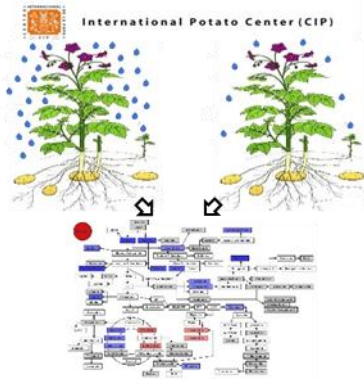


## Graphical Abstract

Margit Drapal, Evelyn R. Farfan-Vignolo, Raymundo O. Gutierrez, Merideth Bonierbale, Elisa Mihovilovich, Paul D. Fraser\*



Physiological and metabolic adaptations were integrated to elucidate the difference in tolerance/ susceptibility to water restriction between five potato clones.

Highlights: (85 characters each)

- Majority of metabolic adaptations were observed in the leaf tissue.
- Late onset of changes in chlorophyll and phenolics confers higher tolerance.
- Genetic composition can overcome the environmental effect.
- Metabolite profiling highlighted distinct metabolic signatures per genotype.

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4 1 **Identification of metabolites associated with water stress responses in *Solanum***  
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11 4 Drapal, M.\*<sup>1</sup>; Farfan-Vignolo, E.R.\*<sup>2</sup>; Gutierrez, O.R.<sup>2</sup>; Bonierbale, M.<sup>2</sup>; Mihovilovich, E.<sup>2</sup>;  
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50 18 Number of Figures: 3 (Fig. 2 and 3 are coloured)  
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53 19 Supplementary data: 2 Figures, 3 Excel files  
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21 **Abstract (200 words)**

22 Water deficiency has become a major issue for modern agriculture as its effects on crop  
23 yields and tuber quality have become more pronounced. Potato genotypes more  
24 tolerant to water shortages have been identified through assessment of yield and dry  
25 matter. In the present study, a combination of metabolite profiling and  
26 physiological/agronomical measurements has been used to explore complex system  
27 level responses to non-lethal water restriction. The metabolites identified were  
28 associated with physiological responses in three different plant tissues (leaf, root and  
29 tuber) of five different potato genotypes varying in susceptibility/tolerance to drought.  
30 This approach explored the potential of metabolite profiling as a tool to unravel sectors  
31 of metabolism that react to stress conditions and could mirror the changes in the plant  
32 physiology. The metabolite results showed different responses of the three plant tissues  
33 to the water deficit resulting either in different levels of the metabolites detected or  
34 different metabolites expressed. The leaf material displayed the most changes to  
35 drought as reported in literature. The results highlighted genotype-specific signatures to  
36 water restriction over all three plant tissues suggesting that the genetics can  
37 predominate over the environmental conditions. This will have important implications for  
38 future breeding approaches.

40 **Keywords:** *Solanum tuberosum* L., Solanaceae, metabolite profiling, plant response,  
41 water restriction, tolerance, phenolic compounds

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# 43 1. Introduction

44 Our changing climate, with more frequent fluctuations in temp. and precipitation levels,  
45 are affecting the yield and quality of crops, such as potato, one of the most important  
46 crops in the world (Bates et al., 2008). The limitation of water, like many other abiotic  
47 stresses, is a particular issue in many crop plants and mobilises multiple sectors of the  
48 plant metabolism as part of its stress response. Hence, presently it is difficult to  
49 associate a single key metabolite with the environmental adaptation to abiotic stresses  
50 (André et al., 2009; Sánchez-Rodríguez et al., 2011). The unpredictability of weather  
51 conditions, especially at critical periods such as plant emergence and tuberisation, can  
52 affect the plant development and lead to dramatic yield losses (Martínez et al., 1992;  
53 Obidiegwu et al., 2015). This emphasises the need for improved potato varieties that  
54 can withstand those environmental stresses (Evers et al., 2010).

55 Tolerance to water limitation in plants is a well-studied topic and has been intricately  
56 linked to photosynthesis and carbohydrate metabolism (Zingaretti et al., 2013). The  
57 response pattern of plants to water restriction is regulated by the intensity, duration and  
58 rate of progression of imposed water restriction and consists of a combination of  
59 physiological and cellular adaptations (Jefferies et al., 1993; Obidiegwu et al., 2015).

60 Most of the water restriction related changes can be detected in the leaves, the energy  
61 production organ of plants. One common response in leaves under mild to moderate  
62 water limitation is to lower their stomatal conductance, thus helping to maintain the  
63 water balance of the plant. Under prolonged water restriction, the cell turgor is affected  
64 despite the stomatal closure which can be counteracted through osmotic adjustment.  
65 Thus, the osmotic potential of the cell is lowered through accumulation of osmolytes

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66 such as amino acids, sugars and polyols inside the cell (Evers et al., 2010; Morgan,  
67 1984; Zingaretti et al., 2013). A side effect of stomatal closure is a reduction in net  
68 photosynthesis due to limited intracellular CO<sub>2</sub> levels (Chaves et al., 2003) which in  
69 consequence leads to an increase in photorespiration, a lower carbon fixation and  
70 ultimately in a reduced partitioning of assimilates affecting the tuber initiation and  
71 bulking (Loon, 1981).

72 Water limitation, equal to many stress conditions, causes a rise of reactive oxygen  
73 species (ROS) which can lead to oxidative damage within the cells. Genes implicated in  
74 the detoxification of ROS were identified as up-regulated in potato upon drought stress  
75 (Watkinson et al., 2008). Most of these genes are responsible for increased production  
76 of metabolites (e.g. phenylpropanoids), with antioxidant properties, able to scavenge  
77 ROS (Cruz de Carvalho, 2008; Dixon et al., 1995).

78 Metabolite profiling approaches have the potential to provide valuable phenotypic  
79 information on a broad range of metabolites across metabolic pathways that contribute  
80 to plant processes and responses to external stimuli or stresses (Jespersen et al.,  
81 2015). Capturing metabolite levels over a resolved time-course, can provide successive  
82 snapshots of the metabolism which provides a dynamic overview of the current  
83 metabolic status of the plant. This methodology is useful in investigating the complex  
84 interacting mechanisms of cellular metabolic pathways in response to perturbations  
85 such as drought stress (Kim et al., 2007).

86 The need for more tolerant potato varieties that can withstand fluctuation in water  
87 availability and still deliver good yields and quality has becomes the focus of modern  
88 breeding programs. For a better understanding of the different drought sensitivity of five

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89 potato breeding clones, those clones were subjected to water restriction followed by a  
90 metabolite profiling approach to establish global changes arising across sectors of  
91 metabolism. The present study focused on untargeted and targeted analysis of  
92 metabolite features in leaves at 30 days after water restriction initiation (DAWRI) and  
93 leaves, roots and tubers at 70 DAWRI of five potato genotypes under a control  
94 treatment (CT) and water restriction (WR). Additionally, physiological properties of the  
95 plants, including osmotic potential (OP), photosynthetic activity ( $A_{sat}$ ) and several others,  
96 were measured to monitor adaptive responses of the clones. Collectively, these data  
97 will enable (I) the assessment of the different responses to water restriction throughout  
98 the plant, (II) link physiological and metabolic properties and (III) connect the degree of  
99 tolerance to key metabolites and/or biochemical pathways.

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## 2. Results and Discussion

### 2.1 *Physiological and agronomical measurements defining tolerance/susceptibility*

The WR treatment was started 28 days after sprout transplanting in order to expose the plants to water stress during tuberisation and bulking and evaluate the resulting metabolic changes throughout the potato plant. PCA analysis was used to identify the physiological traits with the greatest contribution to the five genotypes varying in tolerance to drought. Physiological traits with the highest eigenvalues (top 15%) were considered to have the greatest impact for a component. The biplot (Supplementary Fig.1B) of physiological/agronomical data (loadings) and genotypes (scores) highlighted a clear clustering between CT and WR treatments. The loadings HI of dry and fresh weight, CCI at all time points and OP at 0 to 15 DAWRI were associated with the WR cluster and the loadings biomass of fresh and dry weight,  $A_{\text{sat}}$  and  $g_s$  at 15 DAWRI and RWC at 35 and 70 DAWRI were correlated with the CT cluster. This was to be expected as well-watered plants have more excess water available (RWC) leading to “normal” plant performance /biomass production, whereas plants under water-stress focus their intracellular processes on tuber development instead of leaf production resulting in a higher HI (Obidiegwu et al., 2015).

