Seasonal Alteration in Amounts of Lignans and Their Glucosides and Gene Expression of the Relevant Biosynthetic Enzymes in the Forsythia suspense Leaf

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Lignans of Forsythia spp. are essential components of various Chinese medicines and health diets. However, the seasonal alteration in lignan amounts and the gene expression profile of lignan-biosynthetic enzymes has yet to be investigated. In this study, we have assessed seasonal alteration in amounts of major lignans, such as pinoresinol, matairesinol, and arctigenin, and examined the gene expression profile of pinoresinol/lariciresinol reductase (PLR), pinoresinol-glucosylating enzyme (UGT71A18), and secoisolariciresinol dehydrogenase (SIRD) in the leaf of Forsythia suspense from April to November. All of the lignans in the leaf continuously increased from April to June, reached the maximal level in June, and then decreased. Ninety percent of pinoresinol and matairesinol was converted into glucosides, while approximately 50% of arctigenin was aglycone. PLR was stably expressed from April to August, whereas the SIRD expression was not detected from September to November. In contrast, the UGT71A18 expression was found from August to November, but not from April to July. The SIRD expression was prominent from April to May, not detected in June to July, and then increased again from September to November. These expression profiles of the lignan-synthetic enzymes are largely compatible with the alteration in lignan contents. Furthermore, such seasonal lignan profiles are in good agreement with the fact that the Forsythia leaves for Chinese medicinal tea are harvested in June. This is the first report on seasonal alteration in lignans and the relevant biosynthetic enzyme genes in the leaf of Forsythia species.

Key words biosynthetic enzyme; Forsythia; lignan; lignan glycoside

Lignans have a variety of clinically and dietarily important biological activities. Lignan biosynthesis is well studied in Linum, Forsythia and other plants. These plants are used as Chinese medicines and health diets. Forsythia, commonly known as golden bell flower, is a perennial woody plant, and consists of a large number of varieties, including F. suspensa. In Forsythia, major lignan biosynthesis pathways have been determined (Fig. 1). Pinoresinol, a basal lignan, is stepwisely reduced into secoisolariciresinol via lariciresinol by pinoresinol/lariciresinol reductase (PLR) or glucosylated by UGT71A18. Secoisolariciresinol is converted into matairesinol by secoisolariciresinol dehydrogenase (SIRD). Furthermore, matairesinol is glucosylated or consumed for the biosynthesis of arctigenin by unidentified enzymes (Fig. 1). Arctigenin is also glucosylated to arctiin by an unidentified glucosyltransferases.

Lignans and their glucosides in Forsythia fruits and leaves exhibit diverse biological activities on mammals. Most lignans are uptaken directly, or further metabolized by enteric bacteria to enterolactone and enterodiol, designated as phytoestrogens. There have been increasing reports on the effects of the dietary lignans on reduction of oxidative injury, post-menopause symptoms, and cancer risks. Notably, each lignan was found to exhibit both similar and differential bioactivities in mammals, which underscores the dietary and medicinal significance of specific production of a targeted lignan, instead of lignan mixtures. These findings suggest a potential increase in the demands of isolated lignans for the development of health diets, supplements, or drugs, and the requirement of efficient acquisition of a lignan from source plants. Thus, elucidation of seasonal profiles in lignan amounts and the biosynthetic enzymes is expected to develop novel regulatory systems for efficient production of target lignans in Forsythia species. In this paper, we present the seasonal profiles of pinoresinol, matairesinol, arctigenin, and their biosynthetic enzyme genes (Fig. 1) in the leaves of F. suspense.

MATERIALS AND METHODS

Plant Material and Leaf Sampling The F. suspense plants were cultivated in the outdoor garden at Suntory Business Expert Limited in Osaka, Japan. Eight to thirty F. suspense leaves located at the outermost branches of the middle parts of the fully grown plants were sampled every month from April 5 to November 15 (Table 1). The sampled leaves were cut into small pieces and mixed well. The mixed leaf pieces were aliquoted into 0.2 g and kept at −80°C until use. In addition, the climate conditions for each sampling day are summarized in Supplemental data 1.

Lignan Extraction and HPLC Analysis 0.2 g of the leaf piece was suspended in 1 mL of 80% ethanol and 20% 0.1 M sodium phosphate buffer (pH 4.6) and sonicated twice for 15 s in 4°C. The mixture was centrifuged and the supernatant was evaporated to 200 µL. The remaining phase was adjusted to 1 mL by addition of 0.1 M sodium phosphate buffer. The H2O phase containing the lignan glycoside was digested at 40°C overnight with 6 U mL−1 almond β-glucosidase (Sigma-Aldrich, St. Louis, MO, U.S.A.) in 0.1 M sodium phosphate buffer (pH 4.6). Lignan samples with or without β-glucosidase treatment were dissolved again with 50% aqueous acetonitrile.
trile, and analyzed by reverse-phase high performance liquid chromatography (HPLC) using a Develosil C30-UG-5 column (4.6×150 mm, Nomura Chemical, Aichi, Japan) with an LC-2010A HT system (Shimadzu, Kyoto, Japan) as previously reported.28-30 These extraction and HPLC analysis of lignan contents were independently performed three times.

