Postprint of: This version of the article has been accepted for publication, after peer review (when applicable) and is subject to Springer Nature's AM terms of use, but is not the Version of Record and does not reflect post-acceptance improvements, or any corrections. The Version of Record is available online at: <https://doi.org/10.1007/s00217-023-04390-y>

Postprint of: Kim, Y.M., Lubinska-Szczygeł, M., Polovka, M. et al. Properties of some fruit wines. Eur Food Res Technol (2023)

Metadata of the article that will be visualized in OnlineFirst

Recently we reported about the consumption of red wines from grapes, having several health properties. There are different types of wines that originated from grapes and other fruits. In the present study fruit wines from persimmon, kiwifruit and pomegranate were investigated and compared for theirantioxidant ability, usingcupricion reducingantioxidant capacity (CUPRAC)and the 2,2-diphenyl-1-picrylhydrazyl (DPPH) assays.

To the fruit wines were applied the same methods of investigation as to the traditional ones made from grapes. The results showed the highest antioxidant activity of pomegranate, followed by kiwifruit and persimmon wines. Fourier transform infrared (FTIR) spectroscopy was used in order to correlate these results. The interaction of wine bioactive compounds with the main serum proteins in the human metabolism, such as human serum albumin (HSA), globulin (GL), and fibrinogen (FB), showed that pomegranate wine possesses higher quenching properties than kiwifruit and persimmon wines. All determined fluorescence indices have a direct correlation with the bioactivity of polyphenols and not with the content of alcohol. We hypothesize that the results of the interaction of main human serum proteins with bioactive compounds of wines can be additional predictors of their health properties. The used analytical methods for quality of fruit wines can be applied to a wide range of fruits and vegetables.

Graphical Abstract:

Keywords (separated by '- Polyphenols - Pomegranate - Kiwifruit - Persimmon - Fluorescence - FTIR bands - Antioxidants - Human serum proteins - Quenching ')

Footnote Information

ORIGINAL PAPER

1

Properties of some fruit wines 2

- **Young Mo Kim1 · Martyna Lubinska‑Szczygeł² · Martin Polovka3 [·](http://orcid.org/0000-0001-8398-2713) Blanka Tobolkova3 [·](http://orcid.org/0000-0003-4809-2840)** 3
- **Pitipong Thobunluepop⁴ [·](http://orcid.org/0000-0001-6592-5141) Yong Seo Park5 [·](http://orcid.org/0000-0002-5827-3584) Kyung Sik Ham[6](http://orcid.org/0000-0001-8372-6445) · Yang Kyun Park6 [·](http://orcid.org/0000-0002-9609-064X) Seong Gook Kang⁶ [·](http://orcid.org/0000-0002-8498-120X)** 4
- **Dinorah Barasch7 · Alina Nemirovski7 · Shela Gorinstein7** 5

Received: 22 May 2023 / Revised: 7 October 2023 / Accepted: 13 October 2023 6

© Springer-Verlag GmbH Germany, part of Springer Nature 2023 7

Abstract 8

- Recently we reported about the consumption of red wines from grapes, having several health properties. There are different **[AQ1](#page-18-0)** 9
- types of wines that originated from grapes and other fruits. In the present study fruit wines from persimmon, kiwifruit and 10
- pomegranate were investigated and compared for their antioxidant ability, using cupric ion reducing antioxidant capacity 11
- (CUPRAC) and the 2,2-diphenyl-1-picrylhydrazyl (DPPH) assays. 12
- To the fruit wines were applied the same methods of investigation as to the traditional ones made from grapes**.** The results 13
- showed the highest antioxidant activity of pomegranate, followed by kiwifruit and persimmon wines. Fourier transform 14
- infrared (FTIR) spectroscopy was used in order to correlate these results. The interaction of wine bioactive compounds with 15
- the main serum proteins in the human metabolism, such as human serum albumin (HSA), globulin (GL), and fibrinogen 16
- (FB), showed that pomegranate wine possesses higher quenching properties than kiwifruit and persimmon wines. All deter-17
- mined fluorescence indices have a direct correlation with the bioactivity of polyphenols and not with the content of alcohol. 18
- We hypothesize that the results of the interaction of main human serum proteins with bioactive compounds of wines can 19 20
- be additional predictors of their health properties. The used analytical methods for quality of fruit wines can be applied to
- a wide range of fruits and vegetables. 21

Graphical Abstract 22 23

Extended author information available on the last page of the article

A1

- Persimmon · Fluorescence · FTIR bands · Antioxidants · 26
- Human serum proteins · Quenching 27

Introduction 28

calluly part of a diet [1, 2). Dietary persistmon (part)

and the main of the distribution and models [3-5]. Kiniching the models and the models are the stationary entrinois and the model and distribute calluly perceptivel Most of the tropical and traditional fruits are important sources of antioxidants, vitamins, and minerals and form a very healthy part of a diet [1, 2]. Dietary persimmon (peel and pulp) was reported to exert hypolipidaemic properties in some hyperlipidemic animal models [3–5]. Kiwifruits exhibit mostly antioxidative and antihypercholesterolemic properties. Analytical and processing methods affect the physicochemical and biological properties of kiwifruit-derived ingredients and depend on different cultivars which were intensively studied and compared with persimmon [6–8]. Pomegranate is rich in colored and colorless phenolic compounds, varied depending on cultivars, and contains a high amount of polyphenols, anthocyanins, catechins, tannins, gallic and ellagic acids, and possesses high health properties [9, 10]. Concerning the antioxidant and healthy properties of presented fruits, the manufacture of wine from fruits, other than grapes, has been developed in recent years [11]. As it was shown above, kiwifruits, persimmons, and pomegranates contain high levels of bioactive compounds, especially polyphenols, and it is important to preserve them in the preparation of wines, juices and other varieties of food products [12–14]. The fruit wines are less popular compared to grape wines, but the raw materials of these wines, are rich in phenolic antioxidants. Polyphenols have a strong bioactivity as an active element in foods, fruits, cereals, vegetables and beverages (beers and wines). Many studies [2, 3, 5, 8] have demonstrated that polyphenols have also strong effects on the vascular system by lowering blood pressure, increasing antioxidant defenses, inhibiting platelet aggregation and low-density lipoprotein oxidation [15–17]. These properties are extensively used in the prevention and treatment of coronary artery disease (CAD) [1–3]. The purpose of the present report was to study the antioxidant, binding and healthy properties of bioactive compounds, which are present in kiwifruit, persimmon, and pomegranate wines, applying the advanced analytical methods, such as FTIR and fruorescence. The antioxidant properties of wines were determined by cupric ion reducing antioxidant capacity (CUPRAC) and the 2,2-diphenyl-1-picrylhydrazyl (DPPH) assays. The polyphenol spectrum of wines was determined by Fourier transform infrared (FTIR) spectroscopy. The binding abilities of wines with human serum albumin (HSA), globulin (GL) and fibrinogen (FB) were estimated by 3D- fluorescence and correlated with their polyphenol contents. **[AQ2](#page-18-1)** 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61 62 63 64 65 66 37 68 69 70 <u>10</u>

