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Huyen, Dao Thi Thanh Graduate School of Bioresource and Bioenvironmental Science, Kyushu University

Van, Dao Thanh Thai Nguyen University of Agriculture and Forestry

Huang, KuangLiang
Department of Horticultural Science, National Chiayi University

Miyajima, Ikuo Institute of Tropical Agriculture, Kyushu University

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#### Distribution and Composition of Flavonols in the Flowers of Rhododendron oldhamii Maxim.

#### Dao Thi Thanh HUYEN¹, Dao Thanh VAN², KuangLiang HUANG³ and Ikuo MIYAJIMA\*

Institute of Tropical Agriculture, Kyushu University, Fukuoka 812–8581 Japan (Received November 9, 2015 and accepted November 19, 2015)

Rhododendron oldhamii Maxim. is an endemic evergreen azalea in Taiwan and recently planted in Japan. This species is distributed from 150 m to 2,700 m above sea level. Red–flowered species with petal color ranging from orange to red normally contain only cyanidin series anthocyanins and no flavonols in their petals. However, the color of blotches on the upper lobe of the petals or at the base of the funnel is slightly reddish–purple that suggested the presence of flavonols. In R. oldhamii, cyanidin 3–arabinoside and cyanidin 3–galactoside are two major anthocyanins in both upper and lower petals but flavonols were only detected in the upper petals, specifically in the blotches areas, by HPLC analysis. Two major flavonols were identified as quercetin glycosides. The pH value of reddish–purple blotches was slightly higher than that of surrounding areas and lower petals. These results indicated that co–pigmentation between cyanidin glycosides and quercetin glycosides pigments associated with higher pH condition, causing reddish–purple color for blotches in R. oldhamii flowers.

 $\textbf{Key words}: \text{ anthocyanin, copigmentation, flavonol, } Rhododendron \ old hamii$ 

#### INTRODUCTION

Rhododendron oldhamii Maxim., classified in the subgenus Tsutsusi, is a subtropical broadleaf species endemic to Taiwan and was scientifically described for the first time in 1870 (Hsieh et al., 2013). This species is distributed from 150 m to 2,700 m above sea level, and the population size of it is the largest among all Rhododendrons in Taiwan. It is commonly used as important ornamental plants for gardens, street plantings or flowerpots because of the beauty of flowers and multiple flowering seasons per year.

Rhododendron oldhamii is semideciduous shrubs with red, brick-red or orange five-petal-lobe flowers and reddish-purple blotches on the upper insides of the petals (Fig. 1). Pigment components of some red flowered species such as R. simsii, R. indicum and R. oldhamii flowers have been reported to be cyanidin 3-galactoside and cyanidin 3-arabinoside as two major anthocyanins in whole flower petals (Hang et al., 2011). However, the development of the reddish-purple color of blotches in R. oldhamii has not been well analyzed and explained. In azaleas, it is well-known that co-pigmentation between anthocyanins and flavonols has a bluing effect for flower color (Asen et al., 1971; De Loose, 1978). Reddish-purple blotches of R. oldhamii suggest the existence of kind of co-pigments, which should be revealed.

The purpose of this study is to clarify the distribution and composition of pigments in reddish–purple



Fig. 1. Full opened Rhododendron oldhamii flowers. Bar indicates 1 cm.

blotches of R. oldhamii petals by anatomical and chromatographic methods.

#### MATERIALS AND METHODS

#### Plant materials

Full–opened R. oldhamii flowers were collected at Kyushu University greenhouse. A part of fresh petals was used for microscopic observation of the cross–sections. The remaining was separated into upper and lower petals, and each part of petals was boiled at  $100^{\circ}\mathrm{C}$  for 5 seconds and immediately cooled in water. Then the samples were dried in the forced convention oven overnight at  $50^{\circ}\mathrm{C}$ . The dried samples were stored in a desiccator at  $4^{\circ}\mathrm{C}$  until using for pigment analysis.

