



Title	Suppression of red color development associated with anthocyanin accumulation in the epicarp of grape (<i>Vitis labrusca</i> x <i>vinifera</i> cv. Ruby Roman) berries caused by air temperature in daylight periods higher than 33 degrees C during maturation
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1 **Suppression of red color development associated with anthocyanin accumulation**
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3 **by air temperature in daylight periods higher than 33°C during maturation**

4

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18

19 **Abstract**

20 The effect of high air temperature on the anthocyanin accumulation in the epicarp
21 of 'Ruby Roman' grape (*Vitis labruscana* × *vinifera*) berries was examined. Seventeen
22 types of anthocyanin were detected from the epicarp tissue of the matured berries using
23 HPLC. Since a strong positive correlation ($P < 0.005$, $R^2 = 0.937$) was confirmed
24 between the degree of color density evaluated using a color chart and the total content
25 of anthocyanins quantified separately, it seems possible to trace anthocyanin
26 accumulation using the color chart on each berry during maturation at different air
27 temperatures. Four conditions were designed for cooling the fruit clusters with a
28 spot-cooling system, which can keep air temperature 4–5°C lower for 60–70 days after
29 the full-bloom, i.e. consecutive (0:00–24:00) cooling, daytime (6:00–18:00) cooling,
30 nighttime (18:00–6:00) cooling and no-cooling (control). The degree of color density
31 showed statistically higher ($P < 0.05$) values in the consecutive-cooling treatment than
32 that of the control through the post-cooling duration (0–15 days after cooling). To
33 clarify the effect of temperature zones with or without daylight on the anthocyanin
34 accumulation in the epicarp, percentage of exposure time to each temperature zone of
35 < 21 , 21–24, 24–27, 27–30, 30–33, 33–36 and $36^\circ\text{C} \leq$ were calculated for both daylight
36 and dark periods, separately. Correlations of the Δ time % (the amount of difference in
37 percentage of exposure time from the control) with the Δ color degree (the amount of
38 difference in degree of color density from the control) in the daylight period were
39 positive at temperature zones of $< 30^\circ\text{C}$, but turned to negative at those of $30^\circ\text{C} \leq$. The
40 fact that correlation coefficient was statistically significant ($P < 0.05$) at the
41 temperature zones of 27–30°C (positive) and 33–36°C (negative) in the daylight period
42 shows that the threshold where the anthocyanin accumulation starts to be suppressed
43 exists within 30–33°C.

44

45 Keywords

46 Color chart; degree of color density; fruit maturation; HPLC; spot-cooling system.

47

48 **1. Introduction**

49 The 'Ruby Roman' grape (*Vitis labruscana* × *vinifera*) is a brand–new tetraploid
50 cultivar bred and registered in 2007 by Ishikawa prefecture, Japan, known as a progeny
51 of 'Fujiminori' which was open-pollinated in 1994 (Shima et al., 2006). The berries of
52 the cultivar have red color on the epicarp and are consumed mainly as sophisticated
53 fresh fruits. Thus, the epicarp color of the fruit is one of the cultivar's most important
54 qualities. Accumulation of red color in the epicarp commonly starts during the
55 veraison period, 40–50 days after the full–bloom, when the young hard fruits turn soft
56 and begin to enlarge. On the production of grapes with red or black epicarp color, it
57 has been pointed out that high air temperature during the veraison or later causes
58 insufficient color development of the epicarp (Winkler, 1962), which is related to the
59 suppression of anthocyanin accumulation (Kliewer, 1970; Mori et al., 2004; Yamane et
60 al., 2006). In this case, the accumulation of anthocyanins was strongly affected by the
61 local temperature of atmosphere around berries (Tomana et al., 1979; Koshita et al.,
62 2007) and cultivars (Kliewer and Torres, 1972). Recently, Gao-Takai et al. (2019)
63 clearly demonstrated that the accumulation and component of anthocyanins, and gene
64 expression related to the anthocyanin biosynthesis in the epicarp of grape berries
65 depended on the temperature. Shinomiya et al. (2015) reported that color development
66 of the epicarp of 'Kyoho' (*V. labrusca* × *vinifera*) became poor when the average daily
67 air temperature exceeded 27°C. However, details on the relation of high temperature,
68 such as threshold of temperature, duration of exposure time, with or without light, and
69 so on, had not yet been clarified.

70 With anthocyanin analysis using chemical methods, plant tissues must be
71 homogenized during the extraction procedure. This fact indicates that the process of
72 anthocyanin accumulation into the epicarp is untraceable using the same berries
73 attached on a vine by utilizing chemical methods. Matsuda et al. (2020a) showed that a

74 positive linear correlation was observed between the degree of color density, evaluated
75 using a color chart created for the 'Ruby Roman' berries specifically, and the content of
76 anthocyanins, quantified as cyanidin 3–glucoside equivalent using colorimetric
77 determination. The color chart seems viable for tracing anthocyanin accumulation
78 using the same berries on a vine without sampling. Furthermore, since there are
79 different kinds of anthocyanin molecules contained in the epicarp tissue of grape
80 berries (Li et al., 2013), colorimetric quantification might be too rough. So, we tried to
81 quantify anthocyanins separately using HPLC in this study, and compared the degree
82 of color density with the total content of anthocyanins, so that we can confirm if degree
83 of color density is viable for tracing anthocyanin accumulation in the epicarp of the
84 berries maturing in different conditions.