72

73

74

76

83

Materials and methods

Materials

The chemicals 6-Hydroxy-2,5,7,8-tetra-methylchroman-2-carboxylic acid (Trolox), 1,1-diphenyl-2-picrylhydrazyl (DPPH), lanthanum(III) chloride heptahydrate, $CuCl₂ × 2H₂O$, 2,9-dimethyl-1,10-phenanthroline (neocuproine), quercetin, tannic acid, human serum albumin (HSA), fibrinogen (FB), globulin (GL), phosphate buffer and Folin-Ciocalteu reagent (FCR) were purchased from Sigma (St. Louis, MO, USA) and Fluka Chemie GmbH (Buchs, Switzerland). 75 77 78 79 80 81 82

Wine samples

Wines were bought in Israel and South Korea and were used in the present investigation study. Each kind of wine was purchased in the amount of five samples in several places, but from the same year of vintage and showed the same shelf life. The bottles with a range of alcohol in the same volume were frozen at – 80 °C to assess their antioxidant status and bioactivity. Pomegranate dry wine with 13.8% alcohol of 2018 vintage was purchased from Rimon wineries, Israel. Persimmon wine (Persimun wine (Regular) with 12.0% alcohol was delivered from Agricultural Corporation Cheongdo Persimun wine, Cheongdo, Gyeongsangbuk-do, Korea. Kiwi (Darae) wine with 8.0% alcohol was made from kiwis (chamdarae in Korean) grown in an environmentally friendly manner from farms in Sacheon, Gyeongsangnam-do (South Gyeongsang Province). As kiwis are consumed after ripening, a bottle of Darae Wine is made after going through a 10 months maturation. The samples were produced by Darae wine shop, GyeongnamSacheon-siMiryong-Gil, Korea. Samples for analysis were taken out of the refrigerator and were diluted according to the methods explained below. 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 100 101 102 103

Determination of bioactive compounds

104

105

The total phenolic amount (TP) was measured by using the Folin–Ciocalteu method [\[18](#page-15-10)], using 250 μL of wine mixed with 1000 μ L of sodium carbonate (7.5%) and 1250 μ L of Folin–Ciocalteu's (10% in water) reagent. The mixture was incubated for 15 min at 50 °C in the dark (water bath) and measured at 765 nm, using a spectrophotometer (Hewlett-Packard, model 8452A, (Rockville, MD, USA). Gallic acid was used as the standard, and the results were expressed as milligrams of gallic acid equivalent per liter (mg GAE/L). 106 107 108 109 110 111 112 113 114

Experient and typerator and experient of the liquid minimal selection of the particular temperature of the state of th The anthocyanin content (AC) in wines was measured in aliquots of 250 μL of the wine sample which was poured into a tube with 2 mL of potassium chloride solution (0.025 M), adjusted to pH 1 with concentrated HCl. The mixture was incubated at room temperature for 20 min. In another tube, $250 \mu L$ of wine was mixed with 2 mL of sodium acetate solution (0.4 M, pH 4.5) and incubated at room temperature for 20 min. The absorbance of an aliquot of 300 μL of each wine sample was measured at 520 and 700 nm. The results were expressed as milligrams of cyanidin 3-glucoside equivalent per L (mg C3G/L) [19, 20]. The total tannins (TNs) were estimated by using spectrophoto-metric measurements of 0.5 mL of wine, where 3 mL of a 4% methanol vanillin solution and 1.5 mL of concentrated hydrochloric acid were added. The mixture was allowed to stand for 15 min. The absorption of the samples and a blank against water was measured at 500 nm [20, 21]. Some bioactive compounds, such as quercetin, were determined with a high-performance liquid chromatography HPLC system [20, 22]. A volume of 50 mL of each of the wine samples was extracted three times with 25 mL of diethyl ether and then three times with 25 mL of diethyl acetate, and the organic fractions were combined. After 30 min of drying with anhydrous $Na₂SO₄$, the extract was filtered through a Whatman-40 filter and evaporated to dryness in a rotary evaporator. The residue was dissolved in 2 mL of methanol/water (1:1, *v/v*) and analyzed by HPLC. A Waters (Milford, MA, USA) chromatograph equipped with a 600-MS controller, a 717 plus autosampler, and a 996 photodiode-array detector was used [20]. For the HPLC analysis, an aliquot $(50 \mu L)$ was injected into the column and eluted at the temperature of 20 °C. Total ascorbic acid content [TAAC, mg ascorbic acid (AA) per L] was evaluated in water wine extracts, where 100 mg of the freeze-dried wine sample was extracted with 5 mL water. Then, the CUPRAC method was conducted and formed bis (Nc)-copper (I) chelate was determined spectrophotometrically at 450 nm [23]. 115 116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131 132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147 148 149 150

Antioxidant capacity asssays 151

For the cupric-reducing antioxidant capacity (CUPRAC) assay [[23](#page-15-14), 24] fruit wines were diluted in a ratio of 1:10 (v/v) with dH_2O . About 1.0 mL of each of the three solutions containing 0.010 M Cu (II), ammonium acetate buffer at pH 7.0, and 0.0075 M neocuproine (2,9-dimethyl-1,10-phenanthroline) in EtOH was mixed with 0.5 mL of the appropriately diluted sample together with 0.6 mL of $dH₂O$ in a tube. The reaction mixture was left for 1 h in the dark, and then the absorption was measured at 450 nm [\[20,](#page-15-11) [23](#page-15-14), [24](#page-15-15)]. The antioxidant capacity was measured by using 1, 1-diphenyl-2-picrylhydrazyl (DPPH) in 25 μL of the sample, which was mixed with 180 μL of DPPH radical at 6 mM and measured at 517 nm every 30 s for 10 min. Trolox was used as a standard for two antioxidant methods, and the results were

expressed as millimoles of Trolox equivalent per liter (mmol TE/L) [[20,](#page-15-11) [25\]](#page-15-16). 166 167

> 168 169

182

198

Fourier transform infrared spectra of polyphenols in wines

Total phenols in the investigated fruit wines extracts were studied by IR spectroscopy. 170 171

The fruit wines were evaporated from the amount of alcohol, transferred under the liquid nitrogen in order to prevent their oxidation, and then lyophilized. The dry powder was extracted with methanol (25 mg/mL), then evaporated and lyophilized. The polyphenol extracts were used for FTIR measurements. A Nicolet iS 10 Fourier transform infrared (FTIR) Spectrometer (ThermoScientific Instruments LLC, Madison, (WI, USA)), with the smart iTRTM attenuated total reflectance (ATR) accessory was used to record IR spectra [26]. 172 173 174 175 176 177 178 179 180 181

Fluorometric studies

The properties of bioactive substances in wines were determined by using three-dimensional (3D-FL) fluorescence (model FP-6500, Jasco spectrofluorometer, serial N261332, Tokyo, Japan). The 3D-FL was measured at emission wavelengths between 200 and 795 nm, and the initial excitation wavelength was 200 nm. For comparison of the obtained results quercetin and tannic acid were used [20]. Standard phenolic solutions, such as tannic acid and quercetin were prepared daily by dissolving at a concentration of 10 mM in methanol and then diluting with 10 mM phosphate buffer at pH 7.4. The initial fluorescence intensities of fibrinogen, albumin, and globulin were measured before their interactions with the investigated wines. As mentioned above, the changes in the fluorescence intensities were used in the estimation of the binding activities [17, 20]. 183 184 185 186 187 188 189 190 191 192 193 194 195 196 197

Statistical analysis

All data obtained were calculated on the basis of a statistical analysis of Duncan's multiple range test. Values were mean $s \pm SD$ per liter of 25 measurements, representing the commercial status of the wines and their replicates. Five replications of five wine samples were used. To determine the statistical significance at the 95% interval of reliability, a one-way analysis of variance (ANOVA) was used. 199 200 201 202 203 204 205

