a nutritional perspective. Given that the UDSA database is updated 225 226 regularly through submission of independent analyses from food manufacturers, the minor 227 differences observed may simply reflect seasonal variation in composition. However, there 228 were larger differences in both the current analytical data and the USDA data compared with 229 the UK data provided by McCance and Widdowson, the latter reporting lower energy and starch values. Notably, little information is provided on the sourcing of prune samples 230 231 reported in McCance and Widdowson and so it is unknown whether prune origin could be 232 responsible for differences in starch content. Water content was comparable between the data (30.9-31.1%), therefore any differences were not due to variation in water content. 233 234 Rather, the difference in energy content is explained by differences in the components 235 included in the energy calculation and different conversion factors used.

236

237 In the current study, energy content is calculated based upon the contribution of 'available carbohydrate', fat, protein, fibre and polyols, as per European Union labelling regulations (EC, 238 239 2011). The USDA data includes 'total carbohydrate' in the energy calculation (rather than 240 'available carbohydrate'), and therefore does not take into account the lower energy 241 contribution from fibre and polyols, and this is reflected in the slightly higher energy value published by USDA (240 kcal/100 g) compared with the current analytical data (230 kcal/100 242 243 g). In stark contrast, the UK data from McCance and Widdowson (141 kcal/100 g) excludes fibre and polyols from the energy calculation. 244

245

Prunes contained high levels of sorbitol (11.2-15.5 g/100g), these values being similar to USDA 246 247 values (12.0 g/100g) and other studies in the scientific literature (10.8 g/100g) (Yao et al. 2014). The sorbitol content of prunes is therefore higher than that of its non-dried 248 249 counterpart plums (2.4 g/100g), as well as other non-dried stone fruits such as cherries (0.7 g/100g) and dried fruits such as dried apricots (6g/100g), dried pear (8.1 g/100g) and dried 250 apple (1.9g/100g) (Yao et al. 2014). Some polyols have been shown to induce increases in 251 small intestinal water, although this has not been confirmed for sorbitol. For example, a 252 253 fourfold increase in small intestinal water was observed in healthy individuals 60 minutes 254 following ingestion of 17.5 g of mannitol (Marciani et al. 2010).

Prunes also contained high levels of chlorogenic acid (1.3-3.9 g/100g) and neochlorogenic acid 256 (39.8-92.0 g/100g), particularly those from the USA and France. This reflects data from 257 258 previous studies reporting high levels of phenolic compounds in prunes (Donovan et al. 1998; 259 Stacewicz-Sapuntzakis 2013). These phenolic compounds are partially absorbed in the small 260 intestine and the remainder enter the colon where they undergo biotransformation by the 261 microbiota into caffeic acid and quinic acid (Olthof et al. 2001). A recent systematic review suggests that polyphenols and their degradation products can modulate the gut microbiota 262 263 and have prebiotic effects (Nash et al. 2018). Chlorogenic acid has been shown to inhibit the 264 growth and adhesion of selected gut pathogens to a human gut cell line and to enhance the 265 proliferation and adhesion of the probiotic L. rhamnosus (Parkar et al. 2008). Taken together, it is plausible that the combination of different dietary fibres, sorbitol and polyphenols 266 267 naturally abundant in prunes create a synergistic effect, which in part, may be the reason why 268 prunes are considered beneficial for gastrointestinal health (European Food Safety Authority 269 2014).

270

271 The current analytical data and the USDA data calculate carbohydrate values by difference 272 (subtracting amounts of the other proximates from the total weight), while McCance and Widdowson calculate available carbohydrate using monosaccharide equivalents of each 273 measured component. In the McCance and Widdowson UK data, available carbohydrate is 274 275 equal to total sugars since no polyols, oligosaccharides or starch are reported. Though the reason for the lack of starch in prunes (0.0 g/100 g) reported in the McCance and Widdowson 276 UK data is unclear, it is possible that when analysis was undertaken, prunes of French origin 277 (which in the current analysis contained less starch) were more readily available and 278 279 accessible. Notably, the current analytical data and the data from the USDA database may considerably overestimate the available carbohydrate content by including unmeasured 280 281 components which are not absorbed or not metabolised in the body to produce energy (e.g. sorbitol). In the present analytical data the sum of starch and sugars is 12 g/100g less than 282 the value for available carbohydrate by difference, while in the USDA data the sum of starch 283 and sugars is around 20 g/100g less than the value for total carbohydrate by difference that 284 285 has been used to calculate energy content. True energy values (kcals) for prunes appear to be 286 between 230-240 kcals/100g in accordance with the present analytical data (230 kcals/100g) and the USDA database (240 kcals/100g). This is in contrast to McCance and Widdowson that
presents noticeably lower energy values (141 kcals/100g).

