Article

Chemical mechanism of petal color change in
Oenothera flowers during senescence

Yada Teppabut 1, Kin-ichi Oyama 2, Tadao Kondo 1 and Kumi Yoshida 1,2,*

1 Graduate School of Informatics, Nagoya University, Chikusa, Nagoya 464-8601, Japan;
teppabut.yada@g.mbox.nagoya-u.ac.jp (Y.T.); kondot@info.human.nagoya-u.ac.jp (T.K.)
2 Research Institute for Materials Science, Nagoya University, Chikusa, Nagoya 464-8602, Japan;
oyama@cis.nagoya-u.ac.jp
* Correspondence: yoshidak@i.nagoya-u.ac.jp; Tel.: +81-052-789-5638

Abstract: Oenothera flower petals change color during senescence. When in full bloom, the flowers
of O. tetrapetra are white and those of O. laciniata and O. stricta are yellow; however, the colors
change to pink and orange, respectively, when the petals fade. We analyzed the flavonoid
components in these petals as a function of senescence using HPLC-DAD and LC-MS. In all three
species, cyanidin 3-glucoside (Cy3G) was found in faded petals, and the content of Cy3G increased
in senescence. In full bloom (0 h), no Cy3G was detected in any of the petals, but after 12 h, the
content of Cy3G in O. tetrapetra was 0.97 µmol/gFW and that in O. laciniata was 1.82 µmol/gFW.
Together with anthocyanins, major flavonoid components in petals were identified. Quercitrin was
detected in the petals of O. tetrapetra, and isosalipurposide was found in the petals of O. laciniata
and O. stricta. The content of quercitrin did not change during senescence, but that of
isosalipurposide in O. laciniata increased from 3.4 µmol/gFW at 0 h to 4.8 µmol/gFW at 12 h. The
color change in all the three Oenothera flowers was confirmed to be due to the de novo biosynthesis
of Cy3G.

Keywords: cyanidin 3-O-glucoside; flower senescence; isosalipurposide; Oenothera; petal color
change; quercitrin.

1. Introduction

Flower color is an important characteristic for plants as it is known to be related to pollination
[1–3]. One of the many ways angiosperm plants attract pollinators is floral color changes [2,3].
Various mechanisms of color change have been reported, such as changes in pH [4,5] and losses of
pigment [6], but the most common physiological process is appearance of a pigment, especially an
anthocyanin [2].

Anthocyanins provide the widest range of colors among the three major classes of flower
pigments: anthocyanins, betalains and carotenoids [7,8]. Many studies have explored the biosynthesis
of these pigments [6–11]. In the case of anthocyanin, it is synthesized from phenylalanine, an amino
acid, via a phenylpropanoid [10–13]. The pathway starts with the synthesis of naringenin chalcone
from 4-coumaroyl-CoA and malonyl-CoA by chalcone synthase (CHS). After that, the chalcone is
converted into flavanone, dihydroflavonol and then leucoanthocyanidin [7,8,10,11]. Then,
leucoanthocyanidin is oxidized and glycosylated to afford anthocyanin [7,8,10,11,14].

A large number of plant taxa show floral color changes, and one of them is genus Oenothera,
evening primrose, which is known to undergo a flower color change during senescence. The flowers
of this genus bloom in the evening and fade in the morning. When fully opened, the petals of O.
tetrapetra are white, and then they become pink in the morning (Figure 1a). Those of O. laciniata as
well as O. stricta are yellow, and then they turn orange as they fade (Figure 1b and c). These
phenomena strongly indicate that an anthocyanin is biosynthesized during senescence. However, the
physiological process of color change in *Oenothera* has not been confirmed. We are interested in the petal color change of these flowers and studying the chemical mechanisms of such changes. Petal components were isolated, and the constituents were identified. Then, the components were quantified according to flower fading stage.

**Figure 1.** Flower color change in *Oenothera* petals during senescence. (a) *Oenothera tetraptera*; (b) *Oenothera laciniata*; and (c) *Oenothera stricta*. Scale bars: 1 cm.

2. Results

2.1 Analysis of Petal Components of *O. tetraptera*

As shown in Figure 1a, the petals of *O. tetraptera* bloom in white in the evening at approximately 21:00 and become pink after 12 h. To determine the chemical compounds responsible for the color change, the petals of *O. tetraptera* were collected at the full blooming white stage (0 h) and the faded stage (12 h), and then the petals were extracted with acidic solution (3% TFA in 50% CH₃CN aq.). Each extract was analyzed by 3D-HPLC (Figure 2). In the white petals at 0 h, 2 was the major component, and in the pink petals, which had undergone senescence, peak 1 appeared. Combined with the results of co-chromatography, the spectrum obtained from 3D-HPLC and LC-MS analysis (Figure S1) using an authentic sample, 1 was identified to be cyanidin 3-glucoside (Cy3G, Figure 3) [15]. Using the same procedure, 2 was determined to be quercitrin (quercetin 3-rhamnoside, Figure 3 and S1). This result revealed that the red color change should be due to the appearance of Cy3G during senescence.