For the purposes of distinguishing the fruit wines, the discriminant procedure was realized by means of the Unistat® statistical package (Unistat, London, United Kingdom) using the entire data, involving methods of principal component analysis (PCA), principal component factoring with varimax rotation (PCF) and canonical discriminant analysis (CDA). The convergence criteria of discriminant analysis 206 207 208 209 210 211 212

 \mathcal{D} Springer

were chosen for a standardized proximity matrix with the maximum number of iterations, 50. The following stepwise selection criteria were used: tolerance – 0.001, F statistic: F to enter -3.8416 , F to remove -2.7056 . 213 214 215 216

Results and discussion 217

Bioactivities of wine samples 218

The bioactivities of the main compounds in wines and their antioxidant activities were determined (Fig. 1). As can be seen (Fig. 1c) that pomegranate wine contains the highest amount of polyphenols and anthocyanins, followed by kiwi (Fig. [1](#page-7-0)b) and persimmon wines (Fig. 1a). Such results are in agreement with some recent reports [14], where the amount of total polyphenols in pomegranate wine was similar to the present results (Fig. 1c). Quercetin in pomegranate wine samples was 12.1 mg/L. The amount of anthocyanins was about 105.4 mg C3G/L. The amount of anthocyanin's concentration oscillated between 136 and 23 mg/100 mL for Wonderful and Mollar de Elche juices, respectively [27]. The amount of anthocyanins, as other bioactive compounds, varied in several genotypes and showed different values in the variety of cultivars, as cyanidin-3,5-O-diglucoside and pelargonidin-3,5-O-diglucoside in Santa Tecla population and were 97.64 mg/L and 40.29 mg/L, respectively. The 219 220 221 222 223 224 225 226 227 228 229 230 231 232 233 234 235

amount of bioactive compounds (polyphenols, anthocyanins, and quercetin) was in correlation with the antioxidant activity and showed the value of total antioxidant activity by CUPRAC of 27.7 mM TE/L and by DPPH of 8.16 mM TE/L. These data are comparable with the same report [\[27](#page-15-18)], where the antioxidant activity was estimated at 9.8 mM/L. The total antioxidant activity values ranged between 221.5 and 36.73 μmol TE/100 mL of juice [[28\]](#page-15-19). Such variety in the amounts of bioactive substances can be explained by the differences in cultivars, climate, and production processes. Pomegranate wines usually have high total phenolics and were 1.5 times higher than the amount of phenolics in green tea of 1029 mg/kg [29]. Persimmon wines showed the lowest polyphenol content in comparison with the other two wines (Fig. 1a). The obtained results depend on the cultivar used for wine production $[30]$. Most of the cited studies were conducted with whole persimmon or persimmon peel or persimmon pulp as test materials. It is well known that persimmon fruit contained a large number of components such as condensed tannin and polyphenols [1, 3, 31]. According to some authors, Wistar rats fed with a hypercholesterolemic diet enriched in persimmon (7%) had lower values of plasmatic lipids (cholesterol, triglycerides, LDL) after 4 weeks compared to control rats [1]. As it was shown previously that persimmon, kiwifruit and pomegranate have high antioxidant activities, then the wines prepared from the same fruits possess health properties. Due to the high content 236 237 238 239 240 241 242 243 244 245 246 247 248 249 250 251 252 253 254 255 256 257 258 259 260 261 262

Fig. 1 Bioactive compounds, antioxidant activities and total binding properties of fruit wines **a** persimmon, **b** kiwi, **c** and pomegranate/L. Values are means \pm SD of 5 measurements; Means within bars with different superscripts are statistically different ($p < 0.05$; Student's t test). Abbreviations: GAE, gallic acid equivalent; C3G, cyanidin

3-glucoside equivalent; AA, ascorbic acid; CUPRAC, Cupric reducing antioxidant capacity; DPPH, 1, 1-Diphenyl-2-picrylhydrazyl method; TE, Trolox equivalent; HSA, human serum albumin; GL, human serum globulin; FB, fibrinogen

$\circled{2}$ Springer

of antioxidants, persimmon could be of help in reducing or preventing LDL oxidation and thus the development of atherosclerosis [32]. It was shown in some reports that the high molecular weight of persimmon tannins is responsible for the hypocholesterolemic effect of persimmon fruit and it might exert the hypolipidemic effect, improving the antioxidant profile of human serum [[3,](#page-14-2) [4,](#page-15-20) 33]. It is obvious that the properties of fruits have to be prevented during processing. The high temperature increased the contents of phenolic and polymeric pigment in wine: with polyphenols of 871.3 mg/L, quercetin of 0.04 mg/L, and total tannins of 311.29 mg/L, which are similar to the one shown in this report (Fig. 1a). The values of antioxidant activity by DPPH were slightly lower than in the present report of 775.1–1326.0 μmol/L [34]. It is generally approved that a moderate consumption of fermented beverages prevents metabolic disorders due to the antioxidant properties of phenolic compounds. Persimmon liqueur was prepared from fresh or dry fruit by: (1) extraction with alcohol, (2) fermentation of fresh fruit, and (3) extraction of dry fruit with distilled alcohol from an extract. Alteration in the ratio of raw and dry materials to a solvent, conditions of fermentation, and the degree of distillation resulted in a beverage with high aroma and taste, polyphenols, and proteins. Similar results were obtained with and without fermentation [13, 35]. Persimmon wine may offer nutritional and medicinal value as it contains compounds that may be beneficial to health. Phenolic compounds are important to wine quality as they contribute to antioxidant activity, aroma formation, colloidal stability, and sensory properties. High temperature also induced the increase of total tannins, compounds that contribute to antioxidant activity [36]. Kiwifruit has a beneficial impact on inflammatory processes, atherogenesis, and thrombogenesis, exerting also hypolipidaemic activity, and preventing diabetes development. Such properties were only found for kiwifruit and not for other fruits according to the EVIDENT study [37]. These properties of kiwifruit characterize the wines from this fruit. 263 264 265 266 267 268 269 270 271 272 273 274 275 276 277 278 279 280 281 282 283 284 285 286 287 288 289 290 291 292 293 294 295 296 297 298 299

Kiwifruit possesses similar properties as pomegranate with the highest amount of vitamin C (Fig. 1b). Cited reports described that kiwifruit is used as a juice, vinegar, dried slices, jam, and wine [38, 39]. The quality characteristics of kiwifruit wine made from over-ripened fruit treated with pectinase showed higher values of wine in many aspects such as sensory value, alcohol and total phenolics content, antioxidant activity, minerals, and production yield [[12](#page-15-6)]. Several domestic varieties of kiwifruit were utilized for the processing of wines. It was shown that the wines have high antioxidant activity. The effects on human health depend on the amount of consumed wines and on their bioavailability. 'Daeheung' had total phenols of 790 mg/L, which was the highest among wines, followed by 'Haenam' and 'Golden King'. 'Daeheung' also showed the highest antioxidant activity (22.55 mMTE/L), while 'Arimold' showed the

lowest one (10.91 mMTE/L). These values are in accordance with the present results (Fig. [1b](#page-7-0)). Two fruit wines (kiwi and persimmon) were delivered from South Korea, where the consumption of fruit wines is high. Various fruits such as raspberry (3,106 233 tons/year), plum (579 tons/year), and mulberry (434 tons/year), and relatively low amounts such as persimmon (83 tons/year) and kiwifruit (70 tons/year) are processed 235 into wine [[40,](#page-16-0) [41\]](#page-16-1). 316 317 318 319 320 321 322 323