For agronomical purposes, it is important that the plant maintains tuber formation and development despite the water dependent reduction in photosynthetic capacity (Legay et al., 2011). Therefore, biomass and HI define the DTI and are distinct indicators to assess differences in tolerance between potato clones (Table 3) (Cabello et al., 2013).

The clones, investigated in this study, were chosen for their abiotic stress properties and showed similar HI to previous studies (e.g. Ramírez et al., 2015a). All clones had



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124 higher HI values under WR and according to those same values, CIP397077.16 had the  
125 best performance followed by CIP390637.1, CIP392797.22, CIP394611.112 and  
126 CIP395448.1. Interestingly, CIP397077.16 showed no significant difference between the  
127 HI of dry and fresh weight under WR and CT conditions. The total biomass at 70DAWRI  
128 showed that the clones with lower HI values, CIP392797.22, CIP394611.112 and  
129 CIP395448.1, weighed more than the two better performing clones. Hence, the DTI was  
130 calculated to establish a precise ranking of tolerance of the five clones investigated. The  
131 DTI values (Supplementary Fig.1A) described CIP397077.16 as the most tolerant  
132 followed by CIP392797.22, CIP394611.112, CIP390637.1 and CIP395448.1. In this  
133 respect, the DTI results for CIP395448.1 differed from the expected high tolerance  
134 properties established in the breeding program (Table 2) (Mihovilovich et al., 2015).  
135 This indicated that under the growth conditions of the present study the CIP395448.1  
136 clone could not use the available water as effectively as reported previously (Ramírez et  
137 al., 2015b).

138 The underlying metabolic processes of the leaf, tuber and root material were evaluated  
139 with untargeted analysis, revealing between 3000 and 7000 metabolite features, and  
140 with targeted analysis, identifying ~60 compounds including primary (e.g. glucose, malic  
141 acid) and secondary metabolites (e.g. carotenoids, phenolic compounds). The  
142 untargeted analysis was visualised as a PCA plot (Fig. 1) and highlighted the  
143 environmental conditions (CT and WR) as the main differentiator within each of the  
144 three plant parts, as seen for the physiological/agronomical data (Supplementary Fig.1).  
145 This corroborates that the metabolome is linked to the physiological properties and vice  
146 versa (Peng et al., 2015). The least separation was detected within the root material

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(Fig. 1D). CIP395448.1, the only clone with a BW population background, showed a separate cluster from the other four clones within both treatments. This indicated significant differences in the metabolic composition. For a more detailed view of the clustering patterns, PCA analysis was performed of solely CT or WR treated leaf samples at 30 and 70 DAWRI (Supplementary Fig.2). The clusters of the clones within each treatment had the same trends as the combined score plots (Fig. 1A, B). However, leaves under WR at 70 DAWRI showed the least pronounced cluster (Supplementary Fig.2D), which suggested similar metabolic adaptations under stress conditions (e.g. accumulation of phenylpropanoids (Wegener et al., 2015)) despite the genotypic differences. The PCA analysis of leaves and tuber showed a general trend of clusters (Fig. 1A, B) and indicated a similar metabolic composition of the tolerant clone CIP397077.16 and CIP390637.1. Even though, the latter clone showed a lower DTI than expected. This would suggest that the majority of metabolic changes activated under WR are the same between the two clones and a few specialised pathways influenced the better performance of the tolerant clone CIP397077.16. The cluster of the medium-tolerant clone CIP392797.22 was located closely to the cluster of the latter two clones. The sensitive clone CIP394611.112 of the LTVR population clustered the furthest away from the other three LTVR clones and was closest to CIP395448.1, the other sensitive clone with BW population background. This is an interesting finding insofar, as the metabolic differences of the two sensitive clones could be influenced by the different backgrounds and their metabolic similarities could be related to metabolic traits causing the susceptibility to the WR condition.

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169 The metabolic changes, specific to each clone under WR conditions, were elucidated  
170 through the more detailed targeted analysis. For this purpose, a direct comparison of  
171 metabolite levels under WR and CT was performed and revealed a variety of changes  
172 and similarities between genotypes and population background (Fig. 2) which will be  
173 discussed hereafter in connection with physiological traits.

## 174 *2.2 Photosynthetic activity and chlorophyll content under WR*

175 The chlorophyll content was measured as CCI in SPAD units and together with other  
176 photosynthesis related isoprenoids by UPLC. The SPAD units showed significant  
177 differences between clones at all measurements (0, 15, 25 and 70 DAWRI), but  
178 statistical differences between treatments were evidenced at 25 DAWRI, and genotype  
179 by treatment interaction at 70 DAWRI (Supplementary File 1). In average, CCI of plants  
180 subjected to WR were significantly greater than those of the CT at all times points  
181 (Table 3), even before the WR treatment was initiated (0 DAWRI), which need to be  
182 taken into account for the evaluation of further time points. The CCI values for both  
183 plant treatments increased until the last measurement at 70 DAWRI with the exception  
184 of the two tolerant clones (CIP397077.16 and CIP390637.1). Both these clones had  
185 significantly decreased CCI values under WR conditions, whilst the other clones  
186 retained higher values than their controls. All breeding clones but the tolerant clone  
187 CIP397077.16 increased their CCI values significantly under WR at 25 DAWRI. The two  
188 susceptible clones CIP394611.112 and CIP395448.1 also showed a significant increase  
189 in CCI values at 25 DAWRI and maintained the increased CCI values until 70 DAWRI.

190 The UPLC measurements at 35 DAWRI showed no significant change of any of the  
191 photosynthesis related isoprenoids for the four LTVR clones and for CIP395448.1

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192 decreased levels were detected of antheraxanthin and two violaxanthin isomers by less  
193 than 0.1 fold. At 70 DAWRI, CIP395448.1 showed no change of any isoprenoids,  
194 whereas the four LTVR clones showed significantly decreased levels of chlorophyll *a*  
195 and *b*. Only the genotypes of the LTVR population showed increased levels of  
196 phytoene, a carotenoid precursor, by ~1.1 fold at 35 and 70 DAWRI. This could indicate  
197 a switch of GGPP utilisation from the predominant use for chlorophyll synthesis to  
198 carotenoid synthesis (Zingaretti et al., 2013) and correlates to downregulated genes for  
199 chlorophyll synthesis (Legay et al., 2011).

200 A comparison of the two chlorophyll measurements indicated very different results.  
201 However, certain trends of those results become visible when both analysis are used for  
202 the interpretation of adaptations under WR. The adjustment of the photosynthesis  
203 processes is directly linked to the utilisation of available water resources and the carbon  
204 allocation to the tuber (Obidiegwu et al., 2015). The chlorophyll content seemed to be  
205 the determining factor for the photosynthesis processes in the clones studied, as both  
206  $A_{sat}$  and  $g_s$ , known to be lowered under drought to conserve water (Obidiegwu et al.,  
207 2015), did not change for any clone at any time point measured. The mid-tolerant clone  
208 CIP392797.22 had a similar chlorophyll/violaxanthin profile to the tolerant  
209 CIP397077.16 and also showed a decrease of CCI from 25 to 70DAWRI. This might be  
210 a factor for the better performance of CIP392797.22 than the reportedly tolerant  
211 CIP390637.1, which showed a decrease of CCI but far less change of chlorophyll  
212 levels. Furthermore, the chlorophyll content could be a deciding factor for the metabolic  
213 difference of the two susceptible clone. Even though both showed the same trends of  
214 CCI values, the lack of change in chlorophyll levels of CIP395448.1 probably led to a

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215 quicker consumption of already limited water, enhancing the drought stress on the plant  
216 and reducing the tuber formation or development (Zingaretti et al., 2013).