Graduate School of Bioresource and Bioenvironmental Science, Kyushu University, Fukuoka 8128581 Japan

<sup>&</sup>lt;sup>2</sup> Thai Nguyen University of Agriculture and Forestry, Thai Nguyen 252161, Vietnam

<sup>&</sup>lt;sup>3</sup> Department of Horticultural Science, National Chiayi University, Chiayi 60004, Taiwan, ROC.

<sup>\*</sup> Corresponding author: (E-mail: imiyajima@agr.kyushuu.ac.jp).

## Microscopic observation of pigment distribution and absorption spectra of fresh petals

Upper and lower petals of fresh flowers of  $R.\ old-hamii$  were cut into small squares at the center positions and fixed using 5% agar in the petri dish. Then they were sliced using a microslicer (DTK–1000, Dosaka EM; Kyoto, Japan) at a thickness of  $150\,\mu\mathrm{m}$ . After slicing, cross–sections were observed under an optical microscope (Leica DM–2500; Leica Microsystems GmbH, Wetzlar, Germany).

#### **HPLC** analysis

Dried petals (ca. 50 mg) of R. oldhamii were soaked overnight with 50% HOAc–H<sub>2</sub>O. After filtration, analytical HPLC was conducted on a LC–20AD pump (Shimadzu, Kyoto, Japan), using a Cosmosil 5C<sub>18</sub> MS–II column (4.6  $\phi \times 250$  mm; Nakalai Tesque, Kyoto, Japan) at 40°C with a flow rate of 1 mL·min<sup>-1</sup>, and monitoring at 520 nm and 360 nm for anthocyanins and flavonols, respectively. A linear gradient elution was applied for 40 min from 20 to 85% solvent B (1.5% H<sub>3</sub>PO<sub>4</sub>, 20% HOAc, 25% MeCN in H<sub>2</sub>O) in solvent A (1.5% H<sub>3</sub>PO<sub>4</sub> in H<sub>2</sub>O).

#### **Identification of pigments**

Two–dimensional thin layer chromatography (TLC) was carried out on cellulose–coated glass plates (Merck, Darmstadt, Germany) using two mobile phases: BAW (1–BuOH/HOAc/H<sub>2</sub>O, 4:1:2, v/v/v) and 10% HOAc. The plates were observed under the UV light (365 nm), and the color of all spots was recorded. Subsequently, each spot was collected from the TLC plates, and dissolved using MeOH for HPLC analysis.

#### Isolation of major flavonols

Dried petals (ca. 0.7 g) of R. oldhamii were soaked overnight with 100% MeOH. After filtration, preparative HPLC was performed on an LC–6AD system (Shimadzu, Kyoto, Japan), using a Cosmosil  $5C_{18}$  AR column ( $20~\phi \times 250~\text{mm}$ ; Nakalai Tesque, Kyoto, Japan) at  $40^{\circ}\text{C}$  with a flow rate of  $9~\text{mL}\cdot\text{min}^{-1}$ , and monitoring at 360~nm for isolation of major flavonol peaks. A linear gradient elution was applied for 40~min from 50~to 85% solvent B (10%~formic acid, 40%~MeCN in  $H_2\text{O}$ ) in solvent A (10%~formic acid in  $H_2\text{O}$ ). Major peaks were obtained using a fraction collector.

#### Identification of flavonol aglycones

Each purified flavonol was acid hydrolyzed by 2N HCl at 100°C for 90 min. The flavonol aglycones were cochromatographed with authentic standard flavonols aglycones such as myricetin, quercetin and kaempferol, by HPLC with constant flow of 75% solvent A (0.1 M HOAc): 25% solvent B (MeCN). The HPLC system, column, and flow rate were the same as mentioned above (HPLC analysis). Wavelength was set at 360 nm.

#### pH measurement

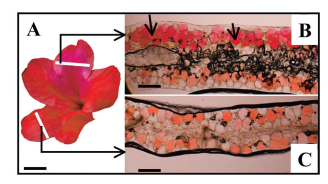
Fresh flowers of *R. oldhamii* were separated into blotches, surrounding areas of botches and lower petals. The pH value of squeezed juice of each part was deter-

mined using a compact pH meter (TWIN pH waterproof B–212; Horiba Ltd., Kyoto, Japan). Five flowers were measured as replications.