324

FTIR spectra of wines

Experient wores: which polyphenols of 871.1 mpl. **FIR spectra of wines**
 Unity polyphenols of 371.29 mpl.
 Unity holyphenols of 311.29 mpl.
 Unity and total canning of 311.29 mpl.
 Unity and to the one shown in th The infrared spectra of persimmon, kiwifruit, and pomegranate wines were measured in the frequency range of 4000–800 cm⁻¹ (Fig. 2a, b, c). Standards such as quercetin and tannic acid (Fig. 2d, e) were done in order to identify the peaks. The O–H and C–H stretching frequencies in the polyphenols are found in the 3500–2600 cm−1 region and C–H stretching vibration occurred in the region of 2900 to 2800 cm−1 (for kiwi and persimmon wines as 2936.76; 2949.87; 2834.26 cm⁻¹). In the region of 2937–2950 cm⁻¹, the CH, $CH₂$, and CH₃ stretching vibrations, derived from carbohydrates and sugars, which are shown for kiwi and persimmon wines [42]. In quercetin, the peak obtained in the range of 3261 cm−1 represented the O–H stretching vibration due to the intra-molecular hydrogen bonding. The band observed at 1662 and 1607 cm^{-1} assigned to carbonyl C=O stretching vibration. The band obtained at 1519 cm^{-1} assigned for NO₂ bending vibration, the peak at 1447–1407 cm⁻¹ for C–O, and the band at 1259–1215 cm⁻¹ allocated to C–O–C of ester for quercetin compound. The prominent peak at $1166-1092$ cm⁻¹ indicated the stretching vibration of the C–O–C group. The presented results were similar to the reported [43]. Pomegranate wine does not show vibration in this region. In tannic acid, one characteristic peak was found at 3344 cm^{-1} which shows the stretching vibration of the O–H group. The peak at 1698 cm^{-1} represented the stretching of the carbonyl (C=O) group. The peak at $1537-1446$ cm⁻¹ allocated the carboxylic acid (O–C–O) and at $1312–1190$ cm⁻¹ showed the bending vibration of the O–H group. The band between 1024 and 956 cm−1 is assigned to the C=O group of molecules. The band at $756-868$ cm⁻¹ is due to the meta-substitution of the aromatic protons. The presented results of the peaks were equal to other reports [\[43,](#page-16-3) [44\]](#page-16-4). The major protein bands include amides I $(C=O$ stretching coupled with N–H bending) vibrations at approximately 1650 cm−1 which appeared in kiwi and pomegranate wines at 1644 and 1648 cm−1 (Fig. [2](#page-9-0)b,c). The in-plane bending vibration of aromatic CH is detected at 1100 cm^{-1} , in kiwi wine and a CO stretching vibration is produced at 1062 cm^{-1} for kiwi and pomegranate wines. For signals with wavelengths smaller than 900 cm⁻¹, the aromatic CH stretching vibration is detected at 891 and 883 cm−1 for kiwi and persimmon wines. The 325 326 327 328 329 330 331 332 333 334 335 336 337 338 339 340 341 342 343 344 345 346 347 348 349 350 351 352 353 354 355 356 357 358 359 360 361 362 363 364 365 366

due to the stretching of the carbonyl C=O group. Peaks at 1062 cm−1 (for pomegranate and kiwi wines) are ascribed to the –COH group of sugars in glycosylated phenols. The 370 371 372

² Springer

37 368 ₃₉

peak at 1019 cm−1 was ascribed to the phenolic C–OH in persimmon wine. The peak at 1735 cm⁻¹ was assigned to the carbonyl C=O stretching band of protonated carboxylic acid, characteristic of the galloyl unit of hydrolysable tan-nins for kiwi wine [\[45](#page-16-5)]. Peaks at 1648 and 1644 cm^{-1} were assigned to the –COO– stretching for pomegranate and kiwi wines. The peak at 1227 cm⁻¹ corresponds to the C–OH of phenols. Peaks at 1163 and 1166 cm⁻¹ were ascribed to the C–OH stretching in glycosylated phenols for pomegranate and kiwi wines. Pomegranate and kiwi wines have in common absorptions at 1062, 1165, 1508, and 1647 cm⁻¹. The FTIR spectra in the region between 1735 and 900 cm⁻¹ can serve as fingerprint and comparison of the investigated wines. 373 374 375 376 377 378 379 380 381 382 383 384 385

Fluorescence properties of wines 386

The antioxidant's strong affinity to human serum proteins and synergism in bioactivity are the main indices in the health application of wines [4, 5, 10, 20]. The chemical, phenolic and antioxidant characteristics of the wines were assessed by measurement of tannins, anthocyanins, and phenolic compounds (Figs. 1, 2). Our results were similar to the wines prepared by classical maceration. These properties of the wine bioactive compounds, mainly polyphenols, in interaction with the main human serum proteins are shown for all investigated samples (Fig. 3). As it was found previously that 387 388 389 390 391 392 393 394 395 396

persimmon wine has the lowest antioxidant activity in comparison with other wines, it is expected from our previous reports that the quenching properties of persimmon wine are lower than in other samples [[7,](#page-15-22) [8,](#page-15-2) [17](#page-15-9), [20](#page-15-11)]. The calculations were done on the basis of decreasing the initial fluorescence intensity (FI) of human serum proteins before interaction with bioactive wine substances (Fig. [3a](#page-10-0), Fig. [4](#page-11-0)a,b,c) and after interaction with wine samples, using the intensity of peaks a and b. 397 398 399 400 401 402 403 404 405

The initial fluorescence intensities for fibrinogen [(FI, Arbitral Units (A.U.) and maximum wavelength (λem/ex) nm] were the following: peak a with $FI = 861.1 \pm 10.4$ and λ em/ex = 229/342. Peak b was estimated as FI = 809.7 \pm 10.3 and λ em/ex = 282/341 (Figs. 3a and 4a). The initial fluorescence intensities for globulin [(FI, Arbitral Units (A.U.) and maximum wavelength (λem/ex) nm] were the following: peak a with $FI = 457.3 + 9.3311$ and λ em/ex = 231/335. Peak b was estimated as $FI = 661.1 \pm 10.3$ and λ em/ $ex = 280/334$ (Figs. 3a and 4b). The initial fluorescence intensities for HSA [(FI, Arbitral Units (A.U.) and maximum wavelength (λem/ex) nm] were the following: peak a with $FI = 643.0 \pm 7.9$ and λ em/ex = 228/353. Peak b was estimated as $FI = 920.1 \pm 10.4$ and λ em/ex = 280/357 (Figs. [3a](#page-10-0) and 4c). The measured data were changing during the interaction of these proteins with wines, tannic acid, quercetin, and ethanol (Figs. 3, 4, 5). It was determined a decrease in 406 407 408 409 410 411 412 413 414 415 416 417 418 419 420 421 422

Fig. 3 Change in the values of **a** fibrinogen/globulin/human serum albumin before the interaction with initial values of fluorescence intensities and after interaction with **b** quercetin, **c** tannic acid, **d** ethanol, **e** persimmon, **f** kiwi, **g** pomegranate wines. Values are means \pm SD of 5 measurements; n=5 samples, each subsampled and analyzed 5 times. Means within bars with the different superscripts

are statistically different $(p<0.05$; Student's t test). Abbreviations: fluorescence; intensity (FI); Arbitral Units (A.U.); fibrinogen (FB); human serum globulin (GL); human serum albumin (HSA), Binding properties (BP, %): binding to FB, to GL and HSA is the % decrease of fluorescence emission intensity of the fractions of the binding sites of the proteins occupied by the ligand to the initial ones