Since the components in *O. tetraptera* petals were identified, quantitative analysis of Cy3G (1) and quercitrin (2) during senescence was carried out. The petals at 0, 4, 7 and 12 h after blooming were collected (Figure 4a), and their reflection spectra were recorded (Figure 4b). The λ_{vismax} of the colored petals was 541 nm at each stage, and the intensity at λ_{vismax} increased during flower development (Figure 4b). This corresponded with the L* value of the CIELAB color coordinate of the petals decreasing and the a* value increasing after blooming (Table 1). In addition, the pH of the
Figure 2. HPLC chromatograms of the extracts of the petals of O. tetrapeta. (a) White petals at 0 h; (b) pink petals at 12 h.

cyanidin 3-glucoside (1) quercitrin (2) isosalipurposide (3)

Figure 3. Chemical structure of the components of Oenothera petals.

Figure 4. Changes in the color and flavonoid components of the petals of O. tetrapeta during senescence. (a) Petal color at each stage; (b) reflection spectra; (c) change in the Cy3G (1) content; (d) change in the quercitrin (2) content. The data displayed are the means ± SE of three replicates (n = 3). Where no error bars are shown, the SE was too small to determine. Different letters indicate significant differences according to Tukey’s HSD test (p < 0.05).
pressed juice was measured, and no obvious changes in pH were observed during senescence (Table 1). This indicates that the color change was not due to a pH change in the petals. With extraction from each petal followed by HPLC analysis, the changes in the contents of Cy3G (1) and quercitrin (2) were quantified (Figure 4c and d). The content of Cy3G increased during flower senescence and reached its highest level (0.97 µmol/gFW) at 12 h after blooming (Figure 4c). In contrast, the content of quercitrin (2) at 0 h after blooming was 13.86 µmol/gFW, which is approximately 14 times more than the highest level of Cy3G, and the content did not significantly change during senescence (Figure 4d).

2.2 Analysis of the Components of the Petals of O. laciniata and O. stricta

Next, the same experiments were done with O. laciniata and O. stricta. These flowers are yellow at full bloom, and then turn orange during senescence (Figure 1b and c). The components of the petals of these two kinds of flowers were extracted and analyzed by 3D-HPLC (Figures 5, S2). As found in O. tetraptera, Cy3G (1) was detected at the senescence stage (Figure 5b, S2b). For structure elucidation of peak 3, the yellow petals of O. laciniata were extracted and peak 3 was isolated. Using MS (Figure S1) and NMR analysis (Figure S3-S8), 3 was identified to be isosalipurposide (chalconaringenin 2'-glucoside, 3, Figure 3) [16,17]. The same compound was detected in petals of O. stricta (Figure S2).

Because the patterns of flavonoids in O. laciniata and O. stricta were almost the same, only the petals of O. laciniata were analyzed to determine the contents of Cy3G (1) and isosalipurposide (3) during senescence. The petals at 0, 4, 8 and 12 h after blooming were collected and extracted for HPLC analysis. The contents of both Cy3G (1) and isosalipurposide (3) were quantified over the course of 12 h after blooming (Figure 6). During senescence, the contents of both 1 and 3 increased with similar significant differences, and the highest contents of the compounds were 1.82 µmol/gFW for Cy3G (1) and 4.83 µmol/gFW for isosalipurposide (3) at 12 h after blooming (Figure 6).

3. Discussion

In this report, the flowers of Oenothera during color change were chemically analyzed. In all three Oenothera species, Cy3G (1) was present in faded flowers, yet no Cy3G was detected at full bloom (0 h). From the quantitative analysis of the flavonoids during flower senescence, increases in Cy3G (1) in both O. tetraptera and O. laciniata petals were observed. This corresponded to the color parameters and the UV/Vis absorption spectra. Therefore, it was concluded that the color change should be due to the de novo synthesis of Cy3G.

