Effect of Light Intensity and Light/Dark Period on Iridoids in *Hedyotis diffusa*

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The purpose of this study was to clarify the effects of light intensity and light/dark period on the concentration of iridoids in the plant *Hedyotis diffusa*, which finds use in herbal medicine. To investigate the effect of light conditions, plants were grown in a chamber with factorial treatments comprising three light/dark periods (14/10 h, 19/5 h and 24/0 h) and two light intensities (photosynthetic photon flux density of 142 and 40 μmol m⁻² s⁻¹). The epigeous dry weight was greatest with the treatments using high light intensity regardless of light period. The concentration of asperuloside (an iridoid) at 24/0 h was noticeably greater than that at 14/10 h or 19/5 h irrespective of light intensity. It was expected that the critical light period for the production of asperuloside would occur between 19 h and 24 h. The content of asperuloside was greatest at 24/0 h under high light intensity. Therefore, high light intensity and a 24/0-h light/dark period can produce high-quality *H. diffusa*.

Keywords: asperuloside, environmental control, *Hedyotis diffusa*, iridoid glycoside, lighting condition, plant factory

INTRODUCTION

Medicinal plants are used as ingredients for crude drugs and herbal medicines. These medicines are traded at inflated prices for traditional primary medical care worldwide; demand for these preparations is increasing and exist throughout the year. In addition, as these plants are used for medicinal purposes, a high-quality and safe product is required. However, most medicinal plants are only available in the wild because methods for their cultivation have not been established. Therefore, it is difficult to provide high-quality and safe medicinal plants throughout the year. In Japan, medicinal plants are imported from foreign countries, with China being a major supplier (Japan Kampo Medicine Manufacturers Association, Crude Drug Commission, 2010). With the increasing demand in Japan for the production of high-quality medicinal plants, methods to cultivate the plants must be developed.

A plant factory is a year-round production system that produces a high-quality, safe and steady supply of vegetables using technological environmental controls and automation. However, such systems for the production of plants are costly and consume considerable energy, which hamper their profitability. This limitation might be resolved by cultivating high-value crops that contain useful substances in higher quantities than those present in crops cultivated in open fields.

*Hedyotis diffusa* (family Rubiaceae) is used in Chinese herbal medicine for treating cancer, urinary infection, appendicitis and bronchitis. This plant’s natural distribution is in eastern Asia in subtropical and tropical zones. The availability of *H. diffusa* depends on harvests of wild crops. A problem encountered in China during harvesting of *H. diffusa* is contamination with other similar cultivars because it is grown with other plants and weeds (Lin et al., 1987; Wee and Keng, 1990). Such contamination, which degrades the quality of herbal medicine, could be avoided if the plant were to be cultivated in a plant factory. The environmental control technology in a plant factory would produce a better quality product than that produced in the open field. However, to date, no cultivation system for *H. diffusa* has been clarified.

Iridoids are an active component in aerial part of *H. diffusa*, existing as iridoid glycosides. *H. diffusa* contains various iridoid glycosides, each one of which has been shown to exhibit different medicinal effects (Niu and Meng, 2013). In particular, numerous iridoid glycosides having an anti-oxidative activity have been reported (Lu et al., 2000; Kim et al., 2005). It is considered that iridoid glycosides exist in the vacuole or cytoplasm of plant cell (Kamata, 2009). Figure 1 shows the biosynthetic pathway of iridoid glycosides (The Pharmaceutical Society of Japan, 2005; Shioi et al., 2009; Takai et al., 2010). The biosynthesis of iridoid glycosides begins with glucose and a promotion of photosynthesis increases the starting material of iridoid glycoside. In addition, biosynthesis of medicinal properties could be enhanced by environmental stimulation to induce protective response of plants (Kozai, 2012). It was commonly reported that a content of anti-oxidant has...
Coumaroyl scandoside methyl ether, which has anti-oxidant activity (Kapadia et al., 1996; Lu et al., 2000; Liang et al., 2008). Because of anti-cancer effects of asperuloside, it showed anti-cancer effects, anti-oxidant effects and anti-angiogenic effects (Lu et al., 2000). The concentrations of these components were enhanced by the increase in light intensity. However, the effect of the light/dark period differed for each iridoid glycoside. Concentrations of asperuloside increased with increasing light period, whereas those of 6-O-p-coumaroyl scandoside methyl ether decreased. It would be difficult to increase the content of all iridoid glycosides by specific environmental conditions. Therefore, we selected asperuloside to limit the scope of quality evaluation conducted in this study. Although only relative concentrations were measured in our previous study (Higashiuchi et al., 2014), absolute concentrations for asperuloside were measured by another extracting method and quantitative analysis in the present study. In addition, the effects of light conditions on growth and absolute content for asperuloside were evaluated.