Fig. 4 Fluorometric measurements in a three-dimensional fluorescence analysis (3D-FL) of **a** fibrinogen, **b** globulin, **c** human serum albumin and tannin acid after interaction with **d** fibrinogen, **e** globulin, **f** human serum albumin. The locations of peaks **a** and **b** are shown in this figure: for fibrinogen with fluorescence intensity of Arbitral Units $(A.U.)$: peak $a=861.1 \pm 10.4$, peak

339 **b**=809.7±10.3; for globulin: peak **a**=457.3±9.3, peak **b**=661.1 \pm 10.3; and for HSA: peak 340 $a=643.0 \pm 7.9$; peak **and the locations of peaks a** and **b** for tannic acid are shown in this (**c**) (for interpretation of the references to color in this figure legend, the reader is referred to the web version of this article)

(Figs. 1b, 3f and 5d, e, f). The total binding properties (%) of pomegranate wines for fibrinogen, globulin, and HSA of peaks a and b were estimated as 78.2 ± 6.3 , 64.8 ± 5.8 , and 47.0 ± 3.9 , respectively (Figs. 1c, 3g, and 5g, h, i). The interactions of quercetin with fibrinogen, globulin, and HSA showed the following total binding properties (%) of peaks a and b such as 20.7 ± 1.1 , 44.6 ± 3.5 , and 15.1 ± 0.8 , respectively (Fig. 3f). The total binding properties (%) of ethanol for fibrinogen, globulin, and HSA of peaks a and b were calculated as 2.9 ± 0.3 , 2.6 ± 0.5 , and 2.4 ± 0.3 , respectively (Fig. [3](#page-10-0)g). All the presented results after the interaction of the main serum proteins were connected to the amount of polyphenols, anthocyanins, tannic acid, and antioxidant activities of the samples. The highest total binding properties were estimated for pomegranate, followed by kiwifruit and persimmon wines (Fig. [1](#page-7-0)). The results of bioactive compounds and fluorescence quenching show that these wines possess multiple properties that have a great potential to be used for human health and show similar data as the used fruits [[8,](#page-15-2)

the fluorescence intensities and a change in the maximum wavelengths during the reaction of fibrinogen, globulin, and HSA with bioactive substances of persimmon, kiwifruit, and pomegranate wines. The total binding properties (%) of quercetin for fibrinogen, globulin, and HSA of peaks a and b were estimated as 20.7 ± 1.1 , 44.6 ± 3.5 , and 15.1 ± 0.8 , respectively (Fig. $3b$). The total binding properties (%) of tannic acid for fibrinogen, globulin and HSA of peaks a and b were estimated as 54.3 ± 4.3 , 42.5 ± 3.8 , and 20.9 ± 1.7 , respectively (Fig. [3c](#page-10-0) and Fi[g.4d](#page-11-0)–f). The total binding properties (%) of ethanol for fibrinogen, globulin, and HSA of peaks a and b were calculated as 2.9 ± 0.3 , 2.6 ± 0.5 , and 2.4 ± 0.3 2.4 ± 0.3 , respectively (Fig. 3d). The total binding properties (%) of persimmon wines for fibrinogen, globulin, and HSA of peaks a and b showed the following data: 59.6 ± 5.7 , 21.6 ± 2.1 21.6 ± 2.1 21.6 ± 2.1 , and 28.8 ± 2.7 , respectively (Figs. 1a, [3e](#page-10-0), and $5a,b,c$ $5a,b,c$). The total binding properties $(\%)$ of kiwifruit wines for fibrinogen, globulin, and HSA of peaks a and b were calculated as 41.7 ± 3.2 , 34.4 ± 4.6 , and 35.7 ± 2.9 , respectively

$\circled{2}$ Springer

Downloaded from mostwiedzy.pl Downloaded fro[m mostwiedzy.pl](http://mostwiedzy.pl)**MOST WIEDZY**

Fig. 5 Fluorometric measurements in a three-dimensional fluorescence analysis (3D-FL) of wines: **a**–**c** persimmon, **d**–**f** kiwi fruit, **g**–**i** pomegranate after interaction with (**a**, **d**, **g**) fibrinogen, (**b**, **e**, **h**)

grape wines and 1.7–3.7 365 mM TE in white wines from

[10](#page-15-4)]. The beneficial effect of fruit- and vegetable-rich diets on cardiovascular health is partly attributed to the effect of their bioactive compounds [\[46](#page-16-6)] and bioactive-rich extracts of kale and pomegranate that are consumed as traditional plant foods of Black Sea area countries were effective in modulating platelet function. Comparison of the present results with the reported ones [[10,](#page-15-4) [20](#page-15-11), [47](#page-16-7), [48\]](#page-16-8) showed the following antioxidant activities: 9.6–29.9 mM TE in red France; 9.2–19.5 mM TE in red wines, and 0.5–1.4 mM TE in white wines from South Africa. The present results of antioxidant activities of 19.4–28.3 367 mM TE by CUPRAC assay showed that the fruit wines had relatively high antioxidant capacities that give a strong possibility of their use as a promising source of phenolic antioxidants. In vitro and in silico interactions of red grape wine polyphenols with human serum albumin, fibrinogen, glutathione peroxidase

pomegranate wines in Fig. 4e, f, and g**,** respectively

3 and C-reactive protein enhance their biological activity

globulin, (**c**, **f**, **i**) human serum albumin, respectively. The locations of peaks **a** and **b** are shown in this figure and for persimmon, kiwi, and

ding to HSA in promegrania wine was shipling have the mail of the luntencence, means the state of the properties of period in HSA in the main carrier in burnan metabolism for dyoublin, HSA thouscone data lod to the success [\[20](#page-15-11)]. A comparison of the binding properties of ethanol and bioactive substances in fruit wines once more showed the natural oxidative properties of polyphenols and their health properties. This can be explained only in the presence of bioactive substances in wines (the total binding properties of wines with fibrinogen ranged from 78.2 to 41.7% in comparison with ethanol of 2.9% [\[48\]](#page-16-8). The highest binding values were with fibrinogen, which is a very important protein and one of the indices of coronary artery disease [[10,](#page-15-4) 49]. The binding to HSA in pomegranate wine was slightly lower than with fibrinogen and globulin, and only in kiwifruit and persimmon wines was slightly higher than for globulin. HSA [\[20,](#page-15-11) [50](#page-16-9)] is the main carrier in human metabolism for drugs, such as antibiotics and a big number of drugs. The binding of HSA resulted in the fluorescence quenching of HSA, as it is in the presented results. Our study for the first time unveils the differential binding properties of kiwifruit and persimmon phytoconstituents with HSA. Although cultivars possess virtually the same amount, the presence of one unique compound significantly alters the binding properties of HSA. The results of fluorescence quenching and molecular docking showed that these fruits possess multiple properties, which have a great potential to be used as functional foods [[7\]](#page-15-22). Synergism was shown in the obtained measurements of quercetin, where the binding was lower than in used wine samples. Oppositely tannic acid showed high results of quenching in comparison with wines, especially persimmon wine which has a high amount of tannins. The binding properties of tannic acid with fibrinogen were higher than in kiwifruit wine but lower than in pomegranate and persimmon wines, which showed equal values for fibrinogen. The interaction between proteins and tannins was strong, as expected, leading to the precipitation of protein-tannin complexes. These results were in agreement with previously published data showing that high levels of precipitation of BSA by tannin can occur at low pH when the tannin to protein ratio is high $[51, 52]$. 479 480 481 482 483 484 485 486 487 488 489 490 491 492 493 494 495 496 497 498 499 500 501 502 503 504 505 506 507 508 509 510 511 512 513 514 515