Reverse Transcription Polymerase Chain Reaction (RT-PCR) Analyses Total RNA was isolated from the sampled leaf blade using an RNeasy Plant Mini Kit (Qiagen, Valencia, CA, U.S.A.). First-strand cDNA was synthesized using 500 ng of total RNA with a reverse transcriptase SuperScript III (Invitrogen, San Diego, CA, U.S.A.). PCR primer sequences of PLR (accession No. AAC49608), SIRD (accession No. AF352735), UG771A18 (accession No. AB524718), and ribosome RNA (accession No. AJ236041) are summarized in Supplemental data 2. PCR was performed at 95°C for 1 min, followed by 30 cycles at 94°C for 20 s, at 52–58°C for 30 s, at 72°C for 1 min, and a final extension at 72°C for 7 min on a thermal cycler (ABI GeneAmp 9700) using ExTaq DNA polymerase (TaKaRa Bio Inc., Kyoto, Japan). PCR products were resolved on 1% agarose gel.

Statistical Analysis Results are expressed as means±S.E.M. for indicated number of observations. Data were analyzed by one-way ANOVA with Dunnett error protection. Differences were accepted as significant for \( p<0.05 \).

RESULTS

Growth of *F. suspensa* under Natural Condition *F. suspensa* leaves generated new buds in early April. On April 5, the buds rapidly developed to 2 to 4-cm length young leaves (Fig. 2). On April 15, the branches developed to 10-cm long, and the leaves gradually further grew to a 5-cm length and 2.5-cm width, and stooped in June. From July to September, no morphological changes in the leaves were observed. In September, the leaves started to turn yellowish at the leaf edge and gradually turned to brown. The leaves were gradually withered from September to November, and then completely dropped. The average weight of the leaves was rapidly increased from April (0.01 g) to September (0.58 g), but decreased from September, ultimately to 0.24 g in November (Table 1).

Seasonal Alteration in Lignan Amounts in *F. suspensa* Leaves The major lignans in *Forsythia* leaves are matairesinol, arctigenin, pinoresinol and their glucosides.1 Total pinoresinol (aglycone and glucosides) amounts were constantly low [1–1.6 mg/g of the dry weight of the leaf (DW)], and statistically significant alteration in amounts was not found (Fig. 3A), compared to matairesinol and arctigenin throughout season (Figs. 3B, C). Pinoresinol aglycone was also consistently detected at low level (approximately 0.2 mg/g DW) from April to September, and further decreased from October. Total matairesinol increased from 10 to 20 mg/g DW for two months as leaves grew (Fig. 3B). The amount of matairesinol peaked in June and then gradually decreased to 2 mg/g DW in November. The matairesinol aglycone content went from 0.2 to 2.0 mg/g DW, and the maximal level was observed in July (Fig. 3B). Total arctigenin content was 12.5 mg/g DW in April and increased to 30 mg/g DW in June, and then decreased to 4 mg/g DW in October (Fig. 3C). These results confirm that matairesinol and arctigenin and their glucosides account for approximately 90% of all lignans in *Forsythia* leaves.

The pinoresinol aglycone ratio to total pinoresinol was almost constant until September, and rapidly decreased from October (Fig. 4A). The matairesinol aglycone ratio consistently increased in April 15 (15%) to November (29%) except for a slight decrease in May (Fig. 4B). The arctigenin aglycone ratio was basically decreased in April 15 (64%) to November (24%) except for a weak (statistically insignificant) increase in May and July (Fig. 4C). It is noteworthy that arctigenin
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aglycone is initially produced at high percentage, compared to the aglycones of pinoresinol and matairesinol. These results reveal that each of the three lignan aglycones shows unique alteration in amounts, and that the arctigenin aglycone ratio most prominently varied of the three lignans throughout the seasons.

Seasonal Gene Expression Alteration in F. suspensa Leaf Subsequently, we examined alteration in gene expression of PLR, SIRD, and UGT71A18 (Fig. 1) in F. suspensa leaves from April to November (Fig. 5). PLR was constantly expressed from April to August, whereas no PLR expression was observed from September to November. SIRD showed the variable expression profiles; it was intensely expressed on April 5 and May 15 with a marked reduction on April 15, and

Fig. 2. F. suspensa Bud and Leaf from April to November

Fig. 3. Lignan Amounts of F. suspensa Leaf from April to November
(A) Pinoresinol, (B) Matairesinol, (C) Arctigenin. Open and filled circles indicate aglycone and total lignan (aglycone + glucosides), respectively. All of the lignans were assessed separately for each sample and expressed as the mean±S.E.M. of three independent experiments (*p < 0.05 vs. April 5, b p < 0.05 vs. each last month).