#### RESULTS AND DISCUSSION

## Distribution of pigmented cells in the petals of R. oldhamii flowers

Blotches of R. oldhamii flowers showed numerous reddish-purple pigmented cells, which accumulated in the adaxial epidermis of upper petals (Fig. 2B). On the other hand, red colored cells were distributed in the epidermis of both upper and lower petals (Fig. 2B, C). The upper epidermis of blotch areas was flat, and the shape of reddish-purple pigmented cells was not different from that of red pigmented cells. Generally, blotches in flowers of Rhododendron species contain stronger colored cells in the adaxial subepidermis rather than in epidermis. (Pecherer, 1992). The same results were observed in the transverse section of red blotches in R. schlippenbachii flowers. While light pink colored cells accumulated in the upper epidermis, red pigments were distributed in subepidermal cells (Yamagishi and Akagi, 2013). In R. oldhamii flowers, reddish-purple pigmented cells are only distributed in adaxial subepidermis of blotches areas.



**Fig. 2.** Cross—sections of flower petals of *R. oldhamii*. A: Whole flower, B: Cross—section of upper petal, C: Cross—section of lower petal.

White lines in the photo A indicate sliced positions. Arrows in the photo B indicate reddish–purple cells. Bars: photo A= 1 cm, photo B and C=  $20 \mu \text{m}$ .

#### Pigment composition in flower petals

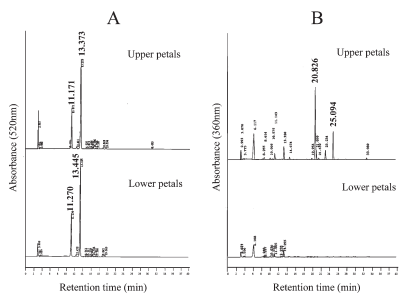
Both upper and lower petals contained two major anthocyanins (Fig. 3A). These two anthocyanin pigments were reexamined and confirmed to be cyanidin 3–galactoside (Cy 3Ga) and cyanidin 3–arabinoside (Cy 3Ar) as previously reported in red flowered R. simsii and R. oldhamii as well (Hang  $et\ al.$ , 2011). At the absorbance of 360 nm, two major peaks were detected in upper petals, while no major peak was detected in lower petals (Fig. 3B). Thus, these two flavonoids seem to be present only in upper petals of R. oldhamii flowers.

To identify these two major flavonoids in upper petals, two-dimensional TLC was carried out. Eight discrete spots appeared in the TLC plates (Fig. 4) and color properties were recorded under visible and UV light

Spot No.	Rf value (×100)		Color in	
	$\mathrm{BAW}^{\scriptscriptstyle{\;1)}}$	10% AcOH	Visible light	UV–light
1	17	22	Red-lilac	Violet
2	48	14	Red-lilac	Violet
3	65	20	Pale brown	Yellow F. 3)
4	76	30	Pale brown	Yellow F.
5	75	48	- <sup>2)</sup>	Blue F.
6	39	50	_	Blue F.
7	64	66	_	Blue F.
8	37	70	_	Blue F.

**Table 1.** Rf values and color properties of spots of pigments extracted from *R. oldhamii* flowers and separated by two-dimensional thin layer chromatography

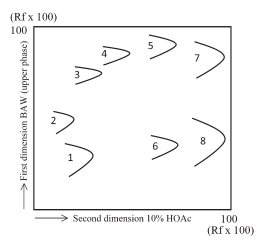
<sup>3)</sup> fluorescent.



**Fig. 3.** HPLC profiles of the extracts from upper and lower petals of R. oldhamii at  $520\,\mathrm{nm}$  (A) and  $360\,\mathrm{nm}$  (B).