Relationship between antioxidant and binding properties of fruit wines

To assess the relationship between antioxidant and binding properties of fruit wines methods of multivariate statistics were applied. Principal component analysis (PCA) brings clear differentiation of eigenvectors according to the wine type (Fig. $6a$). As regards the numerical values, the first two principal components (PC) cumulatively explained more than 91% of the whole dataset variability, with the dominant role of total binding to HSA, CUPRAC, and polyphenols (first $PC_$, $>$ 34% all), and vitamin C, tannins, and total binding to FB (second PC, $>69\%$, $>56\%$ and $>30\%$, respectively). Results of PCA also confirm principal component factoring (PCF) and the stepwise discriminant analysis

 $\circled{2}$ Springer

(CDA, data not presented) resulting in 100% correct classification of fruit wines. The plot of factors (varimax rotation, Fig. [6](#page-14-3)b) shows mutual strong positive correlations among binding properties to main serum proteins and polyphenols, anthocyanins, quercetin, CUPRAC, and DPPH. On the other hand, weak and moderate correlations between binding properties, vitamin C, and tannins are obvious from Fig. [6b](#page-14-3), which correspond with Pearson's correlation coefficients. 530 531 532 533 534 535 536 537

Methods of multivariate statistics were utilized also to process the results of the fluorescence measurements described above. Principal component analysis based on fluorescence data led to the successful differentiation of fruit wines and selected standards (Fig. 6c). First three PCs cumulatively explained more than 81% of the variability of the experimental characteristics, recognizing the variables binding properties and fluorescence intensities of peaks a and b for fibrinogen as the most important for PC1 construction, whereas for the second PC, fluorescence intensity and binding properties of peak a for GL and maximum wavelengths ($\lambda_{\text{em}/\text{ex}}$) of peaks a and b for HSA, and for the third PC, maximum wavelengths ($\lambda_{em/ex}$) of peak b for GL and FB, were identified as parameters with the highest eigenvalues. Similarly to PCA, the stepwise discriminant analysis resulted in 100% correct classification of samples (Fig. [6](#page-14-3)d). By means of the first discriminant function $(DF) > 69.5\%$ of the cases were correctly classified, whereas by the first and second DF > 97.4% of the cases and by the first three DFs, 100% of the cases were classified. As the most discriminating characteristics, in the first DF, fluorescence intensities of peaks a and b for FB were identified. In the second DF, fluorescence intensities of peak b for FB and peak a for GL, and in the third DF, fluorescence intensities of peaks a and b for FB reached the highest discriminant coefficients. 538 539 540 541 542 543 544 545 546 547 548 549 550 551 552 553 554 555 556 557 558 559 560 561 562

Conclusions

In this study, three-dimensional fluorescence spectroscopy in combination with FTIR was used in the investigation of antioxidant profiles in fruit wines. This is the first report showing differences and similarities in fruit wines, using their binding properties. The fluorescence spectral methods, which were applied as a powerful tool showing the quenching properties of intrinsic fluorophores in protein molecules in the presence of fruit wine polyphenols, can contribute to the interaction with drugs. Based on the quenching properties of human serum proteins with wines and recent reports in vivo on human studies, we hypothesize that the used human proteins can be predictors of coronary artery disease (CAD). The applied analytical methods are universal not only for the authentication of fruit wines, but also for a variety of fruits and vegetables. The application of FTIR measurements can be used as a 564 565 566 567 568 569 570 571 572 573 574 575 576 577 578 579

563

 $\overline{\circ}$

Fig. 6 Relationships between antioxidant, spectral, and binding properties of fruit wines. **a,** Principal component analysis of fruit wines based on antioxidant and binding properties of fruit wines. **b,** Plot of factors (varimax rotation) indicating mutual correlation and the importance of antioxidant and binding properties for the fruit wines

fingerprint showing the real composition of the products. The fluorescence spectra, resulting in the binding properties during interaction of different phenolic compounds with human serum proteins, can be used as a fast and reliable in vitro analysis for health-promoting benefits of the food products.

Acknowledgements Thanks from all authors to Dr. Elena Katrich for her assistance in the measuring of some indices in wines. Thanks to Rachel Aviv from the neighborhood wine shop, "Alciolim", Tagore 32, Tel Aviv, Israel, for collecting samples of wines.

Author contributions SG; YMK; ML-S; MP; BT: conceptualization; YMK; YSP; KSH: statistical evaluation; YKP; SGK: data curation; KSH; YKP; SGK: formal analysis; MP; BT: investigation; DB; AN; SG; ML-S: methodology; YSP; KSH: software; ML-S; PT; YSP; DB; AN: validation; SG: supervision; YMK; ML-S; SG; DB; AN; MP; BT: writing—original draft preparation, YMK; SG; ML-S; PT: writing review and editing. All authors have read and agreed to the published version of the manuscript.

differentiation. **c,** Principal component analysis of fruit wines and selected standards based on fluorescence data. **d,** Stepwise discriminant analysis of fruit wines and selected standards based on fluorescence data

Declarations

Conflict of interest The authors declare that they have no conflict of **interests** 599 600

References

- 1. Dembitsky VM, Poovarodom S, Leontowicz H, Leontowicz M, Vearasilp S, Trakhtenberg S, Gorinstein S (2011) The multiple nutrition properties of some exotic fruits: biological activity and active metabolites. Food Res Intern 44:1671–1701 602 603 604 605
- 2. Pasko P, Galanty A, Zagrodzki P, Luksirikul P, Barasch D, Nemirovski A, Gorinstein S (2021) Dragon fruits as a reservoir of natural polyphenolics with chemopreventive properties. Molecules 26:2158 606 607 608 609
- 3. Gorinstein S, Kulasek GW, Bartnikowska E, Leontowicz M, Zemser M, Morawiec M, Trakhtenberg S (2000) The effects of diets, supplemented with either whole persimmon or phenol-free persimmon, on rats fed cholesterol. Food Chem 70(3):303–308 610 611 612 613

