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1	Inhibitory effect of chlorogenic acid on digestion of potato
2	starch
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22 Highlights:

Chlorogenic acid inhibits the digestion of isolated potato starch both competitively and
 non-competitively.

Five commercial potato varieties were analysed for phenolic content and starch
digestibility.

Digestibility is affected by multiple factors including phenolic, dry matter and starch
content.

29 Abstract:

30 The effect of the chlorogenic acid isomer 5-O-caffeoylquinic acid (5-COA) on digestion of 31 potato starch by porcine pancreatic alpha amylase (PPAA) was investigated using isolated 32 starch and cooked potato tuber as substrates. In vitro digestion was performed on five varieties of potato with varying phenolic content. Co- and pre-incubation of PPAA with 5-CQA 33 34 significantly reduced PPAA activity in a dose dependent manner with an IC50 value of about 2 mg mL⁻¹. Lineweaver-Burk plots indicated that 5-CQA exerts a mixed type inhibition as k_m 35 36 increased and V_{max} decreased. The total polyphenol content (TPC) of peeled tuber tissue ranged from 320.59 to 528.94 mg 100g⁻¹ dry weight (DW) in raw tubers and 282.03 to 543.96 mg 37 100g⁻¹ DW in cooked tubers. With the exception of Désirée, TPC and 5-CQA levels decreased 38 39 after cooking. Principle component analysis indicated that digestibility is affected by multiple 40 factors including phenolic, dry matter and starch content.

41

42 Keywords

43 Alpha amylase; starch; chlorogenic acid; enzyme kinetics; polyphenol; digestion; potato;
44 Solanum tuberosum.

46 **1. Introduction**

47 Potato is the third most consumed food crop providing around 5% to 15% of dietary energy, 48 primarily from starch, to various populations around the world. Potato has been labelled with 49 a high glycaemic index (GI) because consumption of some potato products can cause a sharp 50 increase of postprandial blood glucose concentration. However there is significant variation in 51 the GI of various potato products (Weichselbaum, 2010). Many investigators have reported the 52 effect of food processing (industrial or domestic) on the GI of potato (Ek, Brand-Miller and 53 Copeland, 2012). For example, it has been reported that freshly boiled potato (GI 78) and 54 instant mashed potato (GI 87) have a higher GI than French fries (GI 63) (Atkinson, Foster-55 Powell and Brand-Miller, 2008). Furthermore, the incremental area under curve (AUC) of 56 freshly cooked potato decreased by 18% and 30% upon storage in the refrigerator for 1 and 5 57 days respectively. This observation is attributed to the retrogradation of amylose which reduces 58 digestibility (Fernandes, Velangi, and Wolever, 2005). Henry, Lightowler, Strik and Storey 59 (2005) determined the GI of commercially available potatoes in Great Britain and demonstrated 60 that varieties prepared by boiling for 15 min showed a wide variation in GI values, ranging 61 from 56 for Marfona to 94 for Maris Peer. Henry postulated that the difference was related to 62 their texture, with waxy potatoes having a medium GI and floury potatoes having a high GI. 63 We postulate that some of these differences may be attributed to the presence of endogenous 64 polyphenolic substances acting as α -amylase inhibitors in the digestive tract.

Pancreatic alpha amylase is an endoglycosidase enzyme that has a significant role in carbohydrate digestion. It has been shown that inhibitors of α -amylase reduce bioavailability of glucose (Bozzetto et al., 2015). Controlling blood glucose level by α -amylase inhibitors may play a role in preventing hyperglycaemia in patients with diabetes mellitus. Some α amylase inhibitors are naturally present in foods. Potato tubers contain many plant secondary metabolites including phenolics, carotenoids and polyamines. Their content and composition

71 vary according to the variety of potato, conditions of cultivation, cooking and processing 72 methods (Ezekiel, Singh, Sharma and Kaur, 2013, Ryes and Cisneros-Zevallos, 2003). 73 Phenolics present in potatoes include phenolic acids, tannins, lignin, flavonoids, coumarins and 74 anthocyanins (Ryes and Cisneros-Zevallos, 2003). 5-O-caffeoylquinic acid (5-CQA) is an 75 isomer of chlorogenic acid (CGA) which makes up 90% of the phenolic content of potato 76 (Malenberg and Theander, 1985). The concentration of 5-CQA is higher in the skin than in the 77 medulla. Analysis has shown that cooked unpeeled potato contains between 9.1 to 12 mg 5-78 CQA per 100 g fresh weight compared to 0.86 to 6.6 mg for equivalent peeled samples (Mattila 79 and Hellstrom, 2007). The variation in 5-CQA content suggests that different potatoes may 80 inhibit pancreatic amylase to different extents.

