Isolation of isolactarane sesquiterpenes from a *Phlebia tremellosa* culture filtrate and their growth promotion effects on lettuce roots

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The ethyl acetate extract of the culture filtrate of *Phlebia tremellosa* promoted elongation of the lateral roots of lettuce seedlings at 250 µg/mL. We purified two compounds that promote root elongation by using activity-guided chromatographic fractionation. On the basis of spectroscopic analyses, these compounds were identified to be isolactarane sesquiterpenes derived from the dehydrogenation of merulactone, which was previously isolated from the same species. We named the purified compounds phlelactones A and B. Phlelactones A and B promoted primary root elongation at 100–300 and 10–30 µg/mL and the elongation and formation of lateral roots at 300–1000 and 30–100 µg/mL, respectively. © Pesticide Science Society of Japan

Keywords: *Phlebia tremellosa*, plant growth regulator, root development, isolactarane, phlelactone.

Electronic supplementary material: The online version of this article contains supplementary material (Supplemental Tables S1 and S2, and Supplemental Figs. S1–S5), which is available at http://www.jstage.jst.go.jp/browse/jpestics/

Introduction

Plant growth regulators are compounds with biological activities such as the promotion of plant growth, acceleration of matura-

tion, stimulation of fruit coloration and enhancement of toler-
ance to biological and environmental stresses. Plant growth reg-

ulators have attracted increasing attention because of concerns

regarding the adverse effects of global climate change on crop production. Various substances including plant hormones, inor-

ganic compounds, extracts from plants, fermentation products,

and synthetic chemicals, have been shown to have these activi-

ties, and categorized as plant growth regulators.1,2)

Mushrooms are fungal species that form visible fruiting bod-

ies and are considered valuable sources of novel bioactive com-

pounds. The Fungus/Mushroom Resource and Research Cen-

ter (FMRC) at Tottori University has collected and maintained

8,387 mushroom strains belonging to 1,400 species (as of March

2017), which form one of the largest mushroom culture collec-

tions in the world. We have made an extract library of the mush-

rooms. Using this library, we initiated an investigation of the

bioactive compounds present in these strains and have identified

several antimicrobial compounds3,4) and new tyrosinase inhibi-

tors.5) These findings strongly suggested that there remain many

opportunities to discover new specialized metabolites with bio-

logical activity in mushrooms.

Recently, compounds with plant growth-promoting and

-suppressing activities were isolated from mushroom cultures.

Some mushroom species have been known to form a circular

arrangement of fruiting bodies called “fairly rings” on grasslands

and to promote or suppress the growth of surrounding grasses.6)

From the culture filtrate of the fairy-ring-forming mushroom

*Lepista sordida* (Schumach.) Singer, 2-azahypoxanthine (AHX)

and imidazole-4-carboxamide (ICA) have been isolated, and

they showed plant growth-promoting and -suppressing activi-

ties, respectively.7,8) The application of these compounds, which

have been referred to as “fairy compounds,”9) increased the

yields of rice10) and wheat11) in the field.

On the basis of these findings, we screened extracts derived

from the fruiting bodies and liquid cultures of FMRC-preserved

mushroom strains in an attempt to identify novel plant growth

regulators. Hymexazol (3-hydroxy-5-methylisoxazole) and iso-

prothiolane (diisopropyl 1,3-dithiolane-2-ylidenemalonate),

which are known plant growth regulators that increase the yield

of rice, have also been shown to promote root development.12,13)

Thus, we have focused on their promotion of root development

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in plants. Through a screening of extracts of the culture filtrates, mycelia, and fruiting bodies of mushrooms, we found that the extract of the culture filtrate of *Phlebia tremellosa* (Schrad.) Nakasone & Burds. (syn. *Merulius tremellosus* Schrad.) showed this activity. *P. tremellosa* is a white-rot fungus that is distributed in tropical to subarctic zones worldwide. Various isolactarane and sterpurane sesquiterpenes, such as merulidial, merulactone, and tremetriol, have been isolated from the culture filtrates of *P. tremellosa*.14–17) Among them, merulidial has shown antimicrobial activity, mutagenesis activity, and cell toxicity.14,16,18) In the present study, the compounds that promote lettuce root development were isolated from the culture filtrate of *P. tremellosa* and identified.

### Materials and Methods

1. **General**

   1H and 13C NMR spectra and 2D spectra (COSY, HMQC, and HMBC) were recorded using an Avance II instrument (Bruker, Billerica, MA, USA). High-resolution mass spectra were measured using an Exacte mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA), and ESI-MS was conducted using a Quattro Micro API mass spectrometer (Waters, Milford, MA, USA) connected to an Acquity UPLC instrument (Waters).