217 *2.3 OP and RWC and related metabolites affecting tuber formation*

218 Significant differences in RWC between treatments were detected at 35 DAWRI with an  
219 average RWC decrease of 5% under WR and a 7% increase under CT (Table 3). This  
220 trend continued up to 70 DAWRI and showed statistical differences between breeding  
221 clones (Supplementary File 1). WR samples showed a further 18% decrease at 70  
222 DAWRI resulting in RWC values between 60-70%, whereas CT samples maintained  
223 RWC values above 90%. Interestingly, the OP only showed difference between  
224 treatments at 15 DAWRI and remained similar for the remaining experiment between  
225 control and stressed plants and between genotypes (Table 3). CIP394611.112 under  
226 WR showed statistically lower OP values than under CT at 15 DAWRI and maintained  
227 decreasing OP values until the last measurement. The other four clones maintained  
228 their cellular pressure under WR treatment (Supplementary File 1).

229 On the biochemical level, most of the metabolite changes associated with drought  
230 responses (e.g. sugar and osmolytes accumulation (Zingaretti et al., 2013)) were  
231 detected in the leaves of the LTVR accessions. The metabolite changes common to all  
232 five genotypes in leaf were a decrease of citric acid and increases of gluconic acid,  
233 ribitol and metabolites of the phenylpropanoid pathway, in particular phenylalanine and  
234 naringin. The increase of naringin was above 2.5 fold for all genotypes at 70 DAWRI.  
235 The highest increase of naringin in response to water restriction was detected in  
236 CIP397077.16 by 68.2 fold at 70 DAWRI, followed by CIP390637.1 and CIP394611.112  
237 with a 37.8 and 28.1 fold increase, respectively. The increase of phenylalanine and the

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238 decrease of citric acid was less at 70 DAWRI than at 30 DAWRI for all genotypes  
239 except CIP397077.16. The lack of change in osmolytes levels and reduced levels of  
240 phenolic compounds in the sensitive clone CIP395448.1 appears to be a contributing  
241 factor to this clone's comparative inability to adapt to drought conditions. The lack of  
242 osmolytes and phenolic compounds under water deficiency leads to a deformation of  
243 the transportation system of the plant (xylem) and further interferes with its ability to  
244 retain water from the soil (Zingaretti et al., 2013). These phenotypic effects were  
245 observed for the sensitive clone CIP395448.1, as both the OP and RWC values,  
246 displaying cellular pressure and turgidity, were reduced at 25 and 70DAWRI. The  
247 second highest reduction of RWC and OP values was observed for the other sensitive  
248 clone CIP394611.112. Contrary to this, one of the most distinct features of the tolerant  
249 clone CIP397077.16 was a delayed increase of phenylpropanoid formation in the  
250 leaves. This delayed increase of phenolic compounds in combination with the induction  
251 of one specific phenolic compound (naringin, a glycoside of naringenin) could infer a  
252 more efficient ROS scavenging mechanism of this genotype under drought. Activation  
253 of naringenin chalcone synthase has been shown to play a particular role for tolerance  
254 of potato plants to stress conditions (Vasquez-Robinet et al., 2008). Furthermore, the  
255 high levels of phenolic ROS scavengers could delay leaf senescence induced by  
256 oxidative stress and aid the photosynthetic processes leading to a better carbon fixation  
257 and resource allocation to the tuber which corresponds to the better DTI performance of  
258 this clone compared to the others (Obidiegwu et al., 2015; Rivero et al., 2007;  
259 Schippers et al., 2007).

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260 Another metabolic trait related to tuber formation/development is the allocation and  
261 conversion of sucrose to starch. Within the leaf samples, the clones of the LTVR  
262 population showed increased levels of saccharides such as fructose, glucose, inositol  
263 and ribitol of which most had an above 2.5 fold increase at 70 DAWRI. No change of  
264 sucrose levels was detected in leaf of any of the genotypes. The components of the  
265 TCA cycle from succinic to malic acid were increased from ~2.0 to above 2.5 fold at 30  
266 and/or 70 DAWRI, contrary to the 0.5 fold decrease of citric and isocitric acid. The clone  
267 CIP395448.1 showed very little metabolite changes, which included a decrease of citric  
268 and isocitric acid by 0.6-0.7 fold at 30 DAWRI and a 1.7 fold increase of ribitol and 0.7  
269 fold decrease of mannose at 70DAWRI. The metabolite levels in the LTVR clones  
270 suggested an active turn-over from fructose to the TCA cycle which was reported for a  
271 resistant accession as a result of higher mitochondrial metabolic activity (Vasquez-  
272 Robinet et al., 2008). Within the tuber samples, no change of glucose/fructose or  
273 intermediates of the TCA cycle could be seen (Fig. 3). These leaf and tuber results  
274 suggest that only the yield but not the development of the tuber was affected, as seen  
275 by the biomass/HL results and previously described for an acclimated accession of  
276 *Solanum tuberosum ssp. andigena* (Watkinson et al., 2008).

#### 277 *2.4 Phenylpropanoids and other metabolites in tuber*

278 Stress-induced metabolism can vary between different plant parts (Dixon et al., 1995)  
279 and the range of metabolites identified in tubers, 45 in total (Fig. 3), varied as expected  
280 from the leaf material. The intermediates of the phenylpropanoid pathway varied not  
281 only in content but also in composition between the leaves, tuber and roots. The three  
282 phenylpropanoids, naringenin, rutin and umbelliferone, detected with increased levels in

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283 the leaf material of WR treated plants, showed no change or even decreases in the root  
284 material. An exception was the susceptible clone CIP395448.1 which showed an  
285 increase of naringin in the roots. This metabolic adaptation could be related to the  
286 clone's BW resistance as naringin and other flavonoids in roots are known to confer  
287 antimicrobial properties against soil-borne bacteria (Szoboszlay et al., 2016). In tuber,  
288 only umbelliferone was detected next to other phenolic compounds such as chlorogenic  
289 acid, as reported previously (Navarre et al., 2013). The genotypes CIP397077.16 and  
290 CIP395448.1 had increased levels of caffeic acid, ferulic acid, chlorogenic and  
291 cryptochlorogenic acid. CIP392797.22 only had increased levels of caffeic and ferulic  
292 acid, whereas the remaining two clones showed no change in phenylpropanoids. The  
293 importance of phenolic compounds, despite the different expression in the three plant  
294 parts, was shown through the phenylalanine content which was induced (2.6 to 4 fold) in  
295 all genotypes in leaf and tuber and supports the hypothesis that drought stress is  
296 perceived differently in each plant part and activates tissue specific regulators (Dixon et  
297 al., 1995; Sánchez-Rodríguez et al., 2011).

298 Amino acids were the second metabolite group with significant changes in tuber.  
299 Glutamine, a central component in plant nitrogen metabolism, was increased by 2.3 to  
300 3.0 fold in four of the five genotypes, whereas CIP394611.112 produced proline at 25.5  
301 fold under WR conditions. Those ratios of both glutamine and proline were not detected  
302 in the leaf material. Other amino acids, including leucine, valine, threonine, isoleucine,  
303 methionine, tryptophan and serine, were increased in CIP392797.22, CIP394611.112  
304 and CIP390637.1 by 1.9 to 7.4 fold changes and in the case of CIP390637.1 leucine  
305 showed an increase by 15.3 fold. Only the changes of tryptophan were consistent



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306 between tuber and leaf material at 70 DAWRI. These changes of amino acids in  
307 combination with phenylpropanoid results indicated a negative correlation between the  
308 content of free amino acids and phenolic compounds in tuber (Fig. 3). This could be  
309 related to induced protein degradation due to lack of ROS protection (Krasensky et al.,  
310 2012) or an increase of free amino acids acting as osmolytes in addition to the ROS  
311 protection through phenylpropanoids (Yancey, 2001).