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290 Limitations and strengths

291 The major limitation of this study was that, due to financial constraints, we did not analyse 292 multiple prune samples from each country of origin that would have enabled both withincountry variation and between-country statistical comparisons to be performed. In contrast, 293 our approach enabled only between-country variation to be analysed, albeit not statistically 294 295 compared. However, this approach allowed for a wide range of important nutrients and other 296 compounds relevant to health to be included, which we felt outweighed the limitations of 297 pooling samples. Despite the limitation of pooling prune samples from each country of origin, 298 a robust sampling methodology was adopted based upon standards used for food 299 composition databases to ensure high levels of representativeness in each pooled sample, 300 including sourcing from a range of major retail centres in numerous sampling countries.

301

Further limitations include the small number of samples from Argentina, which may therefore 302 303 not be fully representative of Argentinian prunes available across Europe. Any differences 304 attributable to country of origin can only be ascertained by controlling other factors that can 305 influence variation in nutrient composition. The sampling protocol attempted to account for seasonal variation and storage conditions by purchasing samples at the same time of year and 306 307 within use by dates, and minimised changes in composition between purchase and analysis. However, given that all prunes were sampled at point of sale, pre-purchase confounding 308 309 variables such as exposure to heat, light and humidity could not be controlled for. This might be relevant if these factors influence nutrient composition as some previous data suggests, 310 311 however, although such analyses are of important academic and commercial interest, from a practical perspective the consumer cannot currently impact post-harvest/pre-purchasing 312 processing. 313

314

315 Conclusion

The current study provides evidence that small differences in dietary fibre, sorbitol and phenolic content may exist between prunes of different countries of origin. To our knowledge, this is the first study to provide a comprehensive and comparable compositional analysis of

319	prunes of USA (Californian) origin, updating the currently available data reported in the USDA
320	and McCance and Widdowson UK databases. This allows for a more accurate measurement
321	of nutrient intake for future dietary intervention studies. The current study has highlighted
322	the need for thorough and transparent documentation of sampling methods used to produce
323	data for national databases. Furthermore, to eliminate artificial differences in energy content
324	between different databases, carbohydrate values should be expressed using the same
325	method and energy should be calculated using the same conversion factors.
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USA (California)			France				Chile		Argentina			
Sampling	Number of samples		es Weight	Number of samples		Weight	Number of samples		Weight	Number of samples		Weight
country	Purchased	Analysed	contributing to analysis	Purchased	Analysed	contributing to analysis	Purchased	Analysed	contributing to analysis	Purchased	Analysed	contributing to analysis
			125 g			125 g			125 g			
Germany	4	4	(31.25 g per	3	3	(41.6 g per	4	4	(31.25 g per	0	0	-
			sample)			sample)			sample)			
			125 g			125 g			125 g			
Italy	5	5	(25 g per	2	2	(62.5 g per	2	2	(62.5 g per	0	0	-
			sample)			sample)			sample)			
			125 g			125 g			125 g			
UK	5	5	(25 g per	5	5	(25 g per	5	5	(25 g per	0	0	-
			sample)			sample)			sample)			
						125 g						
France	0	0	-	16	8**	(15.625 g	0	0	-	0	0	-
						per sample)						
			125 g						125 g			500 g
Spain	5	4*	(31.25 g per	4	0***	-	4	4	(31.25 g per	5	5	(100 g per
			sample)						sample)			sample)
Total	19	18	500 g	30	18	500 g	15	15	500 g	5	5	500 g

Table 1: Number of prune samples purchased, pooled and analysed, by sampling country and country of origin.

*One sample exceeded the best before date and was excluded ** Only 8 samples required *** Samples all labelled Spanish/French origin and thus excluded