   GC-MS analysis was performed using a GCMS-QP2010 Plus (Shimadzu, Kyoto, Japan) equipped with an Rtx-5MS column (30 m × 0.25 mm, 0.25 µm thickness, GL Sciences, Tokyo, Japan).

   Preparative HPLC was performed with a 10A HPLC system (Shimadzu).

2. **Bioassay**

   A single layer of filter paper was placed in the wells of a 24-well plate, and 300 µL of solutions of mushroom extracts in 1% DMSO was added to the wells. The extract concentration was fixed at 250 µg/mL. In the experiments with fractions of extracts of the culture filtrates from *P. tremellosa*, the samples were dissolved in methanol. The methanol solutions were added to the wells; methanol was removed by air-drying, and then 300 µL of distilled water was added. Four lettuce (*Lactuca sativa* L. cv. Melbourne MT) seeds were placed in a well and incubated at 28°C with 16 hr/8 hr light and dark cycles. After incubation for six days, images of the seedlings were obtained using a scanner (CanoScan LiDE 220, Canon, Tokyo, Japan), and the lengths of the primary and lateral roots and the number of lateral roots were measured using Image J (https://imagej.nih.gov/ij/docs/install/windows.html) software.

### Table 1. 1H (600 MHz) and 13C NMR (150 MHz) data of 1, 2 and merulactone in CDCl3.

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week incubation, the mycelia were inoculated to a malt extract liquid medium (200 mL/flask, 25 flasks) consisting of glucose (30 g/L), peptone (3 g/L), the extract from 50 g/L of malt, and tap water. The cultures were further incubated for 1–3 months and separated into mycelia and culture by filtration. The mycelia were air-dried and extracted by methanol three times. The extracts were combined and evaporated as a mycelium extract. The culture filtrate was partitioned with ethyl acetate three times. The ethyl acetate layers were combined, dried over Na$_2$SO$_4$ overnight, evaporated, and used as a culture filtrate extract.

The fruiting bodies obtained through a mycelial block cultivation method were freeze-dried and well powdered. The resulting powder was successively extracted using ether, dichloromethane, methanol, and water. The obtained extracts were evaporated to dryness and subjected to screening.

4. Liquid culture of $P$. tremellosa

$P$. tremellosa (TUFC11737) is the preserved strain of the FMRC at Tottori University (http://fungusdb.muses.tottori-u.ac.jp/searches/index/en). The strain was obtained from Saihaku-gun in Tottori Prefecture in October 2008. The strain was cultured without shaking at 25°C for 47 days in the dark in 500 mL conical flasks containing a malt extract liquid medium (200 mL/flask, 25 flasks).

5. Purification of active compounds

The culture filtrate (5 L) of $P$. tremellosa was extracted with ethyl acetate (1.7 L) three times. The extract was dried over Na$_2$SO$_4$ overnight and evaporated to dryness. The extract (0.81 g) was subjected to column chromatography using silica gel (85 g, Daisogel IR-60-63/210, Daiso, Osaka, Japan). The column was eluted with mixtures of acetone and hexane. The concentration of acetone was increased from 0 to 60% in increments of 10% with the volume of each fraction being 650 mL. Then the column was eluted with 100% acetone and washed with methanol in volumes of 650 mL. Activity was detected in the 20% acetone fraction (324 mg).

An aliquot of the 20% acetone fraction (289 mg) was subjected to silica gel column chromatography (28.5 g, Daisogel IR-60-63/210). The column was eluted with mixtures of ethyl acetate and hexane. The concentration of ethyl acetate was increased from 20 to 60% in increments of 10%, with the volume of each mixture being 285 mL. Activity was detected in the 50% (17.8 mg) and 60% (60.8 mg) ethyl acetate fractions.

The 60% fraction was subjected to preparative TLC [SiO$_2$ glass plate, 60 F254 silica gel, Merck, Darmstadt, Germany; solvent, ethyl acetate–n-hexane (4:1, v/v)] and divided into six fractions. Fraction 5 corresponded to the UV-active band at $R_f$ values of 0.37–0.46. We subjected this fraction to preparative HPLC because it was active. The HPLC conditions were as follows: column, Cosmosil 5C$_18$-AR-II 10 mm ID×250 mm (Nacalai Tesque, Kyoto, Japan); column temperature, 40°C; flow rate, 3 mL/min; solvent, 30% acetonitrile in water; detection, 280 nm. The major peak was eluted at 4.01 min. The fraction corresponding to this peak was concentrated to yield 1.

Compound 1. 5.9 mg. Positive ESI-MS $m/z$: 247 [M+H]$^+$. HR-
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EI-MS (70 eV) m/z (rel. int.): 246 (35) [M]+, 231 (33), 217 (31), 203 (28), 189 (40), 173 (24), 159 (91), 157 (46), 145 (57), 143 (51), 131 (44), 115 (42), 105 (44), 91 (100), 77 (41), 43 (100).