81 Evidence from a number of in vitro and in vivo studies indicates inhibitory effects of 82 polyphenols on enzymes involved in carbohydrate digestion. It was reported that 5-CQA, 83 quinic acid (QA) and caffeic acid (CA) have mixed-type inhibitory effect against pure porcine 84 pancreatic alpha amylase (PPAA) isomers I and II using p-nitrophenyl-a-D-maltoside as a 85 substrate (Narita and Inouye, 2009). The most potent inhibitor was 5-CQA followed by CA and QA. In an in vivo animal study, oral intake of a 5-CQA solution (3.5 mg kg⁻¹ body weight) 86 87 during a glucose tolerance test lowered the height of glycaemia peaks at 10 and 15 min by 22 88 and 17% respectively compared to control (only glucose) (Bassoli et al., 2008). Rohn, Rawel 89 and Kroll (2002) derivatized PPAA with a number of phenolic compounds and showed that 5-90 CQA reacted covalently with the enzyme and decreased its activity by about 50%. However, 91 the mode of inhibition and potential effect on digestion of native potato starch has not been shown. The aim of the present study was to characterise the effect of 5-CQA on PPAA activity 92 93 in vitro using potato starch as a substrate and to determine the in vitro digestibility of steam 94 cooked potatoes from varieties which vary in their phenolic content.

95 **2. Materials and methods:**

96 **2.1 Potato samples**

97 Tubers from five commercial varieties that differ in skin colour were purchased from food 98 markets in Leeds, UK. Maris piper has creamy flesh and golden yellow skin; Maris peer has 99 creamy flesh and skin; Désirée, Rooster and Mozart have reddish pink skin and light yellow 100 flesh. Three potatoes from each variety were rinsed with water and dried with a paper towel. 101 Then the potatoes were peeled to remove skin and cortex, cut into cubes (1 cm³) and separated 102 into 100 g batches. Potatoes cubes were placed into a steam pan and cooked for 30 min at 103 boiling temperature. Cooked potatoes were mashed by using a fork and used for enzymatic 104 digestion. The remaining raw and steam cooked mashed potato was immediately frozen at -80 105 °C, freeze dried and stored at -80°C for analysis of TPC and phenolic acid composition. 106 Analyses were repeated at least three times with three batches of potato.

107 2.2 In vitro digestion of potato starch in presence of chlorogenic acid

108 The activity of PPAA (16 U/mg, Sigma Aldrich) enzyme on hydrolysis potato starch in the 109 presence and absence of 5-CQA (Sigma Aldrich; PubChem CID 12310830) was examined by 110 the method of Brenfeld (1955) and Kazeem, Adamson and Ogunwande (2013) with some 111 modifications. One percent (w/v) soluble potato starch (Sigma Aldrich) was suspended in 112 20mM sodium phosphate buffer pH 6.9 buffer containing 6.7mM NaCl and gelatinized for 15 113 min at 90 °C then allowed to cool to 37°C before addition of PPAA at a concentration of 0.33 114 U ml⁻¹. The reaction was followed at 37°C for up to 20 minutes. 5-CQA (final concentration 115 1.5 mg mL⁻¹) was either added to the enzyme-substrate mixture at the start of the reaction or 116 was pre-incubated for 10 minutes with the enzyme prior to addition of the substrate. All 117 reactions were carried out in four replicates. Reducing sugar released was measured at two reaction times (5 and 20 min) using the 3, 5-dinitrosalicylic acid (DNS; Sigma Aldrich) 118

119 colorimetric assay (Miller, 1959, Fei et al., 2014). An enzymatic kit was not used due the 120 inhibition of enzymes in the kit by chlorogenic acid. Brayer has shown that maltose is the 121 preferred leaving group for PPAA (Brayer et al., 2000), and therefore maltose was used for 122 generating standard curve to quantify the reducing sugar released. The enzymatic activity of 123 PPAA was determined in the presence of various concentrations of 5-CQA (0.08 - 2 mg mL⁻ 124 ¹). IC₅₀ was calculated as the concentration of 5-CQA required to inhibit 50% of enzyme 125 activity.

126 **2.3 Enzymatic kinetics and mode of inhibition**

Michaelis-Menten kinetic parameters and mode of inhibition of PPAA by 5-CQA was determined from a Lineweaver-Burk plot. One mL of 5-CQA at concentrations ranging from 0 to 2 mg mL⁻¹ was added to a mixture containing 1 mL of starch solution at concentrations from 0 to 6.6 mg ml⁻¹ in the same buffer solution as described in the previous section. The reaction was initiated by addition of a fixed concentration of PPAA (0.33 unit mL⁻¹). The solution mixture was incubated for 5 min at 37 °C. The reducing sugar produced was determined by the DNS colorimetric method as described previously.