Together with anthocyanin, we analyzed the flavonoid components and found that a high level of a flavonol, quercitin (2), was present in white petals of O. tetraptera. The molar ratio of 2 to 1 at 12 h after blooming was more than 13 to 1. The pH of the pressed juice of the O. tetraptera petals was approximately 5.5 (Table 1). At this pH, simple anthocyanins such as Cy3G are not stable, and they are easily hydrated to give colorless pseudobases. However, the high content of quercitin (2) in the petals might stabilize the color of Cy3G by exhibiting a co-pigment effect. On the other hand, the yellow petals of O. laciniata and O. stricta contained a glycosylchalcone, isosalipurposide (3). At 12 h, the molar ratio of 3 to 1 was approximately 2.5 to 1. These results correlated with previous reports on the flavonoid distribution in Oenothera [1,18-20]. In O. laciniata and O. stricta, the orange color in faded
Figure 5. HPLC chromatograms of the extracts of the petals of O. laciniata. (a) Yellow petals at 0 h; (b) orange petals at 12 h.

Figure 6. Changes in color and flavonoid components of the petals of O. laciniata during senescence. (a) Petal color at each stage; (b) change in the Cy3G (1) content; (c) change in the isosalipurposide (3) content. The data shown are the means ± SE of three replicates (n = 3). Where no error bars are shown, the SE was too small to determine. Different letters indicate significant differences according to Tukey’s HSD test (p < 0.05).

petals is developed by mixing yellow chalcone 3 with red Cy3G (1) [16,21,22]. In these petals, Cy3G might also be stabilized with isosalipurposide (3) and other co-existing polyphenolic compounds (Figure 5b).

According to the well-established flavonoid biosynthetic process [8–11,23], anthocyanins and flavonols are produced via a divergent pathway. Dihydroflavonol is the common precursor in the synthesis of both anthocyanidins by dihydroflavonol reductase (DFR) and flavonols by flavonol
4. Materials and Methods

4.1. Plant Materials

The *O. tetraperta* flowers used in this experiment were obtained from the Kochi Prefectural Makino Botanical Garden. The flower buds of *O. tetraperta* were cut, kept in box and transported to Nagoya University within 1 day. Then, flower buds were incubated in a plant growth chamber (14 h-light/10 h-dark cycle) at 25 °C (light) and 20 °C (dark) until sampling. The *O. laciniata* and *O. stricta* were grown at Nagoya University, and the flowers in full bloom were collected and used for the experiment.

4.2. HPLC and Structural Analysis of the Flavonoids in the Petals

In both kinds of *Oenothera*, the blooming flowers were sampled at night (approximately 0-2 h after blooming), whereas the senescent flowers were collected in the morning (approximately 12-18 h after blooming). HPLC analysis was done on small scale according to Yoshida et al. with some modifications [15]. Petals (1 mg) portion were extracted with 20 µl of 3% trifluoroacetic acid (TFA) in 50% aqueous acetonitrile (CH₃CN). The extracts were analyzed by HPLC using a RPAQUEOUS-AR-3 column (2.0 × 150 mm) with linear gradient elution from 10% to 50% aqueous CH₃CN containing 0.5% TFA.

To verify the structure of the flavonoids, LC-MS analysis was performed on a Bruker Daltonics microTOF-QII mass spectrophotometer with an Agilent 1200 Series HPLC system in ESI-positive ion mode with the same HPLC conditions. The extracts were also co-chromatographed with authentic samples to confirm the structure of the compound.

4.3. Quantitative Analysis of the Flavonoids by HPLC

Briefly, all petals of flowers in full bloom were picked and weighed individually. After incubation in a growth chamber for the designed times (0, 4, 7, and 12 h for *O. tetraperta* and 0, 4, 8, and 12 h for *O. laciniata*), the petals were collected. The flavonoids in the petals were extracted and analyzed by HPLC as described above. The content of flavonoids was calculated using a standard curve prepared from the purified compounds [15]. The experiment was performed in triplicate. The obtained data were evaluated by one-way ANOVA with post hoc Tukey’s HSD test (p values ≤ 0.05).

4.4. UV/Vis and Color Parameter Measurements

UV/Vis spectra as well as the color parameters of the *O. tetraperta* petals were measured by a JASCO V-560 UV/Vis spectrophotometer equipped with an integral sphere. The upper edges of the petals were cut into 1.5×1.5 cm squares for use in these analyses.
4.5. Petal pH Measurements

For petal pH measurements, fresh petals of *O. tetraperta* were ground, and then the pH of the obtained petal juice was measured by a pH meter.

4.6. Isolation and Characterization of Isosalipurposide from *Oenothera lacinia*ta

The petals of *O. lacinia*ta (150 g) were extracted 2 times with methanol. The crude extract was concentrated and dried under reduced pressure, and the residue was resuspended with 50% aqueous methanol. After sonification, the mixture was filtered through a PTEE membrane filter (pore size: 0.5 μm) and purified via preparative HPLC with an ODS-HG-5 column (25 mm i.d. × 250 mm) at a flow rate of 15 ml/min. The mobile phases were 0.1% TFA in 5% aqueous CH3CN for 0-5 min, 0.1% TFA in 20% aqueous CH3CN for 5-20 min, 0.1% TFA in 30% aqueous CH3CN for 20-35 min, and 0.1% TFA in 90% aqueous CH3CN for 35-50 min. The fractions were analyzed by HPLC to check their purity.