312 In general and contrary to the leaf material, hardly any changes of saccharides were  
313 detected in tubers under WR compared to CT conditions. Inositol, relevant for  
314 osmoprotection, was increased in the LTVR clones by a fold change of ~2.4 for  
315 CIP392797.22 and CIP394611.112 and ~5.9 fold in CIP390637.1 and CIP397077.16.  
316 The clone CIP395448.1 showed no change of inositol levels, but had a 1.6 fold increase  
317 of another saccharide, sorbose. Overall, CIP395448.1 tuber material showed the least  
318 metabolic changes under WR and differed very little in the metabolite levels from the  
319 tolerant clone CIP397077.16, even though their leaf material showed significant  
320 differences.

### 321 *2.5 Tolerance properties affecting tuber quality*

322 One aspect that has to be taken into consideration by breeders producing more tolerant  
323 potato varieties, is the change in sensory attributes of the tuber such as taste and  
324 texture. In this study, the distinct metabolic features of clone CIP397077.16 highlighted  
325 genetic traits which can overcome the water stress environment more effectively  
326 compared to the other genotypes. These metabolic features included the late onset of  
327 changes in chlorophyll and phenylpropanoid levels in leaf and increased levels of  
328 phenylpropanoid in tuber. The disadvantage of these increased levels in tuber is that

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329 most phenylpropanoids are known to confer bitterness taste properties (Ripoll et al.,  
330 2014). This has been the unforeseen consequences of various breeding programs  
331 where the focus of biotic and abiotic resistance has precluded consideration of  
332 consumable traits preventing the uptake of the varieties into real life scenarios. The  
333 clone CIP392797.22 would be a suitable compromise between harvest loss and taste  
334 properties. This clone showed the second best DTI, a higher total biomass and no  
335 changes of phenylpropanoid levels in the tuber.

336 As illustrated in the present study, the use of metabolite profiling provides detection of  
337 potential taste/aroma influencing compounds within accessions. The incorporation of  
338 these technique into characterisation of germplasm will enable breeding programs to  
339 pyramid multiple traits for resistance, productivity and also consumer preferences.

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### 341 3. Concluding remarks

342 The present study showed that the successful adaptation to water deficit displayed by  
343 several of the selected clones does have a measurable biochemical component. The  
344 adaptation processes are complex and require the interaction of several key pathways  
345 to maintain key cellular processes necessary for “normal” plant development/growth.  
346 The metabolite profiling highlighted that a distinct metabolic signature can only really be  
347 seen in a comparison to other clones as adaptation to drought stresses involves multi-  
348 pathway adaptation rather than the introduction or presence of one novel  
349 compound/marker (Zingaretti et al., 2013). Monitoring metabolite levels over time, to  
350 create a dynamic picture, demonstrated the ability of the tolerant potato clones to fine  
351 tune metabolite compositions to the degree of stress experienced by the plant and its  
352 capacity to deal with this stress condition (Vasquez-Robinet et al., 2008). Overall, the  
353 incorporation of metabolite profiling datasets augments the elucidation of the  
354 biochemical mechanisms underlying the physiological adaptations.

## 356 4. Experimental

### 357 4.1 *Plant material and experimental design*

358 Five potato genotypes differing in their sensitivity to water stress (Table 1) (Cabello et  
359 al., 2012; Carli et al., 2014; Ramírez et al., 2015a) were selected for their similar  
360 vegetative maturity ( $\leq 120$  days) and population background from CIP breeding program.  
361 Four of the clones are part of the lowland tropic virus resistant (LTVR) population  
362 (Bonierbale et al., 2003) and the fifth clone is part of an ancient population bred for  
363 resistance to bacterial wilt (BW) (Table 1) (Mihovilovich et al., 2015).

364 The experiment was conducted under greenhouse conditions from September to  
365 December 2013, at CIP experimental station in Huancayo, Peru (3200 meter above sea  
366 level). Atmospheric temp. and humidity (HOBO U23 Pro v2 Temp./Relative Humidity  
367 Data Logger –U23-001, Onset Computer Corporation, Bourne, MA, USA) were  
368 recorded every 10 min using a mini data –logger (HOBO U12 Outdoor/Industrial Data  
369 Logger, Onset Computer Corporation, Bourne, MA, USA) (Table 2).

370 Tubers of the five genotypes were planted in plastic trays filled with 1:1 rinsed fine –  
371 textured sand and gravel and after 22 days later, thirty vigorous sprouts of each  
372 breeding clone were extracted (cut out from the tubers) and sown individually in 12  
373 inches pots filled with 15kg of the same sand and gravel mixture. Pots were distributed  
374 following a split block design with three replicates of ten pots per genotype; half of them  
375 received a different water treatment. Plants grown from sprouts were watered by drip  
376 irrigation (4L/h) and all fertilisers were injected directly into the irrigation water.  
377 Fertilisation was performed with  $5.18\text{ g l}^{-1}$  potassium nitrate,  $3.36\text{ g l}^{-1}$  ammonium nitrate,  
378  $2.68\text{ g l}^{-1}$  calcium triple superphosphate,  $1.15\text{ g l}^{-1}$  magnesium sulphate,  $120\text{ g l}^{-1}$  fertilon

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379 combi micronutrients, and 96g l<sup>-1</sup> iron quelate. Pest control was carried out with the  
380 application of two insecticides Rovral (active ingredient: Iprodione; Bayer CropScience,  
381 Peru) and Beta baytroid (active ingredient: Beta cyfluthrin; Bayer CropScience, Peru) at  
382 0.16 g l<sup>-1</sup>.

383 All plants were watered every other day for 30 days before the water restriction (WR)  
384 treatment initiation. The WR treatment consisted of applying half of the amount of water  
385 supplied to the control treatment (CT). Irrigation frequency in WR was performed in an  
386 interval of around nine days depending on daily temp. (Table 2) which was repeated  
387 eight times over a total period of 80 days. The CT plants continued to be watered with  
388 the same frequency as the first 30 days.

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## 390 *4.2 Physiological and agronomical measurements*

### 391 *4.2.1 Total biomass, Harvest Index determination and Drought Tolerance Index*

392 At harvest (80 days after water restriction initiation (DAWRI)), total fr. wt and dry wt  
393 were determined. The dry wt of each plant was measured, in triplicate or more, after  
394 drying all the components at 80°C for 3 days in a forced air oven. The harvest index (HI)  
395 was calculated as the ratio of DW tuber (g) to the total DW biomass (g). The drought  
396 tolerance index (DTI) was calculate according to DTI= ((yield under stress x yield under  
397 control)/mean yield under control) (Saravia et al., 2016).

### 398 *4.2.2 Osmotic potential (OP)*

399 Leaf osmotic potential was determined at 0, 15, 35 and 70 DAWRI in leaf discs of 5mm  
400 diameter, taken from the third fully extended leaf of at least three plants per clone and

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401 treatment. Leaf discs were transferred into cryogenic tubes and frozen in liquid nitrogen  
402 for further analysis. The frozen leaves were incubated at 22°C for 30 min in a sealed C-  
403 52 chamber (Wescor Inc., Logan, UT, USA) prior to determination of the osmotic  
404 potential with a dew point microvoltmeter (HR-33T from the same company).

#### 405 *4.2.3 Relative water Content (RWC)*

406 RWC was determined at 0, 15, 35 and 70 DAWRI by weighing the third leaflet from the  
407 youngest fully expanded leaf (third leaf from the apical part) (fr. wt) and placing it in a  
408 4x3 inch Ziploc bag containing distilled water for 24 h. Excess water was removed by  
409 blotting each leaf in a paper towel prior to taking turgid weight (TW). Leaves were then  
410 dried in an oven at 90 °C overnight and reweighed (dry wt). RWC was calculated  
411 according to  $RWC (\%) = [(fr. wt - dry wt) / (TW - dry wt)] \times 100$  (Barrs et al., 1962). The  
412 RWC mean value was established from triplicate measurements of each clone per  
413 treatment.