134 **2.4 Determination of total starch content**

135 Total starch (TS) was determined enzymatically according to the method of Goñi, Garcia-136 Alonso and Saura -Calixto (1997) with some modifications. Raw potato (50 mg) was 137 homogenized in 6 mL of 2M KOH and then agitated using a shaking vortex at room 138 temperature for 30 min. The agitation step was very important to ensure complete solubility 139 of the starch. 3 mL of 0.4M sodium acetate buffer (pH=4.75) was added to the suspension and 140 pH was adjusted to 4.75 by using 3M acetic acid. 60 µl of amyloglucosidase (AMG) from 141 Aspergillus niger (70 U/mg, Sigma Aldrich) was added to the solubilized starch and hydrolysed 142 for 45 min at 60 °C in a shaking water bath. The digestion mixture was centrifuged for 5 min and pH was neutralised with 0.2M NaOH. Glucose in the supernatant was measured using theDNS method. Glucose amount was converted to starch by multiplying by a factor 0.9.

145 **2.5 Determination of free sugar content**

Free sugar content was determined in order to correct the TS value in potato samples. Potato samples, 200 mg of raw or cooked tuber, were homogenized in 6 mL sodium acetate buffer (pH=4.75) and then centrifuged for 10 min. Soluble sugars were determined using the DNS method and high performance anion exchange chromatography with pulsed amperometric detection.

151 2.6 High performance anion exchange chromatography with pulsed amperometric detection 152 (HPAEC-PAD)

153 Sugar solutions (glucose, fructose, sucrose, maltose and maltotriose, all from Sigma Aldrich) were used as standards at concentrations of 0-0.2 μ g mL⁻¹. Samples and standards were spiked 154 155 with internal standard (fucose, final concentration of 0.05 µg mL⁻¹). Samples were filtered 156 through PTFE membrane filters (0.2 µm pore size, Chromacol Ltd) and analyzed by HPAEC-157 PAD (Thermo Fisher DX500 instrument equipped with a GP40 gradient pump, ED40 158 electrochemical detector including gold working and silver reference electrodes and a LC20 159 column oven set at 30°C). The analytical column used was CarboPac PA20 (3×150mm) with 160 guard (3×30mm) with anion exchange capacities of 65µeq/column. The mobile phase was 200 161 mM NaOH and the flow rate was 0.4 mL/min. Injections (10 µL) were made by an AS500 162 autosampler. The elution programme was as follows: isocratic elution with 60mM NaOH from 163 0 to 8 min, followed by increasing gradient up to 140 mM NaOH to 17 min. The concentration 164 was reduced back to 60mM and equilibration was carried out for 6 min.

165 **2.7 Determination of moisture content**

Potato samples fresh and cooked (in triplicate) were weighed and frozen at -80°C then freeze
dried for 48 hours. The moisture content was calculated as percentage of weight loss. Dry
matter (DM) was calculated from the remaining dry yield.

169 **2.8 In vitro starch hydrolysis of steam cooked potato**

170 In order to test the hypothesis that the TPC affects the digestibility of starch in potato varieties,

171 in vitro digestibility of different verities of potato were carried out. Steam cooked tubers from

172 potato varieties (10 mg total starch mL⁻¹) with different TPC levels were digested using PPAA

173 $(0.66 \text{ unit mL}^{-1})$ for up to 180 min at 37°C in a shaking water bath (75 rpm).

In addition, we also performed sequential digestion with AMG (60 µl of AMG (184 U) from
Aspergillus niger, Sigma Aldrich) following PPAA digestion to degrade all digestion products
to glucose. Before AMG digestion, the pH was adjusted to 4.75 and digestion performed for
30 min at 37°C in a shaking water bath (75 rpm). Digestion products were detected using the

178 DNS method and HPAEC-PAD.

179 **2.9 Determination of total polyphenol content (TPC)**

180 The method of extraction was adapted from Shakya and Navarre (2006). Phenolic compounds 181 were extracted in four replicates taken from freeze dried raw and cooked potatoes. Freeze-dried 182 powder (200 mg) was mixed with 1.5 mL of extraction buffer (50% MeOH, 2.5% 183 metaphosphoric acid, 1 mM EDTA, chilled to 4°C) and 500 mg of glass beads (1.0 mm in 184 diameter; Fisher Scientific). Tubes were shaken with a vortex for 10 min at room temperature 185 and then sonicated at 10 °C for 10 min. After sonication, tubes were shaken again with a vortex 186 for 10 min. Tubes were centrifuged at 2500 g at 4°C for 10 minutes and the supernatant was 187 transferred to a clean tube. Extractions were repeated three times and supernatants combined. 188 Samples were kept chilled at all times and not exposed to light. TPC of potato was measured by Folin-Ciocalteau method (Singleton and Rossi, 1965). Potato extract (50 μ L) was mixed with Folin-Ciocalteau reagent (50 μ L, 1 N), sodium carbonate (150 μ L, 20% w/v) and distilled water (750 μ L). After shaking by vortex, the mixture was incubated in the dark at room temperature for 45 min. Absorbance was measured at 765 nm using a spectrophotometer (Cecil, CE 7200 Double Beam UV/VIS Spectrophotometer). Different concentrations of 5-CQA (10-300 μ g mL⁻¹) were used to generate a standard curve.