85 Another problem in investigating the effect of air temperature on anthocyanin
86 accumulation is that it is very difficult to control air temperature around the berries
87 accurately on the vines grown in a plastic greenhouse. To resolve this matter we
88 employed a spot–cooling system which can keep air temperature around the berries at
89 about 4–5°C lower than the outside air temperature. Tarara et al. (2008) also tried to
90 control air temperature around berry clusters of 'Merlot' grape using a forced convection
91 system to produce a dynamic range of berry temperatures under field conditions in both
92 sun-exposed and shaded fruit. We arranged three treatments on air temperature
93 associated with day/night cooling, and compared the color development on cooled
94 berries with that of the control (without cooling). Furthermore, the amount of
95 difference in both exposure time to a specific air temperature zone and the degree of
96 color density against those of the control were calculated in each treatment, and then
97 correlation of these two factors was used to evaluate the effect of air temperature on
98 the accumulation of red color (anthocyanins) in the epicarp.

99

100 2. Materials and Methods

101 2.1. Quantification of anthocyanins using HPLC

102 The berries of 'Ruby Roman' grape categorized into 1–10 degrees of color
103 density using a specific color chart (Matsuda *et al.*, 2020a) were randomly collected in
104 2014 from 8–year–old vines which had been grown in a rain–protected plastic
105 greenhouse without side covers at Ishikawa Agriculture and Forestry Research Center,
106 Kahoku, Ishikawa, Japan. In this case, three berries with the same degree were used for
107 anthocyanin analyses, respectively. An epicarp disk (8 mm in diameter) of the berry
108 was punched out at around the top (style side) of the berry using a cork borer,
109 homogenized using mortar and pestle with 1 mL of 50%(v/v) acetic acid, incubated for
110 12 h at 4°C in the dark, centrifuged at $10,000 \times g$, and then supernatant was filtrated
111 through a polyvinylidene difluoride filter (0.45 μm in pore size, EMD Millipore,
112 Darmstadt, Germany). Anthocyanins were analyzed using a High Performance Liquid
113 Chromatography (HPLC) system (Prominence LC20A instrument, Shimadzu, Kyoto,
114 Japan) as was described by Katayama–Ikegami *et al.* (2016). The conditions for HPLC
115 were as follows: mobile phase, linear gradient analysis of 1.5%(v/v) phosphoric acid
116 (solvent A) and 1.5%(v/v) phosphoric acid, 20%(v/v) acetic acid and 25%(v/v)
117 acetonitrile (solvent B); pump, LC–20AD (Shimadzu); column, Inertsil[®] ODS–2
118 analytical (250 \times 6.0 mm, 5 μm in silica gel particle size; GL Sciences, Tokyo, Japan);
119 temperature, 35°C (CTO–20A column oven; Shimadzu); detector, UV–VIS
120 (SPD–20AV, Shimadzu); absorbance of wave length, 520 nm (A_{520}); flow rate, 0.8 mL
121 min^{-1} ; sample volume, 10 μL . The gradients of solvent A/B were as follows: 0–40 min,
122 linear gradient from 75/25 to 15/85; 40–45 min, 75/25. Depending on the peak area
123 monitored at A_{520} , each anthocyanin of mono glucoside was quantified using an
124 external standard curve for cyanidin 3–glucoside (Cy3G), peonidin 3–glucoside
125 (Pn3G), delphinidin 3–glucoside (Dp3G), petunidin 3–glucoside (Pt3G) and malvidin

126 3–glucoside (Mv3G), respectively. The other anthocyanins of cyanidin 3,5–diglucoside
127 (Cy3,5G), delphinidin 3–(*p*-coumaroyl glucoside)–5–glucoside (Dp3pG5G), cyanidin
128 3–(*p*-coumaroyl glucoside)–5–glucoside (Cy3pG5G), petunidin 3–(*p*-coumaroyl
129 glucoside)–5–glucoside (Pt3pG5G), delphinidin 3–*p*-coumaroyl glucoside (Dp3pG),
130 peonidin 3–(*p*-coumaroyl glucoside)–5–glucoside (Pn3pG5G), malvidin
131 3–(*p*-coumaroyl glucoside)–5–glucoside (Mv3pG5G), cyanidin 3–*p*-coumaroyl
132 glucoside (Cy3pG), petunidin 3–*p*-coumaroyl glucoside (Pt3pG), peonidin
133 3–*p*-coumaroyl glucoside (Pn3pG), malvidin 3–*p*-coumaroyl glucoside (Mv3pG), and
134 malvidin 3–acetylglucoside (Mv3aG) were identified by comparing the retention time
135 and elution order with those described by Shiraishi et al. (2007), then quantified as mg
136 of mono glucoside equivalent which have the same aglycone (anthocyanidin) using the
137 standard curves for Cy3G, Pn3G, Dp3G, Pt3G and Mv3G, since no authentic sample
138 was available. Total content of anthocyanins was calculated by adding all of them.