	Method of analysis	Country of origin (per 100 g wet weight)				Country of origin (per 100 g dry weight)			
			Chile	France	Argentina	USA	Chile	France	Argentina
Moisture (g)	Oven drying	30.9	30.5	33.2	28.7	-	-	-	-
Ash (g)	Incineration (muffle furnace)	1.58	1.53	1.36	1.38	-	-	-	-
Energy (kcal)	Multiplying macronutrients by Atwater factors	230	235	228	241	333	337	340	338
Protein (g)	Total N (Dumas, TruSpec analyser) x 6.25	2.5	2.1	1.6	2.0	3.6	2.9	2.4	2.8
Fat (g)	Soxhlet method	<0.2	<0.2	<0.2	<0.2	<0.2	<0.2	<0.2	<0.2
Total carbohydrate (g)	Calculated 'by difference'	65.1	65.9	63.9	68.0	94.1	94.9	95.4	95.3
Available carbohydrate (g)	Calculated 'by difference'	56.9	58.2	55.6	59.9	82.2	83.8	83.0	84.0
Starch (g)	Enzymatic hydrolysis (Megazyme)	6.6	5.7	1.9	6.1	9.5	8.1	2.8	8.5
Total sugars (g)	Ion-exchange chromatography	38.2	41.3	40.7	42.2	55.2	59.4	60.7	59.1
Fructose	Ion-exchange chromatography	14.0	16.2	16.0	16.9	20.2	23.3	23.9	23.6
Glucose	Ion-exchange chromatography	24.2	25.1	24.6	25.3	35.0	36.1	36.8	35.5
Galactose	Ion-exchange chromatography	<0.01	<0.01	<0.01	<0.01	<0.01	< 0.01	< 0.01	<0.01
Lactose	Ion-exchange chromatography	<0.01	<0.01	<0.01	<0.01	<0.01	< 0.01	< 0.01	<0.01
Maltose	Ion-exchange chromatography	<0.01	<0.01	<0.01	<0.01	<0.01	< 0.01	< 0.01	<0.01
Sucrose	Ion-exchange chromatography	<0.01	<0.01	<0.01	<0.01	<0.01	< 0.01	< 0.01	<0.01
Sugar alcohols (g)									
Sorbitol	Ion-exchange chromatography	14.8	13.8	11.2	15.5	21.4	19.9	16.7	21.7
Dietary Fibre (g)									
Total fibre	Enzymatic-gravimetric (AOAC 991.43)	8.2	7.7	8.3	8.1	11.9	11.1	12.4	11.4
Total fibre	Enzymatic-gravimetric (AOAC 2011.25)	12.0	11.5	8.4	8.9	17.4	16.5	12.5	12.5
Insoluble fibre	Enzymatic-gravimetric (AOAC 2011.25)	4.5	5.7	4.0	4.3	6.5	8.1	6.0	6.0
Soluble fibre	Enzymatic-gravimetric (AOAC 2011.25)	7.6	6.3	4.4	4.6	11.0	9.1	6.5	6.4
Total NSP	Englyst et al (1994)	6.2	5.8	5.8	5.9	9.0	8.3	8.7	8.3
Insoluble NSP	Englyst et al (1994)	2.0	1.8	2.0	1.6	2.9	2.6	3.0	2.2
Soluble NSP	Englyst et al (1994)	4.3	4.1	3.9	4.3	6.1	5.8	5.7	6.0
Cellulose	Englyst et al (1994)	0.2	0.3	0.3	0.3	0.3	0.4	0.4	0.4

Table 2: Composition of prunes from the pooled samples from USA, Chile, France and Argentina, purchased in Europe

	Method of analysis	Country of origin (per 100 g wet weight)				Country of origin (per 100 g dry weight)			
		USA	Chile	France	Argentina	USA	Chile	France	Argentina
Lignin	Enzymatic-gravimetric (AOAC 994.13)	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
Fructans	Enzymatic spectrophotometric (AOAC 999.03)	0.3	0.2	0.3	0.3	0.4	0.3	0.5	0.4
Phenolic compounds (mg									
Chlorogenic acid	UPLC with MS-MS	3.6	1.3	3.9	1.3	5.2	1.9	5.8	1.8
Neochlorogenic acid	UPLC with MS-MS	89.3	39.8	92.0	40.3	129.1	57.2	137.3	56.4

UPLC with MS-MS: Ultra-performance liquid-chromatography tandem mass spectrometry

Table 3: Composition of prunes originating from the USA (California), as analysed in the current study, compared with data from the USDA nutrient database and McCance and Widdowson's The Composition of Foods. Values are units per 100 g wet weight.