195 2.10 Separation of phenolic acids using high performance liquid chromatography with diode array detector – mass spectrometry (HPLC-DAD-MS).

197 The HPLC-DAD-MS system consisted of a micro vacuum degasser (Prominence Degasser LC-198 20 A5, Shimadzu), a liquid chromatograph (Prominence Liquid Chromatograph LC-30 AD, 199 Shimadzu), an auto sampler (Prominence Auto Sampler SIL-30 AC, Shimadzu), a diode array detector (Prominence Diode Array Detector system SPD-M20A, Shimadzu), a column oven 200 201 (Prominence Column Oven CTO-20 AC, Shimadzu), a controller (Prominence Controller 202 CBM-20 A, Shimadzu), and an MS detector with electrospray ion source and quadrupole 203 analyser (Liquid Chromatograph Mass Spectrometer LCMS-2020, Shimadzu). The Labs 204 solutions (Shimadzu) software was used to control the LC-MS system and for data processing. 205 The column used for chromatographic separation was an Agilent Zorbax Eclipse plus C18 206 column 4.6 mm \times 150 mm, 5 µm internal diameter. A gradient elution program of solvent A 207 (0.1 % formic acid, 5% acetonitrile and 94% water) and solvent B (0.1 % formic acid, 5% water 208 and 94% acetonitrile) was set as follows: 61-min; linear gradient from 0-51 min from 0% to 209 100% solvent B, isocratic elution from 51.1-56 min with 100% solvent B, linear gradient from 210 56-56.1 min to 0% solvent B and isocratic elution from 56.1-61 min with 0% solvent B. The column temperature was 35°C and flow rate of 0.5 mL min⁻¹ and injection volume 10 µL. The 211 212 diode array detection spectra was recorded at wavelengths of 280, 290, 315, 320 and 330 nm.

213 2.11 Statistical analysis

Statistical analysis was carried out using IBM SPSS statistics version 22 software program for
window. An independent sample t-test was used to compare amylase activity in presence or
absence of chlorogenic acid. One way ANOVA was used to analyse the differences between
varieties in terms of compositon and digestibility, and also to test the effect of cooking on
phenolic content. Differences were considered to be statistically significant when p≤0.05.
Pearson correlations were performed between composition factors and AUC values.

220 Principle component analysis (PCA) was conducted to analyse the relationship between starch

hydrolyzed and TS, DM, TPC and 5-CQA content of potatoes. PCA was performed using
MATLAB software R2015a (MathWorks, Inc.).

223

224 **3. Results and discussion**

225 **3.1 In vitro hydrolysis of potato starch in presence of chlorogenic acid**

226 We evaluated the inhibitory effect of the 5-CQA on PPAA activity using potato starch as 227 substrate at two incubation time periods of 5 and 20 min (figure 1A). The results indicate that 228 5-CQA significantly ($p \le 0.05$) inhibited PPAA at both incubation times, the inhibition was 229 more pronounced at 5 min (25.5% inhibition) compared to 20 min (1.5% inhibition). The 230 difference in inhibition at the two reaction times indicates that 5-CQA is most efficient during 231 early stages of hydrolysis, most probably in a competitive way. Pre-incubation of enzyme with 232 5-CQA for 10 min inhibited the enzyme by 53.8 and 28.3 % at 5 and 20 min respectively. In 233 order to calculate the IC₅₀, PPAA activity was measured in the presence of increasing 234 concentrations of 5-CQA (figure 1B). The IC₅₀ was found to be around 2 mg mL⁻¹ 5-CQA and 235 this in agreement with results of Sun et al. (2016) who reported IC_{50} of 1.96 mg mL⁻¹ for 236 chlorogenic acid against PPAA hydrolysis of maize starch. The enzymatic kinetic properties