139

140 2.2. *Procedures on fruit growing*

141 Experiments were performed in 2016 using two vines of 10-year-old ‘Ruby
142 Roman’ grown in a rain-protected plastic greenhouse without side covers at Ishikawa
143 Agriculture and Forestry Research Center. In this case, the roof of the house was
144 covered in April 2, and the date of the full-bloom and the onset of veraison was May
145 20 and June 30, respectively. To induce artificial parthenocarpy and make berries
146 seedless, all the fruit clusters at the full-bloom were dipped in a solution containing 25
147 ppm of gibberellic acid (GA₃) supplemented with 5 ppm of forchlorfenuron
148 (1-(2-chloro-4-pyridyl)-3-phenylurea, CPPU), then dipped again in the same GA₃
149 solution without CPPU eleven days after the full-bloom. A few days after the second
150 GA₃ treatment, some fruit clusters were thinned to bring the leaf/cluster ratio closer to
151 90, i.e. 3 shoots/cluster. This shoots/cluster ratio followed the guideline for growers of

152 ‘Ruby Roman’ vines, which was set by Ishikawa prefecture government since high
153 quality berries must be produced. In addition, some berries in a cluster were also
154 thinned to set the number of berries as 25/cluster. Then, all the fruit clusters were
155 covered with a BIKOO® perforated plastic bag (30 × 40 cm, Nidaiki Corp., Komaki,
156 Japan) with numerous 0.1 mm pores, attached with a paper umbrella for shading on the
157 top of each fruit cluster. This transparent bag can help picking off the cracked-berries
158 from fruit cluster and confirming color development on the epicarp of berries.
159 Perforation is for preventing temperature rise in the bag. In this case, 13% of light
160 intensity was reduced by the bag which was determined using an illuminometer
161 (RTR-574, T&D, Matsumoto, Japan). Other cultivation managements like fertilization,
162 sprinkling and pest control followed the practice. The above-mentioned procedures are
163 very common in the commercial production of ‘Ruby Roman’ berries.

164

165 2.3. *Utilization of a spot-cooling system to decrease air temperature around fruit* 166 *clusters*

167 The spot-cooling system (utility model registration number 3204876, Japan Patent
168 Office) employed in this study is composed of three units: a spot-cooling apparatus
169 (SUASP1DS, Daikin, Osaka, Japan) equipped with 2.4 kW cooler, which can decrease
170 the blowing air temperature 9.4°C lower than that of the suction air at a flow rate of 6.6
171 m³ min⁻¹; clear plastic bags covering the fruit cluster but not closed; thermal insulated
172 pipes and tubes connecting the apparatus and bags (Fig. 1). In this case, 60 fruit bunches
173 in maximum can be cooled down to 4–5°C lower than the outside temperature by the
174 system, simultaneously.

175 Four experimental groups were designed on cooling the fruit clusters: consecutive
176 (0:00–24:00) cooling; daytime (6:00–18:00) cooling; nighttime (18:00–6:00) cooling;
177 no-cooling (control). Duration of the cooling treatment was from July 19 to July 29

178 (from 60 to 70 days after the full-bloom), since color development on the berries was
179 the most sensitive to air temperature during this period (Matsuda et al., 2020a). Eleven
180 fruit clusters were used for each treatment. The air temperature was monitored at the
181 center of the fruit cluster in bags using a thermometer (RTR-502, T&D, Matsumoto,
182 Japan) and recorded every 15 min. The surface color of the berries was classified into
183 1–10 degrees of color density using the color chart every 5 days from July 19 to August
184 13 (60, 65, 70, 75, 80 and 85 days after the full-bloom).

185

186 *2.4. Statistical analyses of data*

187 The content of anthocyanins and the degree of color density in the epicarp were
188 represented as average \pm SE ($n = 3$ and $n = 11$, respectively). Differences in the degree
189 of color density among cooling treatments were analyzed statistically using analysis of
190 variance (ANOVA) followed by Tukey's multiple range test.

191

192 **3. Results**

193 *3.1. Anthocyanin content in epicarp related to the degree of color density determined* 194 *using the color chart*

195 An example high performance liquid chromatogram of the anthocyanin analysis
196 was shown in Fig. 2. This sample was extracted from epicarp of matured berries,
197 whose color was classified into the degree 9 in density using the color chart. Seventeen
198 anthocyanin types were confirmed, which was the maximum number in the 'Ruby
199 Roman' berries examined. They were composed of 4 cyanidin types, 3 peonidin types,
200 3 delphinidin types, 3 petunidin types and 4 malvidin types (Table 1). The total
201 anthocyanin content per epicarp area was calculated as $33.92 \mu\text{g cm}^{-2}$. Among
202 anthocyanins, the greatest content ($6.87 \mu\text{g cm}^{-2}$) of Cy3G occupied 20.3% of the total,
203 the second greatest ($5.72 \mu\text{g cm}^{-2}$) of Cy3pG occupied 16.9%, and the third greatest

204 (3.26 $\mu\text{g cm}^{-2}$) of Pn3G occupied 9.6%. Similarly, the total anthocyanin content in
205 epicarp of the berries whose color was classified into 1–10 degrees in density was
206 determined by the same manner with three replications, respectively. In this case,
207 correspondence of the color degree to the fruit developmental stage of berries is as
208 follows: degree 1, second stage; degrees 2–7, third stage; degrees 8–10, harvest stage.
209 Of course, the detected types of anthocyanins were smaller than 17 in berries with poor
210 color density (Suppl. 1). Then, the total anthocyanin content was compared with the
211 degree of color density. As a result, a strong positive correlation ($P < 0.005$, $R^2 =$
212 0.937) was confirmed between both factors (Fig. 3). The results show that the
213 anthocyanin accumulation in epicarp of a berry can be monitored using the color chart
214 exactly.