Pure isosalipurposide was structurally analyzed by NMR, MS, UV/Vis and IR techniques and used as a standard sample in the analysis of the flavonoids. 1H and 13C NMR spectra were obtained with a JEOL JNM-ECA-500 spectrometer. Chemical shifts are reported in ppm relative to CD3OD (δ = 3.31 ppm for 1H NMR and δ = 49.0 ppm for 13C NMR) as the reference. High-resolution mass spectra were recorded using a Bruker microTOF-QII electrospray ionization (ESI) spectrometer. IR spectra were obtained from KBr pellets on a JASCO FT/IR-460 plus spectrometer, while UV/Vis spectra were collected by a JASCO V-560 spectrophotometer (path length: 10 mm).

Isosalipurposide (Chalconaringenin 2′-glucoside), 3: ESI-MS (positive mode) m/z = 435 [M+H]+; λmax (ethanol) = 371 nm (ε = 24,900). IR (v/ν) 1626 cm−1. 1H NMR (500 MHz, CD3OD) δ 8.02 (1H, d, J = 15.5 Hz; H-α), 7.67 (1H, d, J = 15.5 Hz; H-β), 7.61 (2H, d, J = 8.6 Hz; H-2, H-6), 6.83 (2H, d, J = 8.6 Hz; H-3, H-5), 6.22 (1H, d, J = 1.7 Hz; H-3′), 6.00 (1H, d, J = 2.3 Hz; H-5′), 5.14 (1H, d, J = 7.5 Hz; H-1′), 3.92 (1H, dd, J = 2.3, 12.6 Hz; H-6′a), 3.74 (1H, dd, J = 5.2, 12.0 Hz; H-6′b), 3.56 (1H, dd, J = 7.4, 9.2 Hz; H-2′), 3.51 (1H, t, J = 8.6 Hz; H-3′′), 3.46 (1H, dd, J = 2.3, 5.2, 12.6 Hz; H-5′′), 3.44 (1H, t, J = 8.9 Hz; H-4″).

13C NMR (125 MHz, CD3OD) δ 194.5 (C=O), 167.8 (C-6′), 165.9 (C-4′), 161.1 (C-2′), 144.2 (C-β), 131.8 (C-2, C-6′), 128.5 (C-1′), 125.9 (C-α), 116.9 (C-3, C-5), 107.5 (C-1′), 101.9 (C-1″), 98.4 (C-5′), 95.7 (C-3′), 78.5 (C-3″, C-5″), 75.0 (C-2″), 71.2 (C-4″), 62.4 (C-6″).

Supplementary Materials: The following are available online at www.mdpi.com/online. Figure S1: The LC-MS spectra of Cy3G (1), quercitrin (2) and isosalipurposide (3) from the extracts of *Oenothera* flowers. Figure S2: HPLC chromatogram of the extracts from petals of *O. stricta*. Figure S3: The 1H NMR spectrum (500 MHz) of isosalipurposide (3) in CD3OD at 25 °C. Figure S4: The 13C NMR spectrum (125 MHz) of isosalipurposide (3) in CD3OD at 25 °C. Figure S5: The COSY spectrum of isosalipurposide (3) in CD3OD at 25 °C. Figure S6: The NOESY spectrum of isosalipurposide (3) in CD3OD at 63 °C. Figure S7: The HMBC spectrum of isosalipurposide (3) in CD3OD at 25 °C. Figure S8: The HMQC spectrum of isosalipurposide (3) in CD3OD at 25 °C.

Acknowledgments: We would like to thank the Kochi Prefectural Makino Botanical Garden for providing *O. tetraperta* flowers and Ms. Natsuko Yoshino of The Nagoya University Museum for her assistance in collecting the *O. lacinia*ta and *O. stricta* samples. Y.T. acknowledges the financial support from the Ministry of Education, Culture, Sports, Science and Technology, Japan (MEXT) scholarship and the Graduate School of Informatics Doctoral Course Student Research Expense Grant, Nagoya University.

Author Contributions: K.Y. conceived and designed the experiments and wrote the paper; Y.T. performed the experiments and wrote the paper; Y.T. and T.K. analyzed the data; and K.O. contributed analysis tools.

Conflicts of Interest: The authors declare no conflict of interest. The funding agencies had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; or in the decision to publish the results.

References


**Sample Availability:** Samples of the compounds 1 and 3 are available from the authors.