237 were examined by applying Michaelis-Menten assumptions. The initial velocity of the reaction 238 (V) was determined at various concentrations [S] of gelatinized potato starch in the absence 239 and presence of various concentrations of the 5-CQA and a fixed concentration of PPAA. A 240 Lineweaver-Burk plot produced linear relationships for 1/V against 1/[S] at various 5-CQA 241 various concentrations (figure 1C). Increasing inhibitor concentrations led to an increase in 242 slope and y-intercept on the vertical axis. The intersection of lines in the second quadrant 243 suggests a mixed type of inhibition of 5-CQA against PPAA. Kinetic parameter V_{max} and k_m were obtained from Lineweaver-Burk plots (table 1); k_m values increased from 1.66 to 2.08 mg 244 245 mL^{-1} starch with increasing inhibitor concentrations while maximum velocity (V_{max}) values decreased from 0.35 to 0.29 mg mL⁻¹ min⁻¹ maltose. These results also suggest that 5-CQA has 246 247 a mixed-type inhibition (competitive and non-competitive) against PPAA when potato starch 248 is used as a substrate. This mixed type inhibition behaviour was previously reported for 5-CQA 249 using p-nitrophenyl-α-D-maltoside as a substrate (Narita and Inouye, 2009). Recently, Sun et 250 al (2016) reported that chlorogenic and caffeic acid have a mixed type inhibitory effect on 251 PPAA hydrolysis of maize starch. These two inhibitors decrease the V_{max} , by interacting more 252 strongly with the enzyme-substrate complex than with the free enzyme (Sun et al 2016, Narita 253 and Inouye, 2011). Rohn, Rawel and Kroll (2002) reported that the incubation of digestive 254 enzymes (PPAA, trypsin, lysozyme) with simple phenolic compounds (CQA, caffeic acid, 255 gallic acid and ferulic acid) for 24 hours resulted in covalent attachment of the phenolic 256 compounds to the free amino groups of the enzymes and in consequence, decreased their 257 activity irreversibly and non-competitively. The non-competitive inhibition of PPAA was 258 reported for millet seed coat extract using potato starch as substrate (Shobana, Sreerama and 259 Malleshi, 2009).

The inhibitory activity of phenolic acids is enhanced with increasing the number of phenolicsub-structures. The inhibitory effect of caffeic acid was enhanced 5-fold by combining with

quinic acid to form chlorogenic acids (Narita & Inouye, 2011). We show that 5-CQA is ableto inhibit PPAA activity on potato starch in both competitive and non-competitive manners.

264 **3.2 Total phenolic content**

The TPC of raw and cooked peeled potato samples is shown in table 2. TPC in fresh peeled potatoes varied from 320.59 mg 100 g⁻¹ dry weight (DW; 5-CQA equivalent) in Rooster to 528.94 mg 100 g⁻¹ DW in Maris Peer. TPC content was in the order Maris Peer > Mozart > Désirée > Maris Piper > Rooster from highest to lowest content. Variety affected TPC content significantly (p \leq 0.05). We did not measure TPC content in the skin of the potatoes, but observe that skin colour is not a good indicator of TPC content in the flesh.

In general, steam cooking affected the TPC in all potato samples. Cooking reduced the TPC in Maris Peer (-35%) and Mozart (-19%) significantly ($p \le 0.05$). Smaller non-significant decreases were observed in Maris Piper (-7%) and Rooster (-12%) (p = 0.26 and 0.20 respectively). However, the TPC content increased in Désirée potatoes by 22% (p=0.04). This variation in the response to cooking has been reported in the literature (Tian, Chen, Ye, Chen, 2016).

277 There are many reports of TPC in different genotypes of potato in peeled, unpeeled, fresh and 278 cooked forms. The TPC varies according to genotype, agricultural practices and method of 279 Ah-Hen, Fuenzalida, Contreras, Vega-Galvez and Lemus-Mondaca (2012) extraction. 280 examined the TPC in some native coloured potatoes from Chiloe Island (Southern Chile) and found that TPC in peeled potato samples varied from 192 to 1864 mg 100g⁻¹ DW (ferulic acid 281 282 equivalent). Lachman, Hamouz, Sulc, Orsak and Dvorak (2008) reported TPC in red and 283 yellow-fleshed potato to range from 296 mg 100 g⁻¹ DW in yellow-fleshed potato to 468 mg 100 g⁻¹ DW (gallic acid equivalent) in purple-fleshed potato. These levels are consistent with 284 285 the results obtained in this study. Recently, Tierno, Hornero-Mendez, Gallardo-Guerrero, 286 Lopez-Pardo and Ruiz de Galarreta (2015) reported that boiling peeled potato for 30 min 287 reduced the TPC of tubers by about 50% and similar results also were obtained by Lemos, 288 Aliyu and Hungerford (2015). Meanwhile, Blessington, Nzaramba, Scheuring, Hale, Reddivari 289 and Miller (2010) reported an increase in TPC after baking, frying and microwaving. Bembem 290 and Sadan (2013) and Burgos et al. (2013) observed an increase in TPC after boiling, steam 291 cooking, microwaving and pressure cooking. Therefore, an increase in level of TPC of Désirée 292 potato in our study after steam cooking is in accordance with some published reports. The 293 increased levels could be due to the release by cooking of phenolic or other compounds (Tian, 294 Chen, Ye, Chen, 2016).