**MATERIALS AND METHODS**

**Raising seedlings**

In this study, raising the seedlings from scions rather than from seeds shortened the cultivation time by a maximum of 10 d. Scions were harvested from the mother plants, which were hydroponically cultured for 6 weeks in a closed chamber, i.e. former 2 weeks for raising the seedlings under 13 μmol m⁻² s⁻¹ photosynthetic photon flux density, 400 μmol mol⁻¹ CO₂ concentration and 70% RH humidity at 14/10 h photoperiod and 25°C air temperature and latter 4 weeks for cultivation under 112 μmol m⁻² s⁻¹ photosynthetic photon flux density at 14/10 h photoperiod and 25°C air temperature to standardize the quality of scions. The seedlings for experimental treatments were grown from scions that were harvested from the mother plants. The seedlings were prepared as shown in Fig. 2. Each 45 mm scion was cut from the top of the mother plant. Four leaves with a midrib length longer than 10 mm were retained on the scion. The cut end of each scion was inserted at a 15 mm depth into a polyurethane foam cube (23.5 × 23.5 × 30 mm) and then placed in a transparent plastic tray (NP box #7; Astage, Niigata, Japan). A 500 mL solution of half-strength Otsuka A prescription (OAT House No. 1 and No. 2; OAT Agrio, Tokyo, Japan) was added to 12 scions in the tray. After covering the trays with polyethylene film to suppress transpiration, they were placed in a growth chamber (NC200SC; Nippon Medical & Chemical Instruments Co., Osaka, Japan). The 48 seedlings (12 seedlings × 4 plastic trays) were grown for 2 weeks in the growth chamber under the conditions shown in Table 1. The photosynthetic photon flux density was measured at the tray base using a quantum metre (Meter: Light Meter LI-250, Sensor: LI-190SA; LI-COR, Lincoln, NE, USA). The light was provided by fluorescent lamps (FL15EX-N-HG, FL20EX-N-HG and GL40EX-N-HG; NEC Lighting, Tokyo, Japan). To suppress transpiration, the photon flux density was set below that used for cultivation. The solution was replaced once a week, and after 1 week, the polyethylene film was nicked by cutting with a knife to acclimatize the seedlings to a lower humidity.

![Synthesis pathway of iridoid glycosides.](image-url)
**Cultivation**

The 20 experimental seedlings were transplanted into the cultivation plate (240 × 360 mm), which was floated onto the 10 L culture solution in the blue box (NF box #13; Astage, Niigata, Japan) to suppress algae growth and then cultivated for 3 weeks in a growth chamber (BAC-130H; Especmic, Aichi, Japan). The culture solution was the same as that used for growing the seedlings. Air was pumped into the solution using an air pump (W-600; Nisso Industry, Osaka, Japan) and an air stone (LS-150A; Kainuma-sangyo, Aichi, Japan).

Three different light/dark period treatments (14/10 h, 19/5 h and 24/0 h) were used, and for each of these, two different light intensity treatments were used (photosynthetic photon flux density 142 μmol m⁻² s⁻¹ and 40 μmol m⁻² s⁻¹) for a total of six treatments. Photosynthetic photon flux density was decreased compared to previous study because of aged deterioration of fluorescent lamps. The growth chamber was divided into two parts, i.e. the upper for the high-intensity light treatments and the lower for the low-intensity light treatments. The inside glass wall of the lower space was covered with black cheesecloth to regulate the photon flux density. Table 2 presents the cultivation conditions. Earlier study demonstrated that drying stress and strong light stress produced by a rapid increase in light intensity and a sudden decrease in humidity during the transition from raising seedlings to transplantation led to poor growth and a reduced iridoid glycoside content (Hisano et al., 2013). Therefore, in the present study, the photon flux density and humidity were changed gradually to reduce the drying and light stresses. Twenty samples were cultivated for each treatment, and the solution was replaced once a week.