 \hat{Z} Springer

598

601

AQ3 585

- 4. Gorinstein S, Leontowicz H, Leontowicz M, Jesion I, Namiesnik J, Drzewiecki J, Park Y-S, Ham K-S, Giordani E, Trakhtenberg S (2011) Influence of two cultivars of persimmon on atherosclerosis indices in rats fed cholesterol-containing diets: Investigation in *vitro* and in *vivo*. Nutrition 27(7–8):838–846 614 615 616 617 618
- 5. Zou B, Li C-M, Chen J-Y, Dong X-Q, Zhang Y, Du J (2012) High molecular weight persimmon tannin is a potent hypolipidemic in high-cholesterol diet fed rats. Food Res Intern 48:970–977 619 620 621 622
- 6. Drzewiecki J, Latocha P, Leontowicz H, Leontowicz N, Park YS, Najman K, Weisz M, Ezra A, Gorinstein S (2016) Analytical methods applied to characterization of *Actinidia arguta, Actinidia deliciosa*, and *Actinidia eriantha* kiwi fruit cultivars. Food Anal Methods 9:1353–1366 623 624 625 626 627
- 7. Kim YM, Park YS, Park YK, Ham KS, Kang SG, Shafreen RMB, Lakshmi SA, Gorinstein S (2020) Characterization of bioactive ligands with antioxidant properties of kiwifruit and persimmon cultivars using *in vitro* and *in silico* studies. Appl Sci 10:4218 628 629 630 631
- 8. Kim Y-M, Abas F, Park Y-S, Park Y-K, Ham K-S, Kang S-G, Lubinska-Szczygeł M, Ezra A, Gorinstein S (2021) Bioactivities of phenolic compounds from kiwifruit and persimmon. Molecules 26:4405 632 633 634 635
- 9. Çam M, Hısıl Y, Durmaz G (2009) Classification of eight pomegranate juices based on antioxidant capacity measured by four methods. Food Chem 112:721–726 636 637 638
- CONDITIONAL TRIVIAL IN THE CONDITIONAL TRIVIAL IN THE CONDITIONAL TRIVIAL IN THE CONDITIONAL TRIVIAL IN THE CONDITION CONDITIONAL TRIVIAL IN THE CONDITION CONTINUES IN THE CONDITION CONTINUES IN THE CONDITION CONTINUES IN 10. Aviram M, Dornfeld L, Rosenblat M, Volkova N, Kaplan M, Coleman R, Hayek T, Presser D, Fuhrman B (2000) Pomegranate juice consumption reduces oxidative stress, atherogenic modifications to LDL, and platelet aggregation: studies in humans and in atherosclerotic apolipoprotein E-deficient mice. Am J Clin Nutr 71:1062–1076 639 640 641 642 643 644
- 11. Chakraborty K, Saha J, Raychaudhuri U, Chakraborty R (2014) Tropical fruit wines: a mini review. Nat Products NPAIJ 10(7):219–228 645 646 647
	- 12. Towantakavanit K, Park Y-S, Gorinstein S (2011) Quality properties of wine from Korean kiwifruit new cultivars. Food Res Intern 44:1364–1372
	- 13. Gorinstein S, Moshe R, Weisz M, Hilevitz J, Tilis K, Feintuch D, Bavli D, Amram D (1993) Characteristics of persimmon liqueur. Food Chem 46:183–188
	- 14. Akalın AC, Bayram M, Anlı RE (2018) Antioxidant phenolic compounds of pomegranate wines produced by different maceration methods. J Inst Brew 124:38–44
	- 15. Paśko P, Tyszka-Czochara M, Namieśnik J, Jastrzębski Z, Leontowicz H, Drzewiecki J, Martinez-Ayala AL, Nemirovski A, Barasch D, Gorinstein S (2019) Cytotoxic, antioxidant and binding properties of polyphenols from the selected gluten-free pseudocereals and their by-products: *In vitro* model. J Cer Sci 87:325–333
	- 16. Ku YG, Kim HC, Bae JH, Kang BS, Nemirovski A, Barasch D, Gorinstein S (2019) Antioxidant capacities and polyphenols in autumn-growing cultivar of Chinese cabbage (*Brassica rapa* L. ssp. pekinensis cv Bulam Plus). Europ Food Res Tech 245:1871–1879
	- 17. Shafreen RMB, Lakshmi SA, Pandian SK, Park YS, Kim YM, Pásko P, Deutsch J, Katrich E, Gorinstein S (2020) Unraveling the antioxidant, binding and health-protecting properties of phenolic compounds of beers with main human serum proteins: in *vitro* and *in silico* approaches. Molecules 25:4962
	- 18. Singleton VL, Orthofer R, Lamuela-Raventós RM (1999) Analysis of total phenols and other oxidation substrates and antioxidants by means of Folin-Ciocalteu reagent. Methods Enzym 299:152–178
	- 19. Lee J, Durst RW, Wrolstad ER, Eisele T, Giusti MM, Hofsommer H, Koswig S, Krueger AD, Kupina S, Martin SK, Martinsen BK, Miller TC, Paquette F, Ryabkova A, Skrede G, Trenn U, Wightman JD (2005) Determination of total monomeric anthocyanin pigment content of fruit juices, beverages, natural colorants, and

wines by the pH differential method: collaborative study. J AOAC Int 88:1269–1278

- 20. Shafreen RMB, Lakshmi SA, Pandian SK, Kim Y-M, Deutsch J, Katrich E, Gorinstein S (2021) *In vitro* and *in silico* interaction studies with red wine polyphenols against different proteins from human serum. Molecules 26:6686
- 21. Broadhurst RB, Jones WT (1978) Analysis of condensed tannins using acidified vanillin. J Sci Food Agric 29:788–794
- 22. Lamuela-Raventos RM, Waterhouse AL (1994) A direct HPLC separation of wine phenolics. Am J Enol Vitic 45:1–5
- 23. Özyürek M, Güçlü K, Bektas Oğlu B, Apak R (2007) Spectrophotometric determination of ascorbic acid by the modified CUPRAC method with extractive separation of flavonoids-La(III) complexes. Anal Chim Acta 588:88–95
- 24. Apak R, Güçlü K, Özyürek M, Karademir SE (2004) Novel total antioxidant capacity index for dietary polyphenols and vitamins C and E, using their cupric ion reducing capability in the presence of neocuproine: CUPRAC method. J Agric Food Chem 52:7970–7981
- 25. Brand-Williams W, Cuvelier M, Berset C (1995) Use of a free radical method to evaluate antioxidant activity. LWT 28:25–30
- 26. Park Y-S, Im MH, Ham K-S, Kang S-G, Park Y-K, Namiesnik J, Leontowicz H, Leontowicz M, Trakhtenberg S, Gorinstein S (2015) Quantitative assessment of the main antioxidant compounds, antioxidant activities and FTIR spectra from commonly consumed fruits, compared to standard kiwi fruit. LWT Food Sci Technol 63:346–352
- 27. Mena P, Gironés-Vilaplana A, Martí N, García-Viguera C (2012) Pomegranate varietal wines: phytochemical composition and quality parameters. Food Chem 133:108–115
- 28. Di Stefano V, Pitonzo R, Novara ME, Bongiorno D, Indelicato S, Gentile C, Avellone G, Bognanni R, Scandurra S, Melilli MG (2019) Antioxidant activity and phenolic composition in pomegranate (*Punica granatum* L.) genotypes from South Italy by UHPLC-Orbitrap-MS approach. J Sci Food Agric 99(3):1038–1045
- 29. Gil MI, Tomas-Barberan FA, Hess Pierce B, Holcroft DM, Kader AA (2000) Antioxidant activity of pomegranate juice and its relationship with phenolic composition and processing. J Agric Food Chem 48:4581–4589
- 30. Denev P, Yordanov A (2013) Total polyphenol, proanthocyanidin and flavonoid content, carbohydrate composition and antioxidant activity of persimmon (*Diospyros kaki* L.) fruit in relation to cultivar and maturity stage. Bulg J Agric Sci 19(5):981–988
- 31. Jiménez-Sánchez C, Lozano-Sánchez J, Marti N, Saura D, Valero M, Segura-Carretero A, Fernández-Gutiérrez A (2015) Characterization of polyphenols, sugars, and other polar compounds in persimmon juices produced under different technologies and their assessment in terms of compositional variations. Analytical methods. Food Chem 182:282–291
- 32. Yaqub S, Farooq U, Shafi A, Akram K, Murtaza MA, Kausar T, Siddique F (2016) Chemistry and functionality of bioactive compounds present in persimmon. J Chem. [https://doi.org/10.1155/](https://doi.org/10.1155/2016/3424025) 2016/3424025
- 33. Pérez-Burillo S, Oliveras MJ, Quesada J, Rufián-Henares JA, Pastoriza S (2018) Relationship between composition and bioactivity of persimmon and kiwifruit. Food Res Intern 105:461–472
- 34. Zou B, Wu J, Yu Y, Xiao G, Xu Y (2017) Evolution of the antioxidant capacity and phenolic contents of persimmon during fermentation. Food Sci Biotechnol 26(3):563–571
- 35. Sokolletowska A, Kucharska AZ, Winska K, Szumny A, Nawirskaolszanska A, Mizgier P, Wyspianska D (2014) Composition and antioxidant activity of red fruit liqueurs. Food Chem 157:533–539
- 36. Suh JH, Virsolvy A, Goux A, Cassan C, Richard S, Cristol JP, Teissèdre PL, Rouanet JM (2011) Polyphenols prevent lipid 743 744 745