295 One of the disadvantages of the TPC assay using Folin-Ciocalteau reagent is the low level of 296 specificity. For instance, the reagent reacts with ascorbic acid and tyrosine, both abundant in 297 potatoes. It has been reported that the ascorbic acid response to the Folin-Ciocalteau reagent is 298 two times higher than gallic acid (Lachman, Hamouz, Sulc, Orsak and Dvorak, 2008). 299 Therefore, results using TPC values need to be interpreted with caution. To address the non-300 specificity, the individual phenolic acids in potato extracts were identified using HPLC-DAD-301 MS (typical chromatogram in supplementary material 1). 5-CQA was the predominant 302 phenolic acid in the flesh of raw and cooked potatoes. Other minor peaks were also observed 303 but not quantified. The content of 5-CQA in raw potato ranged from the lowest in Désirée (10.36 mg 100 g⁻¹ DW) to highest in Maris Piper (29.46 mg 100 g⁻¹ DW) (table 2). In cooked 304 305 tuber, the 5-CQA content was lowest in Rooster (6.51 mg 100 g⁻¹ DW) and highest in Maris 306 Piper (21.24 mg 100 g-1 DW). Similar to the effects of cooking on TPC, steam cooking 307 significantly decreased the 5-CQA level of all potatoes tested ($p \le 0.05$), except Désirée which 308 saw an increase. 5-CQA in cooked Désirée was 1.7 time higher than in raw potato (p≤0.05). 309 The level of chlorogenic acids in 50 unpeeled potato genotypes was reported by Navarre, Pillai, Shakya and Holden (2011) to be in the range 21.9 to 473.0 mg 100g⁻¹ DW. Mattila and 310 Hellstrom (2007) reported that cooked and peeled potato contain 4.13 to 31.2 mg 100 g⁻¹ DW, 311

312 in agreement with the current study. The colour of the skin has been suggested to be associated 313 with TPC content in the flesh (Tierno, Hornero-Mendez, Gallardo-Guerrero, Lopez-Pardo and 314 Ruiz de Galarreta, 2015). However, in this study, this was not found to be the case. Observed 315 colour was not a good indicator of TPC and 5-CQA content in flesh. Rooster has red skin but 316 its flesh showed one of the lowest TPC and 5-CQA contents amongst the varieties investigated. 317 Conversely, Maris Piper has yellow skin and had the highest 5-CQA in cooked flesh. The 318 HPLC results indicate that TPC content overestimates potato phenolic content but is still used 319 in many studies.

320 **3.3 In vitro starch hydrolysis of steam cooked potato**

321 Hydrolysis of potato starch by PPAA was monitored by measuring the reducing sugar produced 322 at different times using the DNS colorimetric method and the maltose content by HPAEC-PAD 323 (typical chromatogram in supplementary material). The amount of sugar detected after PPAA 324 hydrolysis was around 1.5 times higher using the DNS method than using HPAEC-PAD. This 325 can be explained by DNS reacting with oligosaccharides produced by amylase digestion that 326 can react with DNS (Van der Maarel, Van der Veen, Uitdehaag, Leemhuis and Dijkhuizen, 327 2002, Nigam and Singh, 1995, Robyt and Whelan, 1972). AMG was used to digest PPAA 328 digestion products to glucose (figure 2) and in this case DNS and HPAEC-PAD gave similar 329 results (data not shown). Using HPAEC-PAD, free soluble sugars including glucose, fructose 330 and sucrose were detected at low levels at time 0 (supplementary material and table 2). While 331 after hydrolysis by PPAA, maltose and maltotriose were detected as products of digestion 332 (supplementary material) and after AMG digestion, glucose was the only product (not shown). 333 The HPAEC-PAD also showed that the PPAA preparation contains sucrose, most probably as 334 a stabiliser and this needs to be corrected for. Sucrose does react with DNS and therefore it is 335 important to undertake a blank correction of the enzyme preparation. Enzymatic kits were not

used because polyphenols inhibit the kit enzymes, underestimating the carbohydrate content(data not shown).