	Analysis in the current study	,	Databa	ise values	
	Method of analysis in the current study	USA (Californian) (wet weight per 100 g)	USDA (wet weight per 100 g)	McCance & Widdowson (wet weight per 100 g)	
Water (g)	Oven drying	30.9	30.9	31.1	
Ash (g)	Incineration (muffle furnace)	1.58	NR	2.64	
Energy (kcal)	Multiplying macronutrients by Atwater factors	230	240	141	
Protein (g)	Total N content (Dumas, TruSpec analyser) x 6.25	2.5	2.2	2.5	
Fat (g)	Soxhlet method	<0.2	0.4	0.4	
Fatty acids (g)					
Saturated Fatty Acids	Gas chromatography with flame ionisation detection	<0.1	<0.1	NR	
Mono-unsaturated Fatty Acids	Gas chromatography with flame ionisation detection	<0.1	<0.1	NR	
Polyunsaturated Fatty Acids	Gas chromatography with flame ionisation detection	<0.1	<0.1	NR	
Trans-unsaturated Fatty Acids	Gas chromatography with flame ionisation detection	<0.1	NR	NR	
Total carbohydrate (g)	Calculated 'by difference'	65.1	63.9	NR	
Available carbohydrate (g)	Calculated 'by difference'	56.9	NR	34.0	
Starch (g)	Enzymatic hydrolysis (Megazyme)	6.6	5.1	0.0	
Total sugars (g)	Ion-exchange chromatography, HPAEC-PED	38.2	38.1	34.0	
Fructose	Ion-exchange chromatography, HPAEC-PED	14.0	12.5	12.1	
Glucose	Ion-exchange chromatography, HPAEC-PED	24.2	25.5	17.9	
Galactose	Ion-exchange chromatography, HPAEC-PED	<0.01	0.0	NR	
Lactose	Ion-exchange chromatography, HPAEC-PED	<0.01	0.0	0.0	
Maltose	Ion-exchange chromatography, HPAEC-PED	<0.01	0.1	0.0	
Sucrose	Ion-exchange chromatography, HPAEC-PED	<0.01	0.2	4.1	

	Analysis in the current st	Database values			
	Method of analysis in the current study	USA (Californian) (wet weight per 100 g)	USDA (wet weight per 100 g)	McCance & Widdowson (wet weight per 100 g)	
Sugar alcohols (g)					
Sorbitol	Ion-exchange chromatography, HPAEC-PED	14.8	NR	NR	
Dietary Fibre (g)					
Total dietary Fibre	AOAC 991.43	8.2	7.1	NR	
Total dietary Fibre	AOAC 2011.26	12.0	NR	NR	
Insoluble dietary fibre	AOAC 2011.26	4.5	NR	NR	
Soluble dietary fibre	AOAC 2011.26	7.6	NR	NR	
Non-starch polysaccharides	Englyst et al (1994)	6.2	NR	5.7	
Insoluble NSP	Englyst et al (1994)	2.0	NR	NR	
Soluble NSP	Englyst et al (1994)	4.3	NR	NR	
Cellulose	Englyst et al (1994)	0.2	NR	NR	
Lignin	Enzymatic-gravimetric method (AOAC 994.13)	0.009	NR	NR	
Fructans	Enzymatic spectrophotometric (AOAC 999.03)	0.3	NR	NR	
Phenolic compounds (mg)					
Chlorogenic acid	UPLC with MS-MS	3.6	NR	NR	
Neochlorogenic acid	UPLC with MS-MS	89.3	NR	NR	
Minerals					
Calcium (mg)	ICP-OES	45.0	43.0	34.0	
Iron (mg)	ICP-OES	0.7	0.9	2.6	
Potassium (mg)	ICP-OES	622	732	760	
Magnesium (mg)	ICP-OES	47.0	41.0	24.0	
Sodium (mg)	ICP-OES	9.8	2.0	11.0	

	Analysis in the current	Database values			
	Method of analysis in the current study	USA (Californian) (wet weight per 100 g)	USDA (wet weight per 100 g)	McCance & Widdowson (wet weight per 100 g)	
Phosphorus (mg)	ICP-OES	68.1	69.0	73.0	
Zinc (mg)	ICP-OES	0.4	0.4	0.4	
lodine (mg)	ICP-OES	3.0	NR	NR	
Selenium (µg)	ICP-OES	30.0	0.3	3.0	
Vitamins					
Riboflavin (mg)	HPLC	0.0	0.2	0.2	
Niacin (mg)	HPLC	1.1	1.9	1.3	
Vitamin B6 (mg)	HPLC	0.3	0.2	0.2	
Biotin (µg)	Plasmon resonance technology	20.0	NR	Tr	

NR = Not reported

Tr = trace

ICP-OES Inductively Coupled Plasma Optical Emission Spectrometry HPAEC-PED High-Performance Anion-Exchange Chromatography Coupled with Pulsed Electrochemical Detection

HPLC High Performance Liquid Chromatography