**Harvesting and drying**

The epigeous fresh weight was measured after harvest using an electric balance (GF-3000; A&D Company, Tokyo, Japan). Owing to the risk of denaturation of the medicinal properties of plants by high-temperature drying, the samples were dried naturally at 25°C for 7 d after weighing. After drying, the epigeous dry weight was measured. Then, 20 samples were preserved at −20°C in a freezer (KGT-4056HC; NIHON FREEZER, Tokyo, Japan) until component analysis.

**Analysis of H. diffusa components**

A sampling of 0.5 g H. diffusa was repeated 4 times, and each samples contained 3 or 4 plants. Then, the asperuloside concentration was measured by liquid chromatography-mass spectroscopy (LC/MS). Therefore, for each

<p>| Table 1 | Conditions for raising H. diffusa seedlings for experimental treatments. |</p>
<table>
<thead>
<tr>
<th>Factor</th>
<th>Date</th>
<th>Set value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light/dark period (h/h)</td>
<td>1–14 d</td>
<td>14/10</td>
</tr>
<tr>
<td>Air temperature (°C)</td>
<td>1–14 d</td>
<td>25</td>
</tr>
<tr>
<td>Relative humidity (%)</td>
<td>1–14 d</td>
<td>70</td>
</tr>
<tr>
<td>Photosynthetic photon flux density (μmol m⁻² s⁻¹)</td>
<td>1–14 d</td>
<td>13±3.5</td>
</tr>
<tr>
<td>CO₂ concentration (μmol m⁻² s⁻¹)</td>
<td>1–14 d</td>
<td>400</td>
</tr>
</tbody>
</table>

* Dates indicate days after transplanting.
* Mean ± standard deviation measured at the bottom of the plastic tray.

<p>| Table 2 | Conditions for the cultivation of H. diffusa. |</p>
<table>
<thead>
<tr>
<th>Factor</th>
<th>Date</th>
<th>Set value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air temperature (°C)</td>
<td>1–21 d</td>
<td>25</td>
</tr>
<tr>
<td>Relative humidity (%)</td>
<td>1–7 d</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>8–21 d</td>
<td>70</td>
</tr>
<tr>
<td>Photosynthetic photon flux density (μmol m⁻² s⁻¹) at high-intensity treatment</td>
<td>1–4 d</td>
<td>35.7±1.01</td>
</tr>
<tr>
<td></td>
<td>5–7 d</td>
<td>119±4.70</td>
</tr>
<tr>
<td></td>
<td>8–21 d</td>
<td>142±3.14</td>
</tr>
<tr>
<td>Photosynthetic photon flux density (μmol m⁻² s⁻¹) at low-intensity treatment</td>
<td>1–4 d</td>
<td>32.4±2.61</td>
</tr>
<tr>
<td></td>
<td>5–7 d</td>
<td>38.9±1.62</td>
</tr>
<tr>
<td></td>
<td>8–21 d</td>
<td>39.9±1.37</td>
</tr>
<tr>
<td>CO₂ concentration (μmol mol⁻¹)</td>
<td>1–21 d</td>
<td>400</td>
</tr>
</tbody>
</table>

* Dates indicate days after transplanting.
* Mean ± standard deviation measured at the surface of the cultivation plate.
treatment, the sample number of asperuloside concentrations measured was four.

Deionized water was used for liquid extraction, and Milli-Q water was used for LC/MS. Methanol (LC/MS grade) was purchased from J. T. Baker (Pennsylvania, USA), and formic acid (LC/MS grade) was purchased from Sigma-Aldrich (Missouri, USA). Asperuloside was purchased from Standhill Technology (Hong Kong) and was used as a reference standard. The D101 macroporous adsorption resin was purchased from China.

Dried and powdered samples (0.5 g) of *H. diffusa* were extracted twice with 5 mL of deionized water under reflux; each extraction lasted 1 h. After filtration of the aqueous fraction, the filtrate was concentrated to 2 mL. The filtrate was concentrated to dryness by rotary evaporator (RE111; Buchi, Switzerland) and re-dissolved with 2 mL of deionized water. The concentrated extracts were then subjected to D101 macroporous adsorption resin and eluted with water and different concentrations of ethanol. Thirty-five percent ethanol fraction was concentrated to dryness under reduced pressure. The dried fraction was dissolved in 2 mL of water and filtered through a membrane filter (0.22 μm) prior to analysis by high-performance liquid chromatography-electrospray ionization-mass spectroscopy (HPLC/ESI/MS).