215

216 3.2. *Effect of cooling berries on red color development (anthocyanin accumulation)* 217 *in the epicarp*

218 Effects of the cooling treatments, i.e. consecutive–cooling, daytime–cooling and
219 nighttime–cooling, on red color development in the epicarp were shown in Fig. 4. In
220 all cooling treatments, the degree of color density increased gradually along with the
221 maturing of berries through and after the cooling period. Furthermore, the degree of
222 color density in consecutive–cooling treatment was statistically greater ($P < 0.05$) than
223 that of the control (without cooling) from 0 through 15 days after the cooling period.
224 The curves of both the daytime–cooling and the nighttime–cooling treatments existed
225 between the curves of the consecutive–cooling treatment and the control, but not
226 significantly different from the control.

227 During the cooling period (July 19–29), sunrise and sunset were at around 5:00 and
228 19:15, respectively. To evaluate the effect of the air temperature in more detail on the
229 red color development with or without daylight, we employed a concept of temperature

230 zone composed of 7 zones (< 21 , 21–24, 24–27, 27–30, 30–33, 33–36 and $36^{\circ}\text{C} \leq$) of
231 air temperature with 3°C interval basically in this experiment. Then, percentage of
232 exposure time when the monitoring fruit cluster had been exposed to each temperature
233 zone to total time during the experiment were calculated for daylight (5:00–19:15) and
234 dark (19:15–5:00) period, separately (Fig. 5). It must be noticed that daylight and dark
235 periods did not correspond to the periods of daytime– and nighttime–cooling,
236 respectively. Furthermore, the amount of difference in the percentage of exposure time
237 to each temperature zone from that of the control (Δ time %) were calculated in each
238 cooling treatment. Similarly, the amount of difference in average degree of color
239 density (mean value from 0 through 15 days after the cooling period) in each treatment
240 from that of the control (Δ color degree) were calculated. To clarify the effect of the
241 temperature zones on the red color development (anthocyanin accumulation) in the
242 epicarp, correlations of the Δ time % with the Δ color degree at each temperature zone
243 were shown in Fig. 6. During the daylight period, the relationship between the Δ
244 time % and the Δ color degree was linearly positive at the temperature zones less than
245 30°C , but turned negative at the temperature zones greater than 30°C . In this case,
246 correlation coefficient was statistically significant ($P < 0.05$) at temperature zones of
247 27– 30°C and 33– 36°C . By contrast, no certain direction was confirmed between these
248 two factors during the dark period.

249

250 **4. Discussion**

251 The color density of the epicarp in red/black–type grape berries would be affected
252 mainly by the component, kinds and the amount, of anthocyanins and pH in the tissues.
253 The anthocyanin analyses using HPLC in this study revealed that the major kinds were
254 Cy3G and Cy3pG, the cyanidin type both with red color, the sum of which occupied
255 37.2% of the total anthocyanin content. When compared with the other cultivars, the

256 anthocyanin component in the matured ‘Ruby Roman’ epicarp (Table. 1) was similar
257 to that of ‘Steuben’ with the high content of Cy3G, Cy3pG, Dp3G and Dp3pG, but
258 containing some Pn3G and Pn3pG in contrast to ‘Steuben’ (Shiraishi et al., 2007).
259 Furthermore, the ‘Ruby Roman’ epicarp also contained a certain level of Pt3G, Pt3pG,
260 Mv3G and Mv3pG. Thus, the red color of ‘Ruby Roman’ berries seems to be due to
261 this specific composition of anthocyanins accumulated in the epicarp.

262 On the anthocyanin content related to the degree of color density in the ‘Ruby
263 Roman’ epicarp, the correlation coefficient ($R^2 = 0.937$) in the present study was
264 higher than that ($R^2 = 0.745$) in the previous study (Matsuda et al., 2020a) where the
265 anthocyanin content was quantified by colorimetric method. Since anthocyanins were
266 quantified more precisely using HPLC, which showed a strong correlation with the
267 degree of color density (Fig. 3), it seems that monitoring the accumulation of
268 anthocyanins in the ‘Ruby Roman’ epicarp using the color chart is quite exact. Indeed,
269 the color chart has also been employed for estimating anthocyanin content in the
270 epicarp of apples (Marsh et al., 1996), raspberries (Stavang et al., 2015) and grapes
271 such as ‘Kyoho’, ‘Pione’ and ‘Suzuka’ (Sugiura et al., 2018; Yamazaki and Suzuki,
272 1980), even if correlation coefficients were smaller than that of the present study.