338 The area under the curve (AUC) for the in vitro digestion reactions with PPAA and 339 PPAA+AMG were calculated for each potato type (table 3). There were significant ($p \le 0.05$) 340 differences in the AUCPPAA and AUCPPAA+AMG between the varieties. Rooster showed 341 consistently the highest extent of digestion. Meanwhile, Mozart and Maris Peer showed lower 342 levels of digestibility with PPAA than Désirée and Maris Piper, but upon digestion with 343 PPAA+AMG, the opposite trend was observed with Désirée and Maris Piper showing lower 344 levels of digestibility compared to Mozart and Maris Peer. While there was no significant 345 correlation between AUC_{PPAA} and TPC or 5-CQA content there was a significant negative 346 correlation between AUCPPAA+AMG and 5-CQA and between AUCPPAA+AMG and TPC (with 347 Pearson correlation coefficients r = -0.49, p < 0.0001 and r = -0.30, p = 0.0001 respectively).

In this study, the amount of steam cooked potato used as substrate contained 10 mg mL⁻¹ of starch (similar to the first experiments) and around 0.08 mg mL⁻¹ 5-CQA. According to figure 1B, this is a low concentration of 5-CQA, enough to inhibit up to 5.8 % of the alpha amylase activity. This amount of 5-CQA does appear to be enough to inhibit AMG. To better understand the relation between composition factors and starch digestibility, principle component analysis (PCA) was performed.

354

355 **3.4 Principle component analysis (PCA)**

Principle component analysis was conducted in order to determine the relationship between potato composition (TS, DM, TPC, 5-CQA) and percent of hydrolysed starch (as measured using AUC_{PPAA+AMG}). Figure 3 show the loading and scores of the characteristics of the five potato cultivars with the first two PCs explaining 72% of the total variance respectively. PC1 explained 47% and PC2 explained 24% of total variance. For PC1, TS, DM, 5-CQA and TPC had positive loading and starch hydrolysis had negative loading. TS and DM had positive loading for PC2 while 5-CQA and TPC had negative loading for PC2. According to the PCA scores, potato varieties segregated according to their components: Maris Piper showed high DM, high 5-CQA content and low digestibility. Rooster, Maris Peer and Mozart potato appeared in one cluster and showed the opposite characteristics. The digestibility of Désirée appeared to be heavily influenced by TPC, in contrast to the other varieties. It appears that the same single factor does not strongly determine digestibility in all varieties.

368 4. Conclusions:

This study is the first to examine the mechanism of inhibition of pancreatic alpha amylase by 5-CQA using potato starch as a substrate. Kinetic analyses showed a mixed-type inhibition, with stronger inhibition at earlier incubation times (5 min compared to 20 min). In vitro digestion of cooked potato tubers showed that the 5-CQA content in tubers is probably too low to affect PPAA digestion, but a significant effect was observed when AMG was also used. The inhibitory effect of 5-CQA on multiple carbohydrate digestive enzymes needs further investigation.

The results presented in this paper suggest that multiple factors affect potato digestibility, and the effects may be variety specific. Testing a higher number of varieties is required and ultimately testing the digestibility of the varieties in vivo will also confirm whether these observations have biological significance.

380

381

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385

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493 **Table and figure captions**

494

495 Table 1. Kinetic parameters K_m and V_{max} of pancreatic alpha amylase enzyme

496 with increasing concentrations of caffeoylquinic (5-CQA).

497

498					
499 500			[5-CQA] mg mL ⁻¹	km mg mL ⁻¹	V _{max} (mg mL ⁻¹ min ⁻¹)
501			0	1.66	0.35
502			1	1.69	0.32
503			1.5	1.85	0.31
504			2	2.08	0.29
505					
506					
507					
F 00	T . 1 1 3 T (1 1	1.		CC 1 · ·	$(\mathbf{F}, \mathbf{O} \mathbf{O} \mathbf{A})$

Table 2. Total phenolic content (TPC), caffeoylquinic (5-CQA) and dry matter (DM) content 508 509 in raw and cooked potatoes.

510

Potato	$TPC (mg \ 100 \ g^{-1} \ DW)^*$		5-CQA (mg 100 g ⁻¹ DW)		DM (g 100 g ⁻¹ FW)	
varieties	Raw	Cooked	Raw	Cooked	Raw	Cooked
Desiree (R)	445.14±32.89 ^b	543.96±20.68ª	10.36±0.16 ^d	17.83±0.40 ^b	21.04±0.10 ^{ac}	20.08±0.11 ^{abc}
Mozart (R)	524.64±27.10 ^a	425.53±17.36 ^b	16.84±0.59 ^b	11.12±0.31°	16.12±0.81 ^b	19.10±1.13ª
Rooster (R)	320.59±15.34°	282.03 ± 5.07^{b}	13.07±0.70°	6.51±0.23 ^e	22.12±3.13ª	19.52±0.10 ^{abc}
Maris Piper (Y)	375.69±29.06 ^d	349.18±13.58°	29.46±1.19 ^a	21.24±0.42ª	20.67±0.10 ^{ac}	21.47±0.61 ^b
Maris Peer (Y)	528.94±15.72ª	343.83±11.57°	11.12±1.86 ^{cd}	7.40 ± 0.35^d	18.90±0.47 ^{acd}	19.66±0.22 ^{ac}

R= red skin, Y= yellow skin; DW = dry weight; FW = fresh weight

*Chlorogenic acid equivalent

Values are means of four determinations±SD

511 512 513 514 Mean value within a column with different subscript letter ^{a,b,c,d,e} indicate significant differences (one-way ANOVA, $P \le 0.05$).