CD λmax (Δε): 225 (−0.030), 266 (−0.025), 280 (−0.026), 342 (0.0073). NMR data are shown in Table 1 and Figs S1 and S2.

The 50% fraction (17.8 mg) was subjected to preparative TLC [SiO2 glass plate, 60 F254 silica gel, Merck; solvent, ethyl acetate–n-hexane (4 : 1, v/v)] and divided into six fractions. The activity was detected in fraction 5, which corresponded to the UV-active band at Rf values of 0.40–0.51. This fraction was subjected to preparative HPLC under the same conditions as for 1. The major peak was eluted at 4.70 min. The fraction corresponding to this peak was concentrated to yield 2.

Compound 2. 9.7 mg. Positive ESI-MS m/z: 247 [M+H]+. HR-ESI-MS m/z: 246.1326 [M+H]+ (Calc. for C15H19O3: 247.1334). CD λmax (Δε): 245 (−0.022), 263 (−0.019), 283 (−0.021), 349 (0.0078). EI-MS (70 eV) m/z (rel. int.): 246 (48) [M]+, 231 (39), 217 (75), 203 (51), 173 (72), 159 (54), 131 (54), 119 (60), 105 (76), 91 (100), 79 (58), 77 (74), 43 (90), 41 (73). NMR data are shown in Table 1 and Figs. S2 and S3.

Chemical shifts of 1H and 13C NMR data of 1 and 2 are summarized in Fig. S5.

**Results and Discussion**

1. Isolation of compounds

We examined the root growth promotion activities of the extracts of the mycelia and culture filtrates of 33 mushrooms (Table S1) and the extracts of the fruiting bodies of 8 mushrooms (Table S2) at 250 μg/mL. The extract of the culture filtrate and mycelium of *P. tremellosa* exhibited activity, as shown in Fig. 1. Treatment with the extract of the culture filtrate did not affect the length of the primary root (Fig. 1A) but did enhance the elongation of the lateral roots (Fig. 1B) and increase the number of lateral roots (Fig. 1C). On the other hand, the mycelium extract slightly enhanced the elongation of the primary root but did not affect lateral root development. Next, we cultured *P. tremellosa* on a large scale to isolate the active compounds. The filtrate of the *P. tremellosa* culture was extracted and subjected to activity-guided fractionation by column chromatography, TLC and HPLC. Finally, we obtained two active compounds, 1 and 2.

2. Determination of the chemical structures of 1 and 2

The positive ESI-MS analyses of 1 and 2 indicated that they each had a molecular weight of 246. The respective HR-ESI-MS data for 1 and 2 showed protonated molecule peaks at m/z 247.1324 and m/z 247.1326, indicating that their molecular formula were C15H19O3, and thus their hydrogen deficiency indices were 7. Because the isolactarane sesquiterpene merulactone (C15H20O3, M.W. 248, Fig. 1A) was previously isolated from the culture filtrate of this species,17) we speculated that 1 and 2 may be dehy-

Fig. 3. Analysis of the chemical structure of 2. (A) COSY (bold line), HMBC (black arrows from carbon to proton), and NOESY (blue arrows from proton to proton) correlations of 2. HMBC correlations from C-8, C-9, and C-10 are indicated. (B) NOESY correlations on three-dimensional chemical structure of 2. (C) Chemical structure of phlelactone B. The absolute stereochemistry of 2 was not determined. The indicated structure was constructed assuming that the stereochemistry of the 6 and 7 positions was the same as that of merulactone.

Fig. 4. Lettuce seedlings germinated in distilled water (A) and solutions of phlelactone A at 300 μg/mL (B) and phlelactone B at 30 μg/mL (C). The lettuce seeds were sown in the solutions of the phlelacetones and incubated for six days.
drogenated compounds derived from it.

A comparison of the $^{13}$C NMR spectrum of 1 with that of merulactone (Table 1) revealed that the chemical shifts of the signals other than the signal at $\delta_C 80.5$ of 1 were largely unchanged compared with those of merulactone. Thus, the oxygen at the 8 position is likely connected to the carbon corresponding to this signal. This carbon was assumed to be C-1 or C-10 because all of the other $^{13}$C NMR signals were assigned on the bases of $^1$H and $^{13}$C NMR and 2D NMR. In the HMBC spectrum, the signal of C-9 at $\delta_C 41.5$ showed correlations with the $^1$H NMR signals at $\delta_H 2.04$ and $\delta_H 1.23$ (H-10) (Fig. 2B). The H-10 protons were connected to the carbon corresponding to the signal at $\delta_C 42.6$; hence, this signal was assigned to C-10. Accordingly, the signal at $\delta_C 80.5$ was assigned to C-1. Thus, we concluded that 1 is the compound in which the carbons corresponding to C-1 and C-8 in merulactone are bridged by an oxygen atom.