273 It is well known that light intensity play an important role on the anthocyanin
274 accumulation in the epicarp of grape berries. Matsuda et al. (2020b) demonstrated
275 utilizing *in vitro*-cultured ‘Ruby Roman’ berries that the anthocyanin accumulation
276 occurred only when artificial light with luminous intensity of 5,000 lx or more was
277 irradiated from fluorescent tube. In the present study, the light intensity in the green
278 house was kept at more than 5,000 lx from 1h after sunrise to 1h before sunset during
279 the experiment (Suppl. 2). However, we will need to have deeper discussion about the
280 relationship between light intensity and the anthocyanin accumulation.

281 The roles of temperature on the anthocyanin accumulation in grape berries have
282 been discussed in relation to day/night temperature. The discussions were categorized
283 into 4 groups as the accumulation was accelerated by: lower temperature in daytime
284 (Kliewer, 1970); lower temperature in nighttime (Koshita et al., 2007); lower daily
285 average temperature (Shinomiya et al., 2015; Sugiura et al., 2018); the difference
286 between day/night average temperature (Tomana et al., 1979). In the present study, the
287 degree of color density showed statistically higher values in the consecutive-cooling
288 treatment than that of the control through the post-cooling duration (0–15 days after
289 cooling), but the effect of daytime- or nighttime-cooling alone was not clear (Fig. 4).
290 However, it was clearly demonstrated that the relationship between the Δ time % and
291 the Δ color degree was positive at less than 30°C, but negative at greater than 30°C
292 during the daylight period. Thus, our results indicate that high air temperature (30°C \leq)
293 during the daylight period would suppress the anthocyanin accumulation in the epicarp
294 of ‘Ruby Roman’ berries.

295 The concept of temperature zone was employed in this study, since we thought
296 that anthocyanin biosynthesis related to some enzymatic reactions might have the
297 optimum temperature. Another possibility of concerning air temperature is based on
298 the concept of accumulated temperature during the treatments, which was not a divided
299 section like a zone but whole. From this point of view, the average air temperature was
300 calculated from the temperature data (570 and 390 data for daylight and dark period,
301 respectively) determined every 15 min for each cooling treatment, and correlation
302 between the average values and those of the Δ color degree was examined (Table 2).
303 As a result, the correlation coefficient was statistically significant ($P < 0.05$) only in
304 the case of daylight period. This fact clearly indicates the involvement of daylight on
305 the anthocyanin accumulation in the grape epicarp, which was mentioned previously
306 by many researchers (Jeong et al., 2004; Pereira et al., 2006; Matus et al., 2009;

307 Azuma et al., 2012; Koyama et al., 2012). In this case, it was also demonstrated that
308 effect of the difference in the average air temperature between daylight and dark period
309 on the anthocyanin accumulation was not significant.

310 On the critical temperature which suppresses anthocyanin accumulation, it has
311 been pointed out that high temperature during daytime inhibits anthocyanin
312 accumulation in maturing berries (Kobayashi et al., 1967; Kliewer, 1970; Mori et al.,
313 2004). However, since these studies were performed at a constant daytime temperature
314 of 30°C or less set by using a controlled climate room, they couldn't clarify the
315 threshold temperature. The facts that the correlation between the Δ time % and the Δ
316 color degree turned from positive to negative at 30°C, and furthermore the correlation
317 coefficient was statistically significant ($P < 0.05$) at the temperature zones of 27–30°C
318 (positive) and 33–36°C (negative) indicate that the critical point (threshold) where the
319 anthocyanin accumulation starts to be suppressed may exist within 30–33°C (the
320 transitional temperature zone). It is concluded that the anthocyanin accumulation
321 would be suppressed strongly by air temperature more than 33°C during the daylight
322 period. Thus, it is suggested that keeping the air temperature around the fruit clusters in
323 veraison or later at $< 30^\circ\text{C}$ during daylight period is indispensable for fully developing
324 red color on the epicarp of 'Ruby Roman' berries.

325

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417 Japanese with English abstract).
418

419 **Legends of Figure**

420

421 Fig. 1. A picture on cooling fruit clusters of 'Ruby Roman' berries using a spot-cooling
422 system. Cold-air flow was supplied from a cooler via the thermal insulated main
423 pipe (mp), the branch pipes (bp) and the tubes (t) into the clear plastic bags (pb).
424 Length of the branch pipes and the loci where they connected with the main pipe are
425 adjustable.

426

427 Fig. 2. An example high performance liquid chromatogram of anthocyanins detected
428 from the epicarp of matured 'Ruby Roman' grape berries. Cy3,5G: cyanidin
429 3,5-diglucoside, Dp3G: delphinidin 3-glucoside, Cy3G: cyanidin 3-glucoside,
430 Pt3G: petunidin 3-glucoside, Pn3G: peonidin 3-glucoside, Mv3G: malvidin
431 3-glucoside, Dp3pG5G: delphinidin 3-(*p*-coumarylglucoside)-5-glucoside,
432 Cy3pG5G: cyanidin 3-(*p*-coumarylglucoside)-5-glucoside, Pt3pG5G: petunidin
433 3-(*p*-coumarylglucoside)-5-glucoside, Dp3pG: delphinidin
434 3-*p*-coumarylglucoside, Mv3aG: malvidin 3-acetylglucoside, Pn3pG5G: peonidin
435 3-(*p*-coumarylglucoside)-5-glucoside, Mv3pG5G: malvidin
436 3-(*p*-coumarylglucoside)-5-glucoside, Cy3pG: cyanidin 3-*p*-coumaroylglucoside,
437 Pt3pG: petunidin 3-*p*-coumaroylglucoside, Pn3pG: peonidin
438 3-*p*-coumaroylglucoside, Mv3pG: malvidin 3-*p*-coumaroylglucoside.