515

518 Table 3. Composition of peeled potatoes in terms of carbohydrate content (total starch (TS) 519 and free sugars), and area under curve (AUC) of hydrolysed starch in cooked tubers. AUCPPAA 520 is AUC for starch hydrolysed by porcine pancreatic alpha amylase (PPAA) alone, while 521 AUC_{PPAA+AMG} is for starch hydrolysed by PPAA followed by amyloglucosidase (AMG). 522

523

Potato varieties	TS (g 100 g ⁻¹ FW) [*]	Free sugars (g 100 g ⁻¹ FW)**	AUC _{PPAA} g maltose. min ^{****}	AUC _{PPAA+AMG} g glucose. min***
Desiree	13.4±1.44 ^b	2.64±0.06ª	517.35±12.82 ^b	752.59±29.19°
Mozart	17.48±1.22ª	1.06±0.02 ^b	484.05±5.21°	970.41±38.82 ^b
Rooster	13.61±0.80 ^b	1.15±0.01 ^b	599.25±1.31ª	1046.43±48.58ª
Maris Piper	16.97±3.62 ^{ab}	0.98 ± 0.06^{b}	$507.00{\pm}11.51^{b}$	648.22 ± 22^{d}
Maris Peer	15.76±2.44 ^b	1.09±0.02 ^b	486.27±5.30°	1044.82±41.32ª

524 525 FW=Fresh weight *(n=9), **(n=3), ***(n=4)

Mean value within a column with different subscript letter ^{a,b,c,d} indicate significant differences (one-way ANOVA, $P \le 0.05$). 526





Figure 1. Effect of 5-caffeoylquinic acid (5-CQA) on the activity of porcine pancreatic alpha amylase (PPAA) activity using 1% potato starch as substrate. A) Effect of pre-incubation and co-incubation of 5-CQA (1.5 mg mL⁻¹) on the digestion of 1% starch by PPAA (0.33 unit mL⁻

1.5

538	¹) at 5 and 20 minutes incubation. $E = enzyme$, $S = substrate and I = inhibitor$. B) Effect of
539	increasing concentration of 5-CQA on inhibition of PPAA relative to control (no inhibitor) at
540	5 min incubation. Each bar represents mean of four measurements \pm standard deviation. Bars
541	with different letters have significant difference at p<0.05. C) Lineweaver-Burk plot for porcine
542	pancreatic alpha amylase (PPAA) catalysed hydrolysis of increasing concentration of potato
543	starch in the presence of 5-caffeoylquinic acid (5-CQA; 0, 1, 1.5 and 2 mg mL ⁻¹). Each point
544	represents mean of four measurements ± standard deviation.





Figure 2. Starch hydrolysis curves of steam cooked potato samples digested with 0.66 units of 547 porcine pancreatic alpha amylase (PPAA) followed by 180 units of amyloglucosidase (AMG).

- 548 A) Désirée B) Mozart. C) Rooster. D) Maris Piper. E) Maris Peer. Data expressed as amount
- 549 of glucose released and as % of total starch (TS) at each time point as measured by DNS (n=9).
- 550 Error bars represent standard deviation of mean.







Figure 3. Principle component analysis (PCA) using composition and digestibility data from
five potato varieties. Graph shows loading of potato components; total starch (TS), dry matter
(DM), total polyphenol content (TPC), 5-caffeoylquinic acid (5-CQA) content and percentage
total starch hydrolysed (%TSH) and scores of potato varieties according to PCA 1 and 2.

561 Supplementary material

562

563 **Figure 1.**

564 HPLC-DAD chromatogram of A) standard compounds, B) raw potato sample (Desiree) and

565 C) cooked potato sample (Desiree). Peaks were detected at wavelength 330 nm. Peaks; (1) 5-

566 caffeoylquinic (5-CQA) (2) sinapic acid (external standard).



- 583 Supplementary material
- **Figure 2.**
- 585 HPAEC-PAD chromatogram of total sugars present after digestion of a potato sample (Maris
- 586 Piper) for 120 min. Peaks; (1) fucose (external standard), (2) glucose, 3) fructose, 4) sucrose,
- 5) maltose, 6) maltotriose. Glucose, fructose and sucrose were already present at time 0 (not
- 588 shown).