 \mathcal{D} Springer

 $\overline{\circ}$

- abnormalities and arterial dysfunction in hamsters on a highfat diet: a comparative study of red grape and white persimmon wines. Food Funct 2:555–561 746 747 748
- 37. Recio-Rodriguez JI, Gomez-Marcos MA, Patino-Alonso MC, Puigdomenech E, Notario-Pacheco B, Mendizabal-Gallestegui N, de la de la CalFuente A, Otegui-Ilarduya L, Maderuelo-Fernandez JA, de AngelaCaboLaso A, Agudo-Conde C, Garcia-Ortiz L, On behalf of the EVIDENT group (2015) Effects of kiwi consumption on plasma lipids, fibrinogen and insulin resistance in the context 749 750 751 752 753 754
- of a normal diet. Nutr J 14:97 38. Luo A, Liu X, Ren Y, Kou L (2004) Study on brewing technology 755 756
- of kiwi-fruit dry wine. J Chin Inst Food Sci Tech 4:5–11 39. Ma T, Lan T, Ju Y, Cheng G, Que Z, Geng T, Fang Y, Sun X (2019) Comparison of the nutritional properties and biological activities of kiwifruit (*Actinidia*) and their different forms of products: towards making kiwifruit more nutritious and functional. Food Funct 10:1317–1329 757 758 759 760 761 762
- 40. MFDS (2017) Food additives code. Ministry of Food and Drug Safety. Ministry of Agriculture Food and Rural Affairs 763 764
- 41. Cho YS, Kim JJ, Jeon G, Chung M-S, Joo Y, Lee K-W (2021) Total SO₂ levels and risk assessment of wine and fruit wine consumed in South Korea. Food Contr 127:108124 765 766 767
- 42. dos Santos Grasel F, Ferrão MF, Wolf CR (2016) Development of methodology for identification the nature of the polyphenolic extracts by FTIR associated with multivariate analysis. Spectrochim Acta A Mol Biomol Spectrosc 153:94–101 768 769 770 771
- 43. Patle TK, Shrivas K, Kurrey R, Upadhyay S, Jangde R, Chauhan R (2020) Phytochemical screening and determination of phenolics and flavonoids in *Dillenia pentagyna* using UV–vis and FTIR spectroscopy. Spectrochim Acta A Mol Biomol Spectrosc 242:2020118717 772 773 774 775 776
- 44. Mayra A, Castro P, Rodríguez HG (2011) Study by infrared spectroscopy and thermogravimetric analysis of tannins and tannic acid. Rev Latinoam de Quimica 39:107–112 777 778 779
	- 45. Ricci A, Olejar KJ, Parpinello GP, Kilmartin PA, Versari A (2015) Application of Fourier transform infrared (FTIR) spectroscopy in the characterization of tannins. Appl Spectrosc Rev 50(5):407–442

Authors and Affiliations

- 46. Konić-Ristić A, Srdić-Rajić T, Kardum N, Aleksić-Velićković V, Kroon PA, Hollands WJ, Needs PW, Boyko N, Hayran O, Jorjadze M, Glibetić M (2013) Effects of bioactive-rich extracts of pomegranate, persimmon, nettle, dill, kale and *Sideritis* and isolated bioactives on arachidonic acid induced markers of platelet activation and aggregation. J Sci Food Agric 93:3581–3587 784 785 786 787 788 789
- 47. Beer D, Joubert E, Gelderblom WCA, Manley M (2003) Antioxidant activity of South African red and white cultivar wines: free radical scavenging. J Agric Food Chem 51:902–909

799

801 802 803

- 48. Sezer ED, Akçay YD, Ilanbey B, Yildirim HK, Sözmen EY (2007) Pomegranate wine has greater protection capacity than red wine on low-density lipoprotein oxidation. J Med Food 10(2):371–374
- 49. Yoshikawa H, Hirano A, Arakawa T, Shiraki K (2012) Effects of alcohol on the stability and structure of native and disulfide-modified bovine serum albumin. Int J Biol Macromol 50:1286–1291 795 796 797 798
- 50. Zhang Y, Cao Y, Li Y, Zhang X (2022) Interactions between human serum albumin and sulfadimethoxine determined using spectroscopy and molecular docking. Molecules 27:1526 800
- 51. Adamczyk B, Salminen JP, Smolander A, Kitunen V (2012) Precipitation of proteins by tannins: effects of concentration, protein/ tannin ratio and pH. Intern J Food Sci Tech 47(4):875–878
- 52. Mierczynska-Vasilev A, Bindon K, Gawel R, Smith P, Vasilev K, Butt H-J, Koynov K (2021) Fluorescence correlation spectroscopy to unravel the interactions between macromolecules in wine. Food Chem 352:129343 804 805 806 807 808

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations. 809 810

Springer Nature or its licensor (e.g. a society or other partner) holds exclusive rights to this article under a publishing agreement with the author(s) or other rightsholder(s); author self-archiving of the accepted manuscript version of this article is solely governed by the terms of such publishing agreement and applicable law. 811 812 813 814 815

Young Mo Kim1 · Martyna Lubinska‑Szczygeł² · Martin Polovka3 · Blanka Tobolkova3 · Pitipong Thobunluepop⁴ · Yong Seo Park5 · Kyung Sik Ham6 · Yang Kyun Park6 · Seong Gook Kang⁶ · Dinorah Barasch7 · Alina Nemirovski7 · Shela Gorinstein7

- ⁴ Department of Agronomy, Faculty of Agriculture, Kasetsart University, Chatuchak, Bangkok 10900, Thailand
- ⁵ Department of Horticultural Science, Mokpo National University, Muan 534-729, Jeonnam, Korea
- ⁶ Department of Food Engineering, Mokpo National University, Muan 534-729, Jeonnam, Korea
- ⁷ School of Pharmacy, Faculty of Medicine, Institute for Drug Research, The Hebrew University of Jerusalem, 14, 9112001 Jerusalem, Israel

UNCORRECTED

² Springer

Author Query Form

Please ensure you fill out your response to the queries raised below and return this form along with your corrections

Dear Author

During the process of typesetting your article, the following queries have arisen. Please check your typeset proof carefully against the queries listed below and mark the necessary changes either directly on the proof/online grid or in the 'Author's response' area provided below