Fig. 4. Percentage of Aglycone/Total of F. suspensa Leaf from April to November
A) Pinoresinol, (B) Matairesinol, (C) Arctigenin. All of the percentages were assessed separately for each sample and expressed as the mean±S.E.M. of three independent experiments (*p < 0.05 vs. April 5, b p < 0.05 vs. each last month).
decreased from May to August, whereas the expression was restored in September and the expression level was consistent until November. No or markedly low expression of UGT71A18 was detected from April to July, while stable and clear expression of UGT71A18 was detected from August to November. These results indicate the specific gene expression profiles of PLR, SIRD, and UGT71A18.

**DISCUSSION**

We originally investigate seasonal alteration in the amounts of lignan aglycone and glucosides and gene expression of lignan-biosynthetic enzymes in *F. suspensa* leaves from April to November, when the leaves were still on the branches. Pinoresinol, matairesinol, arctigenin, and their glucosides were found to be produced at the maximal level from June to July (Fig. 3), when the leaves have fully grown (Fig. 2 and Table 1). Such maximal production of these lignans is in good agreement with the fact that *Forsythia* leaves for Chinese medicine, health diets and tea are harvested from June to July, given that these lignans are major pharmacologically active components.\(^{19-22}\)

Pinoresinol aglycone and its glucosides were detected at a level (1–1.6 mg/g DW) considerably less than matairesinol and arctigenin plus their glucosides (Fig. 3). Pinoresinol is a specific substrate of PLR (Fig. 1). Moreover, PLR is highly expressed from April to August (Fig. 5). These findings prove that a large part of pinoresinol is consumed for metabolism by PLR from April to August, which is consistent with the high levels of matairesinol and arctigenin during this period (Fig. 3) and with our previous study demonstrating that RNA interference against PLR in *Forsythia* leaf suspension culture cells resulted in a 20-fold greater accumulation of pinoresinol, compared with the intact cells.\(^{20}\) In contrast, PLR is not expressed from September to November, while UGT71A18 is highly expressed (Fig. 5). These lignan amount and gene expression profiles indicate that the pinoresinol metabolism is switched from the PLR-dependent lignan biosynthesis to UGT71A18-dependent glucosylation, leading to the remarkable reduction of major lignan after August.

Total matairesinol increased approximately 2-fold in June, compared to April (Fig. 3), which is in good agreement with intense expression of SIRD during this period (Fig. 5). Total matairesinol was still detected at high levels in July and August without statistically significant aglycone percentage (Figs. 3B, 4B), whereas the SIRD expression gradually decreases during this period (Fig. 5). These data suggest the relatively long stock period of matairesinol glucosides. In contrast, the SIRD expression is upregulated again (Fig. 5), whereas total matairesinol rapidly decreases after August (Fig. 3B). These phenomena are interpreted in two ways. First, matairesinol may be more predominantly consumed for the downstream lignan biosynthesis than newly biosynthesized by SIRD. However, this presumption is highly unlikely, given that arctigenin is also rapidly decreased proportionately to matairesinol (Fig. 3C). Second, newly biosynthesized matairesinol is consumed for some physiological functions specific to this period, although no physiological roles of matairesinol in plants have ever been clarified. Nevertheless, several lignans such as lariciresinol and secoisolariciresinol were found to inhibit germination, seedling, or growth,\(^{21,22}\) implying that matairesinol produced from September to November is involved in the suppression of the leaf growth of *Forsythia* or seedling of other plants around *Forsythia*.

Unlike pinoresinol or matairesinol, arctigenin aglycone is present at high level as a major final lignan product from April to July (Fig. 4). Moreover, arctigenin is most abundantly contained in *Forsythia* leaves through the season. In addition, *Forsythia* suspension culture cells fail to produce arctigenin at a detectable level.\(^{28}\) In combination, these results suggest specific biological roles of arctigenin *Forsythia*. Unfortunately, neither an arctigenin biosynthetic enzyme nor an arctigenin-glucosylating enzyme has so far been isolated in a plant. Molecular and functional characterization of these enzymes, combined with the procedure for generation of transgenic *Forsythia*,\(^{39}\) is expected to provide a crucial clue to understanding the physiological roles of arctigenin.

All of the lignans investigated in this study were accumulated at the high levels from June to August (Figs. 3, 4), when the leaves were also markedly grown (Fig. 2 and Table 1). These results suggest some potential biological roles of lignans in growth or survival of the leaves. To date, no biological roles of lignans in the enhancement of leaf growth have ever been reported. Instead, some lignans including pinoresinol have been shown to exhibit pyrethrum-synergistic or antifeedant effects on insects,\(^{33}\) supporting the view that the lignans abundantly produced during leaf growth participate in defense against defoliators, and thus, assist the growth or survival of the leaves.

Recently, sustainable production of plant secondary metabolites including lignans using plants in closed factories, which is free from climate change risks, has received increasing interest.\(^{33}\) The present data are also expected to contribute to the elucidation of the optimal conditions for efficient production of each lignan by *Forsythia* in closed plant factories. Such studies are currently in progress.

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**REFERENCES**