#### 414 *4.2.4 Chlorophyll content index (CCI)*

415 Chlorophyll content was determined at 0, 15, 25 and 70 DAWRI as a CCI value using a  
416 portable chlorophyll meter (Minolta SPAD-502, Konica Minolta, Sakai, Osaka, Japan). In  
417 each plant, three leaflets of the youngest fully expanded leaf (third node) were used to  
418 measure the transmittance of red (650 nm) and infrared (940 nm) radiation through the  
419 leaf. The average CCI of the three leaflets provided by the average option of the  
420 instrument was recorded as an individual CCI plant value. This was repeated in at least  
421 three plants of each clone per treatment and replication.

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422 *4.2.5 Photosynthetic activity ( $A_{sat}$ ) and stomatal conductance ( $g_s$ )*

423 Light-saturated net CO<sub>2</sub> assimilation rate ( $A_{sat}$ ) and stomatal conductance ( $g_s$ ) were  
424 measured at 15, 25 and 70 DAWRI on the youngest fully expanded leaves, in at least  
425 three plants of each clone per treatment and replication.  $A_{sat}$  and  $g_s$  were measured with  
426 a portable gas exchange system (LI-6400, Li-COR, Lincoln, USA). Leaf chamber  
427 conditions was set up at 385 ppm CO<sub>2</sub>, 23.5 °C (block temp.), a saturated photon flux  
428 density (1500  $\mu\text{mol m}^{-2} \text{s}^{-1}$ ) and ambient relative humidity.

429 *4.3 Metabolite analysis*

430 *4.3.1 Extraction of metabolites*

431 Leaf tissue samples at 30 DAWRI and leaf, tuber and root tissue samples at 70 DAWRI  
432 of both treatments were frozen in liquid nitrogen immediately after harvest and ground  
433 to powder. Samples, including quality controls (pool of all samples, QC), were weighed  
434 (10±0.5 mg) in plastic tubes and extracted as described previously (Nogueira et al.,  
435 2013). Aliquots of the polar and non-polar phase were immediately dried down after  
436 extraction.

437 *4.3.2 GC-MS analysis.*

438 An aliquot of the polar phase (200 $\mu\text{l}$ ) was removed and internal standard ([D4]Succinic  
439 acid, 10 $\mu\text{g}$ ) added before dry down. Dried samples were derivatised as previously  
440 described (Nogueira et al., 2013) and analysed by GC-MS based on literature (Enfissi et  
441 al., 2010), using a 10:1 split mode. Metabolites were identified with respect to an in-  
442 house library based on retention time, retention indices and mass spectrum (Nogueira  
443 et al., 2013) and quantified relatively to the internal standard.

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#### 444 4.3.3 LC-MS analysis

445 Each dried aliquot of the polar phase (700µl) was resuspended in methanol/water (1:1,  
446 100µl) and internal standard (homogenistic acid, 5µg) added. Samples were filtered  
447 using syringe filter (nylon, 0.45µm) before analysis based on a published LC-MS  
448 method (Enfissi et al., 2010) with modification of solvents A (water and 0.1% formic  
449 acid) and B (acetonitrile and 0.1% formic acid). The gradient was held at 100% A for 1  
450 min, followed by a gradient up to 35% B until 18 min and to 95% B until 19 min. The  
451 gradient was then held at 95% B for 4 min and the column returned to the initial  
452 conditions within 1 min and equilibrated for 5 min. Data analysis was performed based  
453 on R package metaMS (Shahaf et al., 2013; Wehrens et al., 2015) with a retention time  
454 window match set to 0.5min.

#### 455 4.3.4 Chromatographic analysis with UPLC

456 For analysis of carotenoids and chlorophylls, an aliquot of the non-polar phase (350µl)  
457 was dried, resuspended in ethyl acetate/acetonitrile (1:9) and analysed as previously  
458 described (Nogueira et al., 2013). Metabolites were identified through specific retention  
459 time and UV/visible light spectrum and quantified from dose-response curves (Fraser et  
460 al., 2000).

#### 461 4.3.5 Data processing and statistical analysis

462 Principal component analysis (PCA) using a correlation matrix to standardise the  
463 variables was performed to identify components which contributed the greatest variance  
464 between genotypes. Statistical analysis was perform using JMP® Pro 12.0.1 (SAS  
465 Institute Inc, Cary, NC) and Simca P 13.0.3.0 (Umetrics, Sweden). Statistical analysis



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466 was performed using the computing environment R v 3.2.4 (The R Foundation for  
467 Statistical Computing, 2016). Changes in identified metabolites between WR and CT  
468 samples were calculated as average ratios. Their significance was established through  
469 pairwise comparison with Student's *t* test for the metabolic data and Fisher's least  
470 significant difference (LSD) for the physiological data ( $P < 0.05$ ). Results for leaves were  
471 displayed in a heat map using Excel 2013 (Microsoft Office, USA). Metabolite changes  
472 in tuber were overlaid with biochemical pathways constructed specifically with  
473 BioSynLab software ([www.biosynlab.com](http://www.biosynlab.com)).

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476 technical assistance. This work was supported by the CGIAR Research Program on  
477 Roots, Tubers and Bananas (RTB).

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4 **480 Tables**

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7 **481 Table 1: Genotypes analysed in the study**

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<b>Population</b>	<b>CIP Number</b>	<b>Variety name (first country of release)</b>	<b>Female parent</b>	<b>Male parent</b>	<b>Drought sensitivity</b>
LTVR	392797.22	UNICA (Peru)	387521.3	Aphrodite	Tolerant
LTVR	394611.112		780280=(P W-88-6203)	676008=(I-1039)	Susceptible
LTVR	390637.1		PW-31	385305.1=(XY.9)	Medium-tolerant
LTVR	397077.16	SARNAV (Uzbekistan)	392025.7=(LR93.221)	392820.1=(C93.154)	Tolerant
BW	395448.1		393617.1=(TXY.11)	BWH-87.344R	Susceptible

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32 **483**

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35 **484 Table 2: Monthly averages of maximum and minimum temp. and relative humidity**  
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37 **485 measured from September to December 2013.**

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	<b>September</b>	<b>October</b>	<b>November</b>	<b>December</b>
Max. temp. (°C)	28.3 ± 0.6	26.6 ± 0.4	26.5 ± 0.4	24.4 ± 0.5
Min. temp. (°C)	5.6 ± 0.5	7.9 ± 0.3	7.5 ± 0.3	8.8 ± 0.2
Average temp. (°C)	14.7 ± 0.3	14.9 ± 0.2	15.1 ± 0.2	14.6 ± 0.2
Max. rel. humidity (%)	90.8 ± 1.1	94.1 ± 0.6	95.3 ± 0.5	97.8 ± 0.4
Min. rel. humidity (%)	19.9 ± 1.3	27.1 ± 1.2	26.3 ± 1.4	36.2 ± 1.3