439

440 Fig. 3. Relationship between the degree of color density and the total anthocyanin
441 content. Values are average \pm SE ($n = 3$). *** $P < 0.005$.

442

443 Fig. 4. Effects of the cooling treatments on red color development in the epicarp of
444 maturing berries. Treatments were as follows: consecutive-cooling (open circle),

445 daytime-cooling (open triangle), nighttime-cooling (closed circle) and no-cooling
446 (control) (closed triangle). Cooling treatments were made from 60 to 70 days after
447 the full-bloom. The values represent average \pm SE ($n = 11$). Where no error bar is
448 visible SE was smaller than the symbols. The difference between the values labeled
449 with different alphabets was statistically significant ($P < 0.05$, Tukey's multiple
450 range test).

451

452 Fig. 5. Percentage of exposure time to each temperature zone associated with the
453 cooling treatments in daylight and dark period, respectively.

454

455 Fig. 6. Correlations between the Δ time % and the Δ color degree in each temperature
456 zone in daylight and dark period, respectively. The Δ time % and the Δ color degree
457 mean the amount of difference in percentage of exposure time and in degree of
458 color density from those of the control, respectively.

459



'Ruby Roman' grape
(*Vitis labruscana* × *vinifera*)

HPLC
17 types of anthocyanin
Total content

×



color chart

strong positive correlation
($P < 0.005$, $R^2 = 0.937$)

Correlations of the Δ time %
(difference exposure time from
the control) with the Δ color
degree

Anthocyanin accumulation

Daylight period



Promotion

Suppression



Threshold

°C 27 30 33 36

Dark period



Not be confirmed

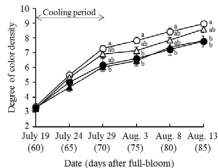
In greenhouse



Spot-cooling system

Spot-cooling system can keep air
temperature 4–5°C lower for 60–70 days
after the full-bloom

- Consecutive (0:00–24:00) cooling (○)
- Daytime (6:00–18:00) cooling (△)
- Nighttime (18:00–6:00) cooling (●)
- No-cooling (control) (▲)



Highlights

Seventeen types of anthocyanin were detected from the epicarp of matured berries

Degree of the epicarp color density was positively correlated with total anthocyanins

Fruit clusters after veraison were grown at four different air temperature conditions

Anthocyanin accumulation was suppressed strongly by $33^{\circ}\text{C} <$ during daylight period

Anthocyanin accumulation starts to be suppressed at a temperature within $30\text{--}33^{\circ}\text{C}$

Table 1. Component of anthocyanins detected from the epicarp tissue of matured 'Ruby Roman' berries.

Anthocyanins	Content ($\mu\text{g}\cdot\text{cm}^{-2}$)	(%)
Cy3G	6.871 ± 0.742^z	20.3
Cy3pG	5.723 ± 0.278	16.9
Cy3,5G	0.700 ± 0.112	2.1
Cy3pG5G	0.558 ± 0.015	1.6
Pn3G	3.261 ± 0.242	9.6
Pn3pG	2.784 ± 0.081	8.2
Pn3pG5G	1.093 ± 0.022	3.2
Dp3G	2.676 ± 0.245	7.9
Dp3pG	2.185 ± 0.181	6.4
Dp3pG5G	0.519 ± 0.051	1.5
Pt3G	1.670 ± 0.170	4.9
Pt3pG	1.239 ± 0.085	3.7
Pt3pG5G	0.795 ± 0.028	2.3
Mv3G	2.119 ± 0.154	6.3
Mv3pG	1.157 ± 0.050	3.4
Mv3aG	0.232 ± 0.005	0.7
Mv3pG5G	0.335 ± 0.019	1.0
Total	33.917	100.0

^zAverage \pm SE ($n = 3$).

Table 2. Correlation between the average air temperature during daylight and/or dark period, the difference, and the Δ color degree.

Cooling treatment	Average air temperature ($^{\circ}$ C)			Difference ($^{\circ}$ C) [A – B]
	Whole day	Daylight period [A] (5:00-19:15)	Dark period [B] (19:15-5:00)	
Consecutive	23.8	25.6	21.2	4.4
Daytime	26.1	26.7	25.2	1.5
Nighttime	25.6	29.0	20.6	8.4
No-cooling (Cont.)	28.6	30.6	25.6	5.0
<i>r</i> value vs. Δ color degree	– 0.765	– 0.956 *	– 0.212	– 0.566

‘ Δ color degree’: see Fig. 6.

‘*r* value’ means Pearson’s correlation coefficient.

* $P < 0.05$.