HPLC measurements were performed using Agilent 1290 Infinity liquid chromatography (Agilent Technologies, USA), comprising an ultra-HPLC binary pump, an autosampler, a column oven and a diode array detector. Separation was performed on a C18 column (250 mm, 4.6 mm internal diameter, 5 μm; GRACE Alltima, USA), and the column temperature was set at 30°C. The eluent system comprised A [0.1% (v/v) formic acid in aqueous phase] and B [0.1% (v/v) formic acid in methanol] with a flow rate of 1.0 mL min⁻¹, which was performed with a linear gradient: 0–20 min, 7–15% B; 20–30 min, 15–19% B; 30–35 min, 19% B and 35–45 min, 19–95% B.

All MS experiments were conducted using a quadrupole time-of-flight mass spectrometer equipped with an ESI interface (Agilent Technologies, USA). Nitrogen was used as the drying gas with a flow rate of 12 L min⁻¹. The drying gas temperature was set at 350°C, and the nebulizer pressure was set at 60 psi. Spectra were recorded in positive ion mode at a spray voltage of 3500 V. The mass scan range was between 50 and 1700 m/z.

Asperuloside content was calculated as a product of the average asperuloside concentration and epigeous dry weight. Statistical tests were conducted using the non-parametric significant difference test. The Wilcoxon test method was adopted for comparison among 2 light intensity treatments, and the S method was adopted when multiple comparisons were performed for comparing among the 3 light/dark period treatments (Shirahata, 1987).

**RESULTS AND DISCUSSION**

Images of the harvested plants for each treatment are shown in Fig. 3. A spindly growth or color degradation was not observed at 14/10 h under low light intensity. Therefore, it was considered that 40 μmol m⁻² s⁻¹ photosynthetic photon flux density was larger than that of light compensation point for *H. diffusa*. The plants grown under high light intensity appeared to be larger than those grown under low light intensity. In contrast to previous research, no noticeable continuous light injury (chlorosis or necrosis) was observed under a continuous light treatment. Light injury results from the accumulation of radical oxygen. In addition, the degree of injury depends on the light intensity and the plant’s anti-oxidative enzymatic activities (Hata et al., 2011; 2012). It was expected that radical oxygen beyond the capacity of anti-oxidative enzymatic activities of *H. diffusa* was not produced even under continuous light with 142 μmol m⁻² s⁻¹ photon flux density; moreover, no light-induced injury occurred.

Figure 4 shows the measurements of epigeous fresh weights, dry weights and percentage dry matter in the 6
light treatments. These measurements show a trend similar to the results obtained in a previous study (Higashiuchi et al., 2014). The mean epigeous fresh weights from the high- and low-intensity light treatments in each light/dark period were significantly different at a 1% significance level. It seems that high light intensity promotes the growth of these plants. Among the light/dark periods, at high or low intensity, the epigeous fresh weights at 19/5 h and 24/0 h were greater than those at 14/10 h. Therefore, the results suggest that a photoperiod of longer than 19 h is required to facilitate plant growth.

The mean epigeous dry weights and percentage dry matter from the high- and low-light intensity treatments in each light/dark period were significantly different at a 1% significance level. These results indicate the presence of more carbon assimilation products under high light intensity than those under low light intensity. Consequently, high light intensity promoted the photosynthetic rate of *H. diffusa*. When the dry weights of plants grown under three different light/dark periods were compared, the dry weight tended to increase as the light period increased. In general, under a long light period, feedback inhibition with starch accumulation or photoinhibition with radical oxygen elicits reduction in photosynthetic rate (Vermeglio et al., 1983; Teng et al., 2006). However, no light injury was observed in this study. Therefore, it seemed that photosynthesis was not inhibited under the light condition. Because of the increase in the time capable of photosynthesis under a long light period, the amount of photosynthetic products produced in a day increased. Therefore, the long light period promoted the generation of photosynthetic products.

The mean percentage dry matter was markedly increased at 24/0 h. It has been reported that continuous light can maintain transpiration of *Metasequoia glyptostroboides* and increase the water loss during 24 h (Yang et al., 2009). Therefore, it seemed that output of water augmented by continuous light led to the increase in percentage dry matter of *H. diffusa* in the 24/0 h light/dark condition.