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8 **Table 3:** Mean  $\pm$  s.d. (n=3) of physiological parameters under control (CT) and water restriction (WR) treatments of five  
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10 breeding clones. Fischer's least significant difference (LSD) values at  $p < 0.05$  were used to compare between treatments  
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12 by clone (significant difference are indicated in bold) and between clones by treatment (different letters indicate  
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14 statistically significant difference). DAWRI= Days after water restriction initiation, RWC= Relative Water Content (%), OP=  
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16 Osmotic potential (bar),  $A_{sat}$ = Photosynthetic activity ( $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ ),  $g_s$ = Stomatal conductance ( $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ )  
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Clones	Treatment	CCI				RWC				$A_{sat}$		
		0DAWRI	15DAWRI	25DAWRI	70DAWRI	0DAWRI	15DAWRI	35DAWRI	70DAWRI	15DAWRI	25DAWRI	70DAWRI
CIP395448.1	CT	42.5 <sup>a</sup> $\pm$ 2.8	49.0 <sup>a</sup> $\pm$ 3.1	50.2 <sup>a</sup> $\pm$ 3.6	50.6 <sup>a</sup> $\pm$ 3.2	91.7 <sup>a</sup> $\pm$ 6.6	88.2 <sup>a</sup> $\pm$ 5.8	92.5 <sup>a</sup> $\pm$ 1.8	90.8 <sup>a</sup> $\pm$ 1.9	10.4 <sup>a</sup> $\pm$ 4.9	9.6 <sup>a</sup> $\pm$ 5.7	9.6 <sup>a</sup> $\pm$ 2.2
	WR	45.7 <sup>a</sup> $\pm$ 3.2	52.7 <sup>a</sup> $\pm$ 3.8	55.2 <sup>a</sup> $\pm$ 4.9	54.9 <sup>a</sup> $\pm$ 8.6	92.9 <sup>a</sup> $\pm$ 4.7	88.8 <sup>a</sup> $\pm$ 2.1	81.2 <sup>a</sup> $\pm$ 9.1	69.1 <sup>a</sup> $\pm$ 13.7	5.4 <sup>a</sup> $\pm$ 5.3	9.6 <sup>a</sup> $\pm$ 2.9	7.2 <sup>a</sup> $\pm$ 3.7
CIP394611.112	CT	41.0 <sup>ab</sup> $\pm$ 2.9	48.4 <sup>a</sup> $\pm$ 1.8	48.7 <sup>a</sup> $\pm$ 1.8	48.3 <sup>a</sup> $\pm$ 2.5	92.4 <sup>a</sup> $\pm$ 6.9	89.1 <sup>a</sup> $\pm$ 4.9	92.3 <sup>a</sup> $\pm$ 2.7	91.8 <sup>a</sup> $\pm$ 2.5	15.3 <sup>a</sup> $\pm$ 1.2	9.8 <sup>a</sup> $\pm$ 7.5	4.4 <sup>b</sup> $\pm$ 3.9
	WR	42.7 <sup>b</sup> $\pm$ 3.1	49.6 <sup>b</sup> $\pm$ 1.3	53.2 <sup>ab</sup> $\pm$ 2.6	55.1 <sup>a</sup> $\pm$ 2.1	92.0 <sup>a</sup> $\pm$ 4.4	85.5 <sup>ab</sup> $\pm$ 5.4	82.2 <sup>a</sup> $\pm$ 7.9	60.6 <sup>b</sup> $\pm$ 5.8	5.9 <sup>a</sup> $\pm$ 4.6	9.8 <sup>a</sup> $\pm$ 7.7	7.1 <sup>a</sup> $\pm$ 5.6
CIP392797.22	CT	37.7 <sup>cd</sup> $\pm$ 2.5	45.2 <sup>a</sup> $\pm$ 1.8	46.3 <sup>a</sup> $\pm$ 1.9	43.3 <sup>b</sup> $\pm$ 2.7	92.5 <sup>a</sup> $\pm$ 7.8	85.8 <sup>a</sup> $\pm$ 5.7	92.3 <sup>a</sup> $\pm$ 3.7	91.3 <sup>a</sup> $\pm$ 2.8	6.8 <sup>a</sup> $\pm$ 4.6	10.5 <sup>a</sup> $\pm$ 10.1	6.0 <sup>ab</sup> $\pm$ 3.3
	WR	39.4 <sup>c</sup> $\pm$ 2.9	45.5 <sup>c</sup> $\pm$ 1.4	50.1 <sup>b</sup> $\pm$ 2.8	47.9 <sup>b</sup> $\pm$ 2.0	92.2 <sup>a</sup> $\pm$ 4.8	81.2 <sup>b</sup> $\pm$ 8.6	78.2 <sup>a</sup> $\pm$ 8.4	69.3 <sup>a</sup> $\pm$ 3.4	8.0 <sup>a</sup> $\pm$ 6.2	9.7 <sup>a</sup> $\pm$ 6.1	7.2 <sup>a</sup> $\pm$ 3.9
CIP397077.16	CT	39.1 <sup>bc</sup> $\pm$ 1.3	43.2 <sup>bc</sup> $\pm$ 1.7	40.1 <sup>b</sup> $\pm$ 1.1	39.7 <sup>bc</sup> $\pm$ 2.3	95.8 <sup>a</sup> $\pm$ 2.8	85.5 <sup>a</sup> $\pm$ 3.0	91.2 <sup>a</sup> $\pm$ 6.0	93.3 <sup>a</sup> $\pm$ 1.3	8.5 <sup>a</sup> $\pm$ 4.0	8.4 <sup>a</sup> $\pm$ 4.5	7.6 <sup>ab</sup> $\pm$ 4.6
	WR	41.5 <sup>b</sup> $\pm$ 1.4	43.8 <sup>cd</sup> $\pm$ 0.9	45.1 <sup>c</sup> $\pm$ 3.3	34.6 <sup>c</sup> $\pm$ 2.7	94.4 <sup>a</sup> $\pm$ 4.1	85.6 <sup>ab</sup> $\pm$ 4.4	81.7 <sup>a</sup> $\pm$ 10.1	60.4 <sup>b</sup> $\pm$ 9.7	11.1 <sup>a</sup> $\pm$ 8.6	9.9 <sup>a</sup> $\pm$ 5.0	8.9 <sup>a</sup> $\pm$ 4.5
CIP390637.1	CT	36.9 <sup>d</sup> $\pm$ 2.3	40.4 <sup>c</sup> $\pm$ 2.5	40.1 <sup>b</sup> $\pm$ 2.4	38.8 <sup>c</sup> $\pm$ 3.3	92.7 <sup>a</sup> $\pm$ 5.0	84.3 <sup>a</sup> $\pm$ 5.5	93.7 <sup>a</sup> $\pm$ 1.0	94.1 <sup>a</sup> $\pm$ 1.9	13.5 <sup>a</sup> $\pm$ 7.8	7.6 <sup>a</sup> $\pm$ 3.7	10.2 <sup>a</sup> $\pm$ 3.0
	WR	38.8 <sup>c</sup> $\pm$ 3.2	42.2 <sup>d</sup> $\pm$ 3.0	45.0 <sup>c</sup> $\pm$ 4.3	36.5 <sup>c</sup> $\pm$ 4.3	93.2 <sup>a</sup> $\pm$ 6.7	82.5 <sup>b</sup> $\pm$ 6.6	79.9 <sup>a</sup> $\pm$ 7.4	72.3 <sup>a</sup> $\pm$ 9.6	10.9 <sup>a</sup> $\pm$ 8.8	9.8 <sup>a</sup> $\pm$ 6.5	9.0 <sup>a</sup> $\pm$ 3.6
LSD values		1.93	2.83	3.93	4.23	4.51	4.83	7.36	8.31	22.87	5.02	5.21