The relative stereochemistry of 1 was determined using its NOESY spectrum (Fig. 2B and 2C). The cross peaks of H-8/H-10, H-8/H-9, and H-5/H-9 indicated that the oxygen bridge of 1 exists on the opposite side of the cyclopropane ring. As 1 has not been reported to our knowledge, we referred to 1 as phlelactone A (Fig. 2D).

A comparison of the $^1$H and $^{13}$C NMR spectra of 2 (Table 1) with those of merulactone suggested that the oxygen atom at the 8 position is connected to the carbon corresponding to the signal at $\delta_C 80.4$, which is assigned to C-1 or C-10, again because the chemical shifts of the other signals were similar to those of merulactone. The HMBC spectrum showed a correlation between C-9 at $\delta_C 40.8$ and H-10 at $\delta_H 1.86$ and 1.48, which are connected to the carbon corresponding to the signal at $\delta_C 41.8$ (Fig. 3A). Thus, this carbon was identified as C-10. Accordingly, the carbon signal at $\delta_C 80.4$ was assigned to C-1, and C-8 and C-1 of 2 are connected by an oxygen bridge, similar to 1.

The relative stereochemistry of 2 was determined using its NOESY spectrum (Fig. 3B and C). The spectrum showed a cross peak for H-8/H-9 but not for H-5/H-9. Thus, the oxygen bridge was considered to be present on the same side as the cyclopropane ring, as shown in Fig. 3C. This compound has not been reported to our knowledge, so we referred to 2 as phlelactone B.

Phlelactones A and B are isomers of each other and classified...
as isolactarane sesquiterpenes. From *P. tremellosa*, isolactarane sesquiterpenes such as merulidial and merulactone, as well as a biosynthetically related sterpurane sesquiterpene, temretiol, have been identified.\(^1\) The phlelactones are likely biosynthesized via a common pathway to those of these sesquiterpenes although we cannot exclude the possibility that the phlelactones were formed from merulactone or related compounds by a non-enzymatic oxidation reaction during the long culturing period. The absolute stereochemistry of phlelactones A and B was not determined. Assuming that the stereochemistry of C-6 and C-7 of phlelactones A and B are the same as those of merulactone, the stereochemistry at C-8 of phlelactone A is the same as that of merulactone, but that of phlelactone B is different. No compound with the same stereochemistry at that at C-8 of phlelactone B has been found in *P. tremellosa*.

3. *Growth promotion effects of phlelactones A and B on primary and lateral lettuce roots*

As shown in Fig. 4, both phlelactones A and B showed promoting effects on the development of lettuce roots. Phlelactone A enhanced primary root growth at 100 and 300 µg/mL with approximately 2.8-fold increases, while it inhibited root elongation almost completely at 3,000 µg/mL (Fig. 5A). The compound promoted the elongation and generation of lateral roots at 300 µg/mL but was inhibitory at higher concentrations (Fig. 5B). Phlelactone B promoted primary root elongation at lower concentrations than phlelactone A (Fig. 5A). At higher concentrations, phlelactone B also inhibited primary root elongation. The most effective concentration of phlelactone B for the elongation of lateral roots was 100 µg/mL (Fig. 5B). Treatments with phlelactones A and B tended to increase the number of lateral roots at the lower concentrations, although the effect was statistically significant only at 300 µg/mL of phlelactone A.

The crude extract of the *P. tremellosa* culture filtrate showed promotion activity on root development at 250 µg/mL. Pure phlelactones A and B showed the marked activity at 100–300 µg/mL. Thus, the activity in the crude extract could not be accounted for by phlelactones A and B alone; other active compounds may be present in the extract.

The initiation and development of roots are controlled by various endogenous signals.\(^{20}\) Auxin is known to be the central player that controls root development. A high level of auxin induces lateral root growth and represses primary root elongation. Lateral roots are more sensitive than primary roots to exogenously applied auxin.\(^{20}\) Along with auxin, cytokinin\(^{21,22}\) and abscisic acid\(^{23,24}\) affect root development. Recently, the mode of action of isoprothiolane, which shows root growth promotion effects on plants, was analyzed in Arabidopsis by using plant hormone-related mutants and chemical inhibitors of plant hormones.\(^{25}\) The analysis showed the involvement of auxin, jasmonic acid, and ethylene in the root growth promotion by isoprothiolane. To determine the mode of action of the phlelactones, it is important to investigate their effects on the hormonal control of root development.

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**References**