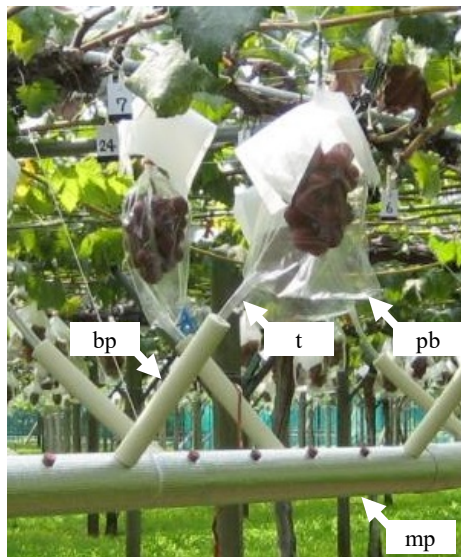


Fig. 1

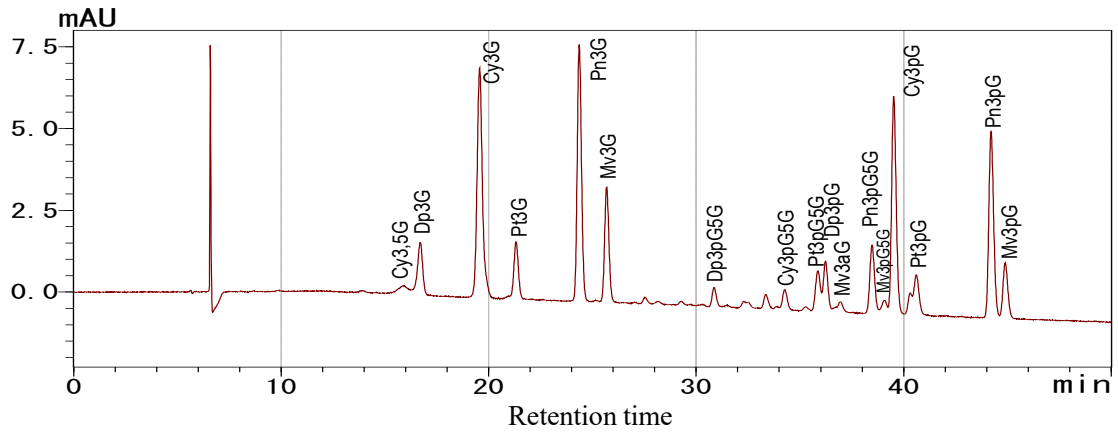


Fig. 2

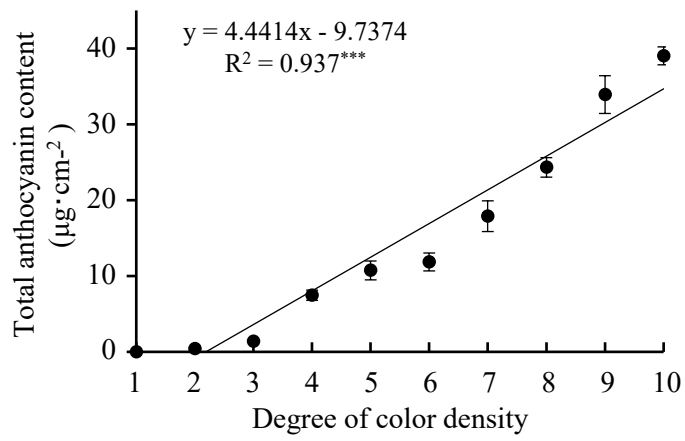


Fig. 3