Clones	Treatment	OP				Total Biomass		Harvest Index		$g_s$		
		0DAWRI	15DAWRI	35DAWRI	70DAWRI	fr. wt	dry wt	fr. wt	dry wt	15DAWRI	25DAWRI	70DAWRI
CIP395448.1	CT	-24.2 <sup>a</sup> $\pm$ 2.2	-24.0 <sup>a</sup> $\pm$ 3.8	-21.7 <sup>a</sup> $\pm$ 3.5	-26.0 <sup>a</sup> $\pm$ 3.0	750.5 <sup>c</sup> $\pm$ 482.2	84.0 <sup>bc</sup> $\pm$ 54.4	31.9 <sup>c</sup> $\pm$ 22.2	29.8 <sup>c</sup> $\pm$ 20.1	0.025 <sup>c</sup> $\pm$ 0.01	0.020 <sup>a</sup> $\pm$ 0.02	0.037 <sup>a</sup> $\pm$ 0.02
	WR	-21.5 <sup>a</sup> $\pm$ 2.7	-20.7 <sup>a</sup> $\pm$ 4.5	-21.2 <sup>b</sup> $\pm$ 4.9	-27.4 <sup>a</sup> $\pm$ 4.6	207.0 <sup>a</sup> $\pm$ 106.3	24.0 <sup>a</sup> $\pm$ 10.3	40.5 <sup>d</sup> $\pm$ 25.6	44.6 <sup>b</sup> $\pm$ 30.6	0.017 <sup>a</sup> $\pm$ 0.03	0.036 <sup>a</sup> $\pm$ 0.03	0.019 <sup>a</sup> $\pm$ 0.01
CIP394611.112	CT	-21.5 <sup>a</sup> $\pm$ 3.8	-19.3 <sup>c</sup> $\pm$ 3.7	-16.9 <sup>b</sup> $\pm$ 3.7	-21.3 <sup>b</sup> $\pm$ 4.8	1043.0 <sup>a</sup> $\pm$ 817.7	101.9 <sup>a</sup> $\pm$ 66.1	43.2 <sup>b</sup> $\pm$ 16.8	51.5 <sup>b</sup> $\pm$ 15.2	0.063 <sup>bc</sup> $\pm$ 0.02	0.032 <sup>a</sup> $\pm$ 0.04	0.035 <sup>a</sup> $\pm$ 0.03
	WR	-20.4 <sup>a</sup> $\pm$ 2.9	-18.4 <sup>a</sup> $\pm$ 3.2	-20.1 <sup>b</sup> $\pm$ 4.2	-24.9 <sup>a</sup> $\pm$ 3.3	247.6 <sup>a</sup> $\pm$ 87.9	29.2 <sup>a</sup> $\pm$ 10.8	61.4 <sup>c</sup> $\pm$ 25.8	69.4 <sup>a</sup> $\pm$ 18.9	0.020 <sup>a</sup> $\pm$ 0.02	0.017 <sup>a</sup> $\pm$ 0.02	0.012 <sup>a</sup> $\pm$ 0.01
CIP392797.22	CT	-21.5 <sup>a</sup> $\pm$ 3.8	-20.7 <sup>bc</sup> $\pm$ 2.1	-20.3 <sup>a</sup> $\pm$ 2.6	-26.0 <sup>a</sup> $\pm$ 2.8	1025.2 <sup>ab</sup> $\pm$ 721.7	100.7 <sup>ab</sup> $\pm$ 61.8	49.0 <sup>b</sup> $\pm$ 15.3	58.2 <sup>b</sup> $\pm$ 12.7	0.021 <sup>c</sup> $\pm$ 0.03	0.022 <sup>a</sup> $\pm$ 0.04	0.016 <sup>a</sup> $\pm$ 0.02
	WR	-21.9 <sup>a</sup> $\pm$ 4.7	-19.5 <sup>a</sup> $\pm$ 3.1	-21.1 <sup>b</sup> $\pm$ 3.9	-27.3 <sup>a</sup> $\pm$ 4.2	255.9 <sup>a</sup> $\pm$ 104.7	29.7 <sup>a</sup> $\pm$ 9.5	64.3 <sup>bc</sup> $\pm$ 23.2	71.2 <sup>a</sup> $\pm$ 17.3	0.023 <sup>a</sup> $\pm$ 0.02	0.016 <sup>a</sup> $\pm$ 0.02	0.018 <sup>a</sup> $\pm$ 0.02
CIP397077.16	CT	-22.4 <sup>a</sup> $\pm$ 2.6	-23.1 <sup>ab</sup> $\pm$ 2.2	-22.3 <sup>a</sup> $\pm$ 4.4	-27.2 <sup>a</sup> $\pm$ 3.4	799.1 <sup>c</sup> $\pm$ 632.6	76.6 <sup>c</sup> $\pm$ 46.3	65.7 <sup>a</sup> $\pm$ 21.9	73.9 <sup>a</sup> $\pm$ 17.3	0.120 <sup>ab</sup> $\pm$ 0.13	0.021 <sup>a</sup> $\pm$ 0.03	0.014 <sup>a</sup> $\pm$ 0.01
	WR	-19.8 <sup>a</sup> $\pm$ 3.0	-18.3 <sup>a</sup> $\pm$ 3.5	-24.7 <sup>b</sup> $\pm$ 3.3	-27.5 <sup>a</sup> $\pm$ 3.4	229.4 <sup>a</sup> $\pm$ 89.6	28.7 <sup>a</sup> $\pm$ 10.7	71.6 <sup>ab</sup> $\pm$ 22.4	78.7 <sup>a</sup> $\pm$ 15.6	0.026 <sup>a</sup> $\pm$ 0.03	0.040 <sup>a</sup> $\pm$ 0.04	0.009 <sup>a</sup> $\pm$ 0.01
CIP390637.1	CT	-25.0 <sup>a</sup> $\pm$ 3.0	-20.8 <sup>bc</sup> $\pm$ 2.4	-20.2 <sup>a</sup> $\pm$ 3.4	-25.2 <sup>a</sup> $\pm$ 2.9	832.1 <sup>bc</sup> $\pm$ 613.0	70.8 <sup>c</sup> $\pm$ 43.7	61.0 <sup>a</sup> $\pm$ 18.4	68.8 <sup>a</sup> $\pm$ 17.1	0.154 <sup>a</sup> $\pm$ 0.22	0.018 <sup>a</sup> $\pm$ 0.02	0.049 <sup>a</sup> $\pm$ 0.03
	WR	-23.7 <sup>a</sup> $\pm$ 4.2	-19.9 <sup>a</sup> $\pm$ 1.3	-21.6 <sup>b</sup> $\pm$ 4.1	-24.7 <sup>a</sup> $\pm$ 4.0	227.8 <sup>a</sup> $\pm$ 99.6	29.1 <sup>a</sup> $\pm$ 17.7	72.5 <sup>a</sup> $\pm$ 22.2	74.4 <sup>a</sup> $\pm$ 18.9	0.028 <sup>a</sup> $\pm$ 0.04	0.034 <sup>a</sup> $\pm$ 0.03	0.031 <sup>a</sup> $\pm$ 0.04
LSD values		4.06	2.88	2.15	3.36	206.92	17.1	7.73	9.61	0.072	0.024	0.037

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4 492 **Figure Legends**  
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9 494 **Figure 1.** PCA analysis of leaf, tuber and root at 30 and 70 days of the water  
10 restriction and control treatment. Score plot was based on the metabolite  
11 composition of leaf (A,B), tuber (C) and root (D) of five potato genotypes at 30  
12 (A) and 70 (B-D) days under control treatment (CT, grey) and water restriction  
13 (WR, white). Metabolite variables were obtained from untargeted analysis of the  
14 polar extracts.  
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26 501 **Figure 2.** Heatmap displaying significant differences of metabolite levels in leaf  
27 (30 and 70 days) of five potato genotypes under water restriction compared to  
28 control treatment conditions. Metabolites were determined from targeted  
29 analysis of polar and non-polar extracts. Fold changes are indicated as  
30 decrease (red), no change (grey) and increase (blue). Fold changes higher than  
31 2.5 are indicated as blue with black dots.  
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43 508 **Figure 3.** Pathway display of significant changes in metabolite levels in tubers  
44 of five potato genotypes under water restriction compared to control treatment  
45 conditions. Metabolites were determined from targeted analysis of polar  
46 extracts. Fold changes are indicated as decrease (red), no change (grey) and  
47 increase (blue). Metabolites not detected in the targeted analysis are indicated  
48 in white.  
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4 515 **Supplementary data**