(Table 1). Spot numbers 1 and 2 expressed red-lilac color under visible light and violet under UV light. This result suggested that they were two major anthocyanins (Cy 3Ga and Cy 3Ar). In addition, spots 3 and 4 appeared pale brown under visible light, and showed yellow florescence under UV light. Spethmann (1980) investigated flavonoids of Rhododendron flowers, and reported that some flavonol glycosides, such as quercetin 3-arabinoside, quercetin 3-rhamnoside or kaempferol 5-methylether, appeared brown or pale brown under visible light and colored greenish-yellow to yellow florescence under UV light on TLC plate. These findings suggested that spots 3 and 4 are flavonols. The HPLC analysis of spots 3 and 4 was carried out after collection and extraction from TLC plate. From HPLC analysis, spots 3 and 4 coincided with two major peaks in upper petals (Fig. 3B).

Next, these two major peaks in upper petals were isolated by preparative HPLC and acid hydrolyzed using 2N HCl. HPLC analysis showed that two major peaks



**Fig. 4.** Two-dimensional thin layer chromatogram of MeOH extract from upper petals of *R. oldhamii* flowers. BAW (1-BuOH/HOAc/H<sub>2</sub>O, 4:1:2, v/v/v). See Table 1 for color properties of spots.

<sup>&</sup>lt;sup>1)</sup> 1-BuOH/HOAc/H<sub>2</sub>O = 4:1:2, v/v/v

<sup>2)</sup> colorless.

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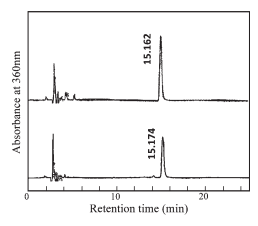
**Table 2.** HPLC retention time of standard flavonols and flavonol aglycones of *R. oldhamii* flowers

Aglycones	Retention times (min)		
Myricetin	8.0		
Quercetin	15.2		
Kaempferol	28.8		
Peak 1	15.2		
Peak 2	15.2		

**Table 3.** pH value of the flower parts of *R. oldhamii* 

Flower parts	pH value	
Blotches	3.14 a <sup>1)</sup>	
Surrounding areas of blotches	2.18 b	
Lower petal	2.26 b	

 $<sup>^{1)}</sup>$  Values with different letters are significantly different at  $P{<}0.05$  by Tukey's test.



**Fig. 5.** HPLC tracing of acid hydrolysates of two major flavonols isolated from the petals of *R. old-hamii*.

have same flavonol aglycones at the retention time of 15 min (Fig. 4). In comparison to authentic standard samples, such as myricetin, quercetin and kaempferol, the aglycones of two major peaks were identified as quercetin (Table 2).

Asen et al. (1971) determined the cause of the difference in color expression between 'Red Wing' azalea and an orange sport of this cultivar. The orange color of the mutant was due to cyanidin glycosides, whereas the color of 'Red Wing' azalea was due to the same cyanidin glycosides co-pigmented with quercetin glycosides. Carmine red color of some cultivar of azalea species, such as 'Vuyk's Scarlet', is in effect produced by the addition of flavonol (quercetin-glycosides) to the red pigments cyanidin or peonidin (Heursel, 1987). Thus, quer-

cetin glycosides co-pigmented with cyanidin glycosides to develop 'bluing effect' causing reddish-purple blotches in *R. oldhamii* flowers.

#### pH value of petal parts

The pH value of the pressed juice from various positions of same flowers was quite different. The pH of reddish–purple blotches was 3.14. On the other hand, that value of surrounding areas of blotches and lower petals were 2.18 and 2.26, respectively. The same results were also reported by Stewart *et al.* (1975) in fuchsia cv. Fanfare flowers. The pH of pink calyx was 4.1, while that of the strong red corolla was 3.7. The pH of reddish–purple blotch areas was higher than that of red petal parts. However, petal color is not only determined by pH value, but also by co–pigments or other inorganic substances, such as Al<sup>3+</sup> (Yoshida *et al.*, 2003).

In conclusion, these results indicated that co-pigmentation between cyanidin glycosides and quercetin glycosides pigments associated with higher pH condition, causing reddish-purple color for blotches in *R. oldhamii* flowers.

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