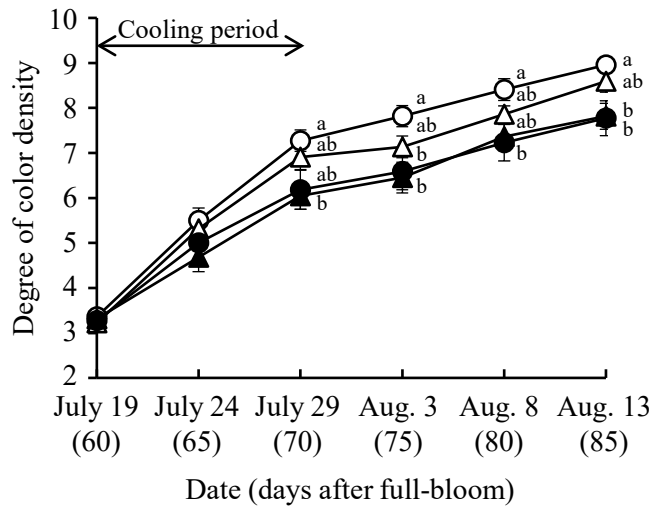


Fig. 4

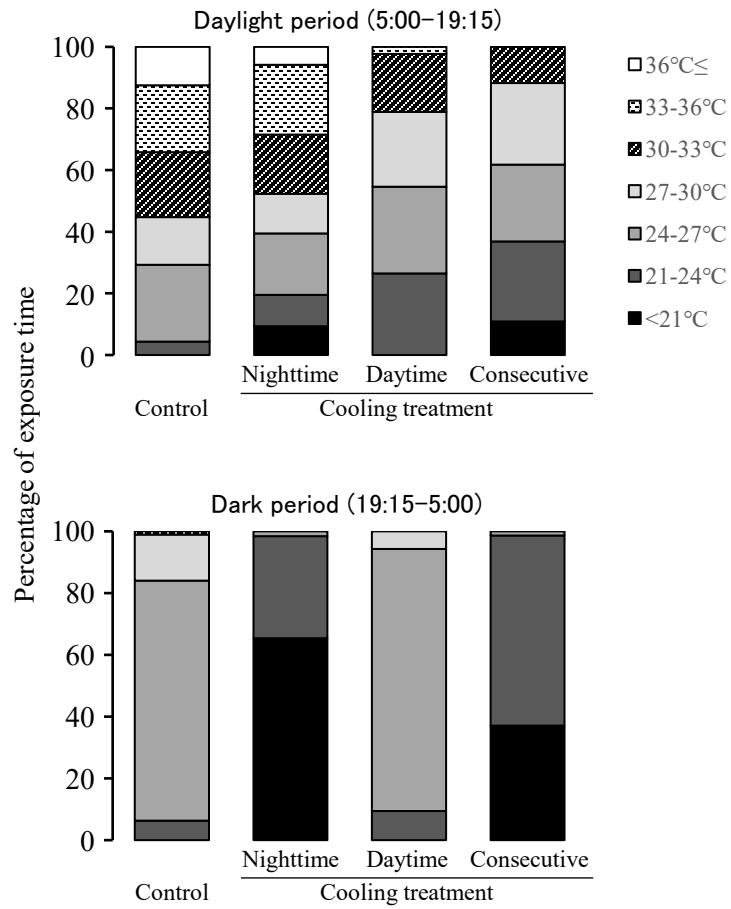


Fig. 5

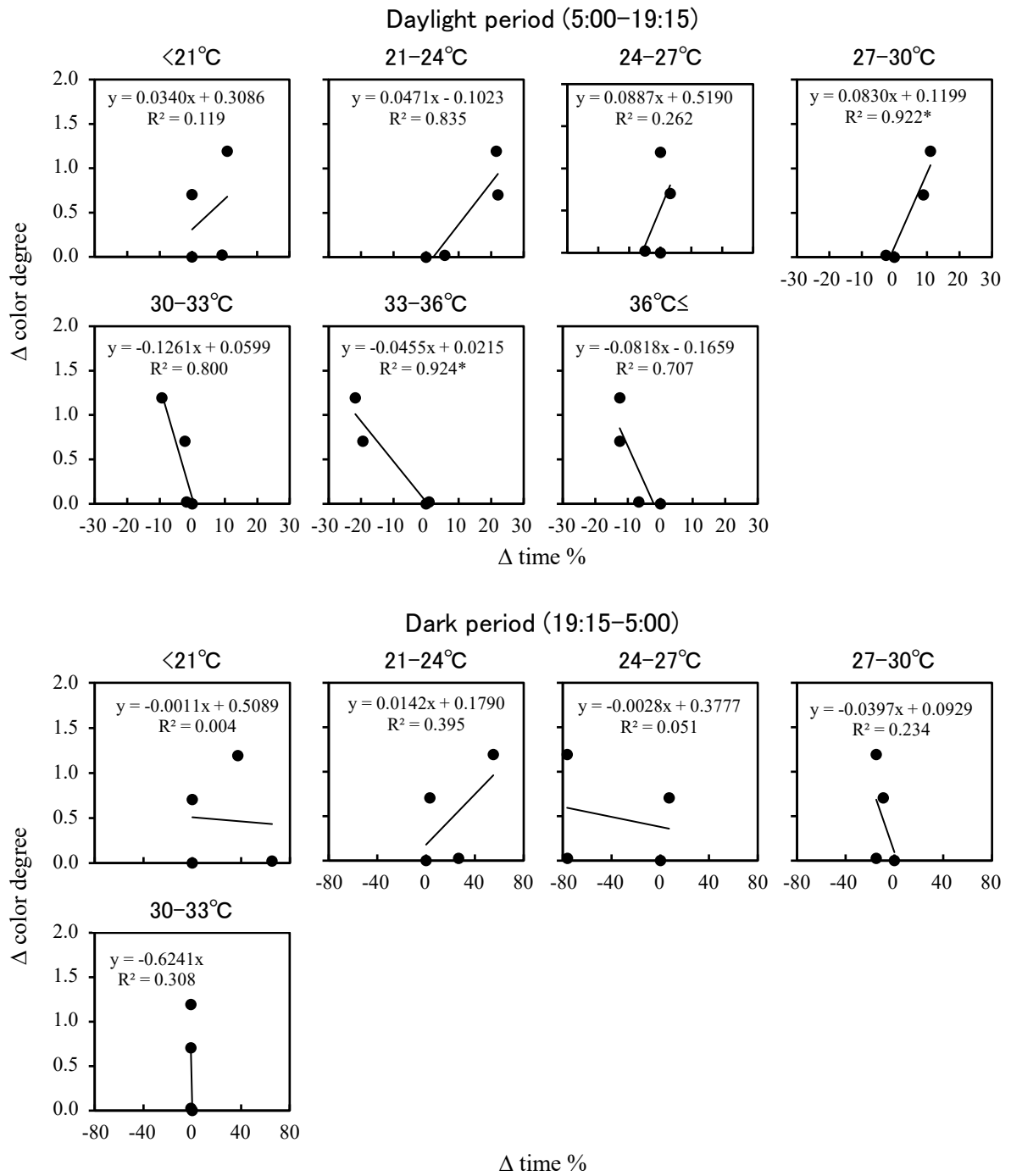


Fig. 6