Figure 5 shows the relationship between the epigeous fresh weight or epigeous dry weight and accumulated photon flux density in a day. The accumulated photon flux density was calculated by multiplying the photoperiod by the photon flux density on or after the eighth day after transplantation. Epigeous fresh weight increased with the increase in accumulated photon flux density up to approximately 10 mol m⁻² d⁻¹ and decreased at approximately 12 mol m⁻² d⁻¹. Existence of peak value in the fresh weight was recognized. This relationship could be approximated by a quadratic function with a high value of coefficient of determination, and a maximum fresh weight at 10.5 mol m⁻² d⁻¹ was derived from the approximated curve. Epigeous dry weight linearly increased with the increase in accumulated photon flux density. It was inferred that the accumulated photon flux density that maximized the epigeous dry weight would be greater than the maximum
value obtained in this study.

Figure 6 shows a chromatogram from one of the samples (Fig. 6A) and the asperuloside reference standard (Fig. 6C). The concentrations of asperuloside in *H. diffusa* were simultaneously quantified by a validated method. A calibration curve (ranging from 3.125 μg mL⁻¹ to 100 μg mL⁻¹ with [M+Na]⁺ at 437 m/z) with a good determination coefficient (R² > 0.998) was constructed using MS peak areas of the reference standards versus their concentrations (Fig. 6B and C). The intra- and inter-batch precision was satisfied with a relative standard deviation of < 7.8% and an accuracy of 98.7–108.5%.

Figure 7 presents the mean asperuloside concentrations on the basis of dry weights and asperuloside contents of *H. diffusa*. The mean asperuloside concentration was significantly greater under high light intensity than that under low light intensity with respect to each light/dark period at a 5% significance level. Higher dry weights and higher concentrations of asperuloside were obtained from the condition of high light intensity. In other species, an increase in the secondary metabolite concentration associated with increasing photosynthetic photon flux density was reported (Mosaleeyanon et al., 2005). It seems that high light intensity promoted photosynthesis of *H. diffusa*, which produced glucose used for biosynthesis of secondary metabolites. In addition, high intensity light would enhance biosynthesis of the asperuloside. Therefore, the asperuloside concentration increased under high light intensity treatments. As for the comparison among the light/dark periods, the asperuloside concentration at 24/0 h was noticeably higher than that at 14/10 h or 19/5 h. Figure 8 shows the relationships among asperuloside concentration (Fig. 8A), asperuloside content (Fig. 8B), asperuloside

Fig. 6  Chromatograms of a sample. Total ion chromatogram (TIC) of *Hedyotis diffusa* (A); 437 m/z extracted ion current (EIC) of *H. diffusa* (B); 437 m/z EIC of the asperuloside standard compound (C).
Asperuloside content increased exponentially with the increase in accumulated photon flux density (Fig. 8B). High light intensity and a 24/0 h light/dark photoperiod were able to maximise the epigeous dry weight and asperuloside concentration; therefore, the content of asperuloside was the greatest at 24/0 h under high light intensity. In addition, the efficiency of asperuloside production per unit accumulated photon flux density in a day was highest at 24/0 h under high light intensity (Fig. 8C). Therefore, the increase in dry weight and the promotion of asperuloside biosynthesis are both necessary for efficient production of a high asperuloside content in *Hedyotis diffusa*. Consequently, high light intensity treatment (142 μmol m⁻² s⁻¹) and a 24/0 h light/dark photoperiod could promote high concentration of asperuloside and yield of *H. diffusa*.

**CONCLUSION**

High light intensity treatment (142 μmol m⁻² s⁻¹) and 24/0 h light/dark photoperiod successfully maximised the content of asperuloside because these light conditions induced both high epigeous dry weight and asperuloside concentration. In some cases, a suitable condition for the biosynthesis of secondary metabolites is not suitable for plant growth. However, in the case of *H. diffusa*, an optimal lighting condition, which can promote both growth and biosynthesis of asperuloside was specified. It seems that in *H. diffusa*, the production of anti-oxidants, including asperuloside, are affected profoundly by light period. The control of photoperiod is useful for an increase in concentration of medicinal properties. In addition, cultivation under an optimum accumulated photon flux density can obtain the highest yield of dry matter. The lighting conditions optimised both light period and accumulated photon flux density, making it possible to maximise efficiently the content of medicinal properties in plants. The results of this study could improve the profitability of plant factories producing *H. diffusa*.

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