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9 517 **Supplementary Fig. 1.** Drought tolerant index (DTI) of tuber yield (A) of five  
10 genotypes and PCA displayed as biplot (B) of control treatment (CT, grey) and  
11 518 water restriction (WR, white) treatments including scores (genotypes) and  
12 519 loadings (physiological measurements). Physiological traits were measured over  
13 520 the whole duration of the experiment.  
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23 523 **Supplementary Fig. 2.** PCA to assess metabolite composition of leaf of five  
24 potato genotypes at 30 (A,B) and 70 (C,D) days under control treatment (CT;  
25 A,C) and water restriction (WR; B,D) treatment. Metabolites variables were  
26 524 obtained from untargeted analysis of polar extracts.  
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35 528 **Supplementary File 1.** Effect of treatment, genotypes, and their interaction on  
36 physiological parameters measured in 5 breeding clones at different times  
37 529 points Values are ANOVA Mean square and P -values are shown.  
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45 532 **Supplementary File 2.** Data of untargeted analysis of leaf, root and tuber  
46 material under control treatment (CT) and water restriction (WR) conditions at  
47 533 30 and 70 DAWRI.  
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54 536 **Supplementary File 3.** Average ratios and *P*-Values of targeted analysis under  
55 water restriction (WR) conditions compared to the control (CT).  
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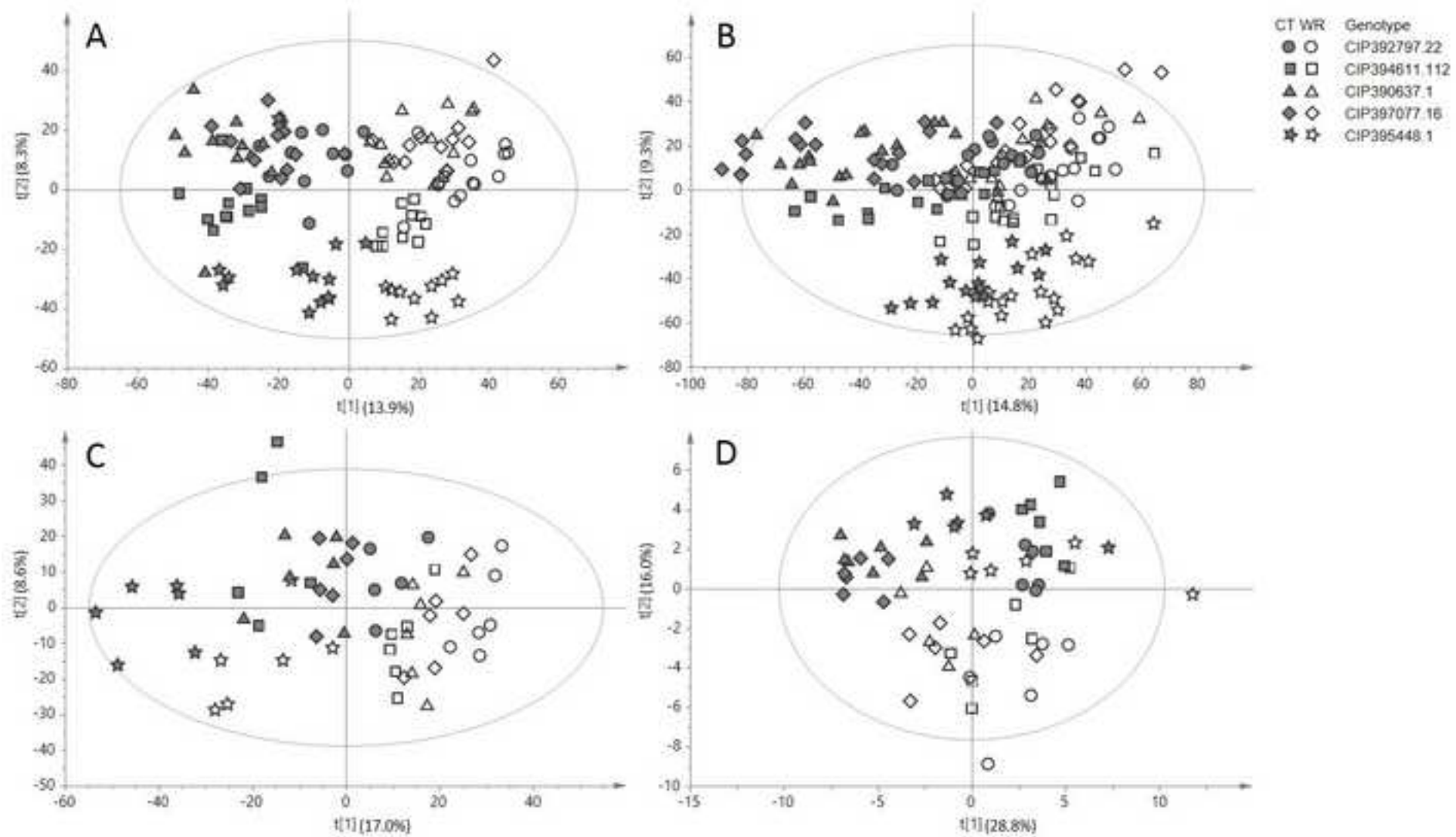
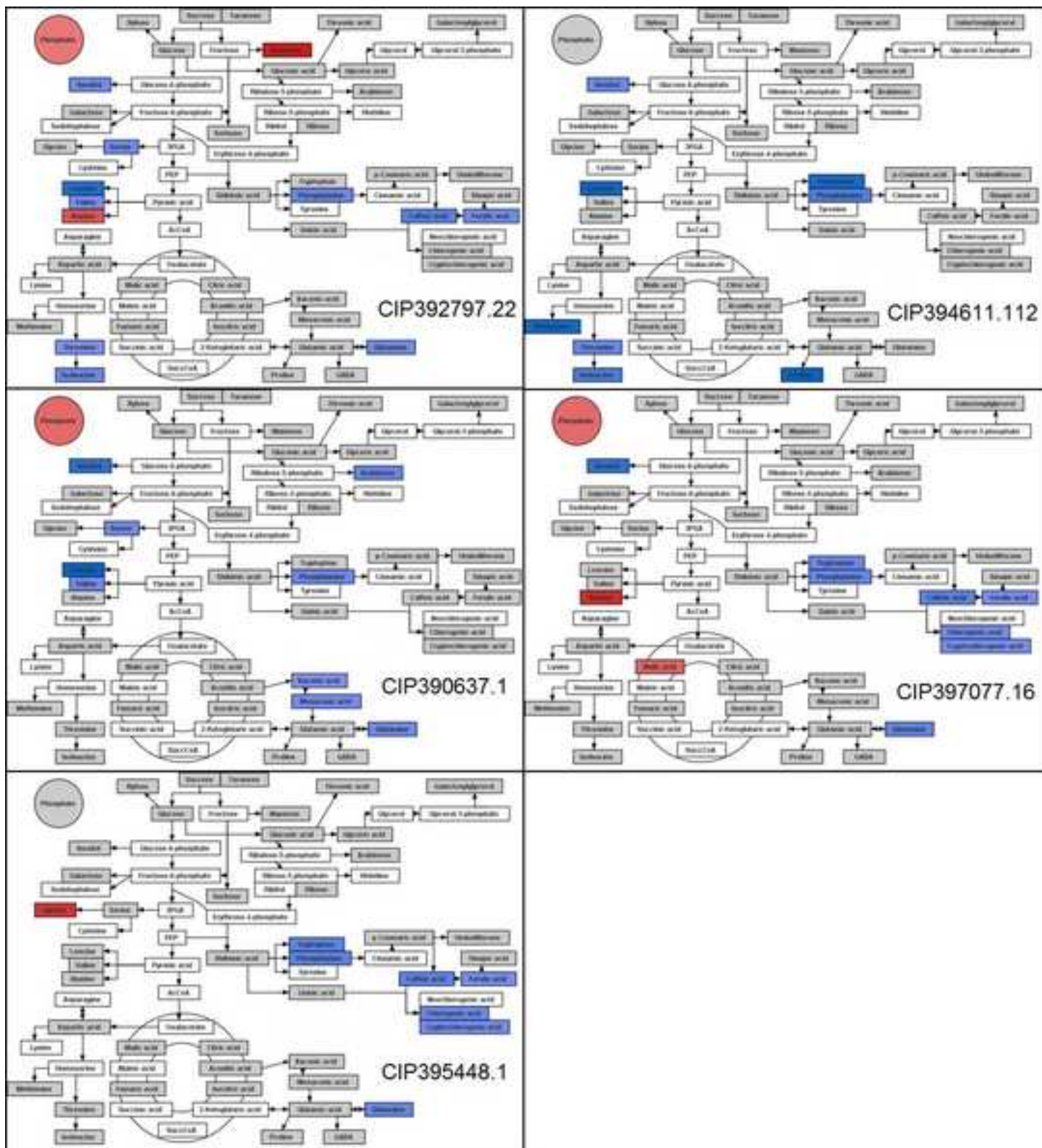




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