Quercetin 3-glucoside and Quercetin 3-rhamnoside Found in the Reddish-purple Blotch Areas of Rhododendron simsii Planch. Flowers

Huyen, Dao Thi Thanh
Institute of Tropical Agriculture, Kyushu University | The Graduate School of Bioresource and Bioenvironmental Sciences, Kyushu University

Ureshino, Kenji
Institute of Tropical Agriculture, Kyushu University | Faculty of Agriculture, University of the Ryukyus

Van, Dao Thanh
Institute of Tropical Agriculture, Kyushu University | Thai Nguyen University of Agriculture and Forestry

Miyajima, Ikuo
Institute of Tropical Agriculture, Kyushu University

http://hdl.handle.net/2324/1799301
Quercetin 3–glucoside and Quercetin 3–rhamnoside Found in the Reddish–purple Blotch Areas of Rhododendron simsii Planch. Flowers

Dao Thi Thanh HUYEN¹, Kenji URESHINO², Dao Thanh VAN³ 
and Ikuo MIYAJIMA⁴*

Institute of Tropical Agriculture, Kyushu University, Fukuoka 812–8581 Japan
(Received October 28, 2016 and accepted November 4, 2016)

The flowers of Rhododendron simsii have reddish–purple blotches in the upper inside of the red petals. HPLC analysis indicated that two major flavonols present in the blotch areas besides anthocyanins. Quercetin 3–glucoside and quercetin 3–rhamnoside were identified by co–chromatographic TLC, HPLC and LC–MS analyses.

Key words: Flavonols, quercetin 3–glucoside, quercetin 3–rhamnoside, Rhododendron simsii

INTRODUCTION

In Vietnam, some accessions of wild Rhododendron simsii are found in northern and central mountainous area, where this species grow near riverside or streamside at 800–1,400 m altitude (Ho, 1991). All of these accessions show only red colored funnel shaped flowers with reddish–purple blotches in upper inside of the petals (Fig. 1). In our continued work on pigmentation of R. simsii flowers, co–pigmentation of anthocyanin–flavonol, and the effect on flavonol for coloration of reddish–purple blotches of R. simsii flowers was revealed (Huyen et al., 2016). In this report, two major flavonols were detected in the upper petals of R. simsii flowers (Fig. 2). However, the flavonols isolated from reddish–purple blotches of R. simsii flowers have not yet been identified.

Flavonols of some wild evergreen azalea species were effectively analyzed by High Performance Liquid Chromatography (HPLC) and Thin Layer Chromatography (TLC) (Harborne, 1959; Miyajima et al., 1997, 2001; Sakata et al., 1991). In this study, chromatographic analyses, and the flavonol properties in the flowers of several red flowered evergreen azalea species were discussed.

Fig. 1. Rhododendron simsii flower having red corolla with reddish–purple blotches on the upper inside of the petals. Bar indicates 1 cm.

Fig. 2. HPLC profiles of flavonols extracted from the upper petals of R. simsii.
MATERIALS AND METHODS

Plant materials

Fresh flowers of wild *R. simsii* distributed in Than Uyen district, Lai Chau province, Vietnam were collected at anthesis. Flowers were separated into upper and lower petals, and each part of petals was boiled at 100°C for 5 seconds, and immediately cooled in water. Then, the samples were dried in a forced convention oven through overnight at 50°C. The dried samples were stored in a desiccator at 4°C until pigment analysis.

Isolation and purification of major flavonols

Dried petals (ca. 0.7 g) of *R. simsii* were soaked overnight with 100% MeOH. After filtration, preparative HPLC was performed on an LC–6AD system (Shimadzu, Kyoto, Japan), using a Cosmosil 5C18 AR column (20 φ x 250 mm; Nakalai Tesque, Kyoto, Japan) at 40°C with a flow rate of 9 mL·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 H₂O) in solvent A (10% formic acid in H₂O). Major peaks were collected using a fraction collector.

Co–chromatography by TLC analysis

After isolating from preparative HPLC, each purified flavonol fraction was dried by evaporator and diluted by drops of MeOH. Each pigment was spotted on 10 cm x 20 cm cellulose–coated glass plates (Merck, Darmstadt, Germany). In addition, four authentic standard quercetin glycosides: quercetin 3–glucoside (Qu 3Gl, isoquercetin), quercetin 3–galactoside (Qu 3Ga, hyperoside), quercetin 3–rutinoside (Qu 3Rt, rutin) and quercetin 3–rhamnoside (Qu 3Rh, quercitrin) were also spotted in order to compare with two major flavonols. Thin layer chromatography analyses were performed by three mobile phases according to the method of Harborne (1959): BAW (1–BuOH/AcOH/H₂O, 4:1:2, v/v/v), 100% H₂O and 15% AcOH. Then all spots were observed under UV light (365 nm) and *Rf* values were also recorded.

Co–chromatography by HPLC analysis

In order to compare the retention times of two major flavonols and four authentic standard quercetin glycosides (same as mentioned above), HPLC analyses were conducted in different solvent systems: (solvent system I) A (1.5% H₃PO₄) and B (1.5% H₃PO₄: 20% AcOH: 25% CH₃CN), B conc.: 20%–85% at 40 min; (solvent system II) A (4% H₃PO₄) and B (CH₃CN), B conc.: 15%–30% in 30 min; (solvent system III) A (0.1 M AcOH) and B (CH₃CN) with constant flow. The wavelength of the analyses was 360 nm. Other HPLC conditions (pump, column, column oven temperature, and flow rate) were same as described by Huyen et al. (2016).

LC–MS analysis

Two purified major flavonols were analyzed using LC–MS by TSK gel ODS–80Ts QA 2.0 φ x 150 mm column. A constant flow was applied for 30 min at 20% solvent B (MeCN) in solvent A (0.1 M AcOH) with a flow rate of 0.2 mL·min⁻¹ and 40°C of column temperature.

Flavonol constitution of the flowers in several red flowered evergreen azalea species

Approximately 50 mg of dried upper and lower petals from *R. simsii* (Iriomotejima Island, Japan), *R. indicum* (Yakushima Island, Japan), *R. oldhamii* (Alishan, Taiwan, ROC) and *R. scabrum* (Okinawa Island, Japan) were soaked overnight with 50% HOAc–H₂O, respectively. After filtration, analytical HPLC was conducted in the same condition as described by Huyen et al. (2016).

RESULTS AND DISCUSSION

By using preparative HPLC, two major flavonols were isolated and purified. The result of co–chromatography by TLC showed that the *Rf* value of flavonol 1 (F1) was almost same as Qu 3Gl or Qu 3Ga in three different solvent systems and could not be distinguished. Flavonol 2 (F2) seemed to be quercetin 3–rhamnoside (Table 1). However, HPLC analysis results indicated the differences in retention times between two standard flavonols Qu 3Gl and Qu 3Ga, and the retention time of F1 was close to that of Qu 3Gl. Thus, F1 and F2 were almost coincided with quercetin 3–glucoside and quercetin 3–rhamnoside, respectively (Table 2). This result was supported by the molecular weight analysis. Table 3 showed that F1 [465.09 g/mol (M + H⁺)] and F2 [449.10 g/mol (M + H⁺)] had concurred molecular weight to that of standard Qu 3Gl and Qu 3Rh, respectively. The chemical struc-

| Table 1. TLC analysis of F1, F2 and four authentic standard quercetin glycosides in different solvent systems |
|---------------------------------|----------|---|---|
| Flavonol glycosides             | BAW      | H₂O | 15% AcOH |
| Flavonol 1                      | 69       | 06  | 27      |
| Flavonol 2                      | 84       | 12  | 40      |
| Quercetin 3–glucoside (isoquercetin) | 68     | 07  | 29      |
| Quercetin 3–galactoside (hyperoside) | 64     | 09  | 31      |
| Quercetin 3–rutinoside (rutin)  | 61       | 35  | 53      |
| Quercetin 3–rhamnoside (quercitrin) | 84     | 13  | 40      |

* 1–butanol: AcOH: DW (4: 1: 2, v/v/v)
Quercetin 3–glucoside and Quercetin 3–rhamnoside Found in Rhododendron simsii Flowers.

In Rhododendron species, four quercetin glycosides were identified as quercetin 3–arabinoside, 3–rhamnoside, 3–galactoside and the 3–rhamnoside of quercetin 5–methyl ether (Harborne, 1962). De Loose (1969) reported that quercetin glycosides (3–rhamnoside or 3–galactoside) seem to be popular in the flower of Red Wing Azalea (R. simsii hybrids). However, this is the first report about the presence of quercetin 3–glucoside and 3–rhamnoside in flowers of wild R. simsii distributed in Vietnam.

The flavonol composition of several wild red flowered species was also surveyed by HPLC analysis. The results showed that upper petals of these species also contained two major peaks and coincided with two major flavonols (Qu 3Gl and Qu 3Rh), which were identified in the blotch areas of R. simsii flowers from Vietnam in this study (Fig. 4).

From the results, quercetin 3–glucoside (isoquercetin) and quercetin 3–rhamnoside (quercitrin) present as the two flavonols in the reddish–purple blotch areas of flowers of R. simsii and several red flowered species. These two flavonols play an important role on co–pigmentation phenomenon. Reddish–purple color of blotch areas is due to quercetin 3–glucoside and quercetin 3–rhamnoside co–pigmented with cyanidin 3–galactoside and cyanidin 3–arabinoside. This detailed understanding about pigmentation of flowers will contribute necessary knowledge for flower color breeding in the further study using Vietnamese wild R. simsii as the important genetic resources.

### REFERENCES

De Loose, R. 1969 The flower pigments of the Belgian hybrids of Rhododendron simsii and other species and varieties from Rhododendron subspecies obtusum. Phytom, 8: 253–259


Fig. 4. HPLC profiles of extracts from upper and lower petals of *R. simsii* and several red flowered evergreen azalea species. Qu 3Gl: quercetin 3-glucoside; Qu 3Rh: quercetin 3-rhamnoside.