Release of acid phosphatase from extraradical hyphae of arbuscular mycorrhizal fungus *Rhizophagus clarus*

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Running title: Acid phosphatase from extraradical hyphae
Abstract
Arbuscular mycorrhizal (AM) fungi enhance plant uptake of available phosphorus (P) from soil through their extraradical hyphae. The mechanism underlying this P uptake enhanced by AM fungi is the increase in the surface area for absorption of available P. Little is known about utilization of unavailable P by AM fungi. We investigated whether extraradical hyphae of AM fungi release ACP. Sterilized Andosol was packed in pots that were separated into the mycorrhizal and hyphal compartments with a nylon net of 30 µm pore size. Seeds of *Allium fistulosum* L. were inoculated or uninoculated with the AM fungus *Rhizophagus clarus*. Mullite ceramic tubes were buried in the soil of each compartment, and soil solution was collected. *A. fistulosum* L. and *Linum usitatissimum* L. inoculated with *R. clarus* were grown in sand culture and *in vitro* monoxenic culture, respectively. Uninoculated *A. fistulosum* L was grown in hydroponic culture to collect root exudate. The soil solution, hyphae extracts, root extract, and root exudates were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis. Shoot P concentration, shoot P content, and shoot dry weight were higher in the inoculated treatment than in the uninoculated treatment. Activity staining of the gel revealed that ACP activity at 187 kDa was observed the soil solution in the inoculation treatment and hyphal extract collected from sand culture and *in vitro* monoxenic culture, but neither in the exudate of non-mycorrhizal roots grown in the hydroponic culture nor in the root extracts irrespective of mycorrhizal status. Those results provide strong evidence that the corresponding activity in the soil solutions in soil culture is of *R. clarus* CK001 origin. These findings suggested that the fungus releases ACP from extraradical hyphae into the hyphosphere.
Key words: arbuscular mycorrhizal fungi, hyphal exudate, soil solution, acid phosphatase, compartment pot
INTRODUCTION

Phosphate fertilizer is produced from phosphate rock. Despite the fact that phosphate fertilizer demand is projected to increase, global phosphate fertilizer production is estimated to peak around 2030, and phosphate rock may be depleted 50-100 years later (Cordell et al. 2009). Two strategies in tackling depletion of phosphate rock are considered. One is to develop plants that can recycle phosphorus (P) more efficiently within their body. The other one is to develop plants that can take up phosphate more efficiently from the soil. Concentration of plant-available phosphate i.e. free inorganic orthophosphate (P_i) in the soil is usually very low in most ecosystems, which frequently limits plant growth. On the other hand, large parts of phosphate in the soil are present as plant-unavailable forms: sparingly soluble inorganic phosphate and organic phosphate. In particular, organic phosphate accounts for 20-80% of total phosphate in the soil (Dalal 1977), and thus conversion (hydrolysis) of organic phosphate into plant-available Pi is a key to sustainable food production.

Plants and fungi possess many genes encoding acid phosphatase (EC 3.1.3.2, ACP) in the genomes, e.g., 29 genes in the plant Arabidopsis thaliana (Wang et al., 2014) and at least 7 genes are expressed in R. clarus (e.g. Kikuchi and Ezawa, 2012, unpublished data). Secretion of ACP into rhizosphere/hyphosphere, however, has been demonstrated only in limited ranges of plant (e.g., Tadano and Sakai 1991; Tatarifar and Claassen 1988) and fungi (e.g., Crowther et al. 2011; Tatarifar et al. 1988; van Aarle and Plassard 2010). In plants, secreted ACP hydrolyzes organic phosphate, increases the Pi pool in the rhizosphere, and thus enhances P_i uptake (Wasaki et al. 2009; Maruyama et al. 2012; Robinson et al. 2012).

AM fungi associate with 80% of land plants and enhance P_i uptake of plants from the
soil through their extraradical hyphae (Smith and Read 2008). It has been considered that the increase in surface area for Pi uptake is the main mechanism underlying the enhanced Pi uptake in the associations. In addition to the mechanism, the involvement of AM fungal ACP in the enhanced Pi uptake has also been proposed. Tarafdar and Marschner (1994) employed the compartment culture system in which the mycorrhizal (i.e. roots + hyphae) and hyphal compartments were separated by a 30-μm nylon mesh and observed higher ACP activity in the hyphal compartment in the presence of an AM fungus *Glomus mosseae*, suggesting that the fungus secreted ACP from extraradical hyphae. Koide and Kabir (2000) demonstrated hydrolysis of organic phosphate in the hyphal compartment of the *in vitro* two-compartment monoxenic culture of *G. intraradices* with a concurrent increase in root P content, suggesting that plants can access to organic phosphate via fungal ACP secretion. Joner and Johansen (2000) found that fungal ACP was mostly associated with the cell wall and released ACP was undetectable, which was also reproduced by Olsson *et al.* (2002).

In plants, not only cell wall-bound ACP but also released ACP plays a significant role in organic P hydrolysis in the rhizosphere. In *Arabidopsis thaliana*, two purple acid phosphatases encoded by *AtPAP12* and *AtPAP26* are released from the roots (Tran *et al.*, 2010), and the corresponding knockout mutants showed poorer growth than the wild type in the presence of organic P (fructose-6-phosphate) as the sole P source in the medium, suggesting that ACP released from the roots greatly contributes to plant growth (Wang *et al.*, 2014). Although these observations led us to hypothesize that AM fungi also release ACP into the soil solution, a technical breakthrough was necessary for the collection and detection of released ACP.

The exudation of organic acid by extraradical hyphae of an AM fungus has been
successfully demonstrated in the soil solution collected using mullite ceramic tubes in a two-compartment culture system (Tawaraya et al., 2006). In the present study, this collection technique was employed for the qualitative assessments of AM fungal ACP exuded in the soil solution to test the hypothesis that AM fungi exude soluble ACP from extraradical hyphae.

MATERIALS AND METHODS

Two-compartment culture for collection of soil solution

Andosol was collected from a native pasture in Haguromachi, Yamagata prefecture, Japan. The soil was air-dried in glass house, sieved on a 2 mm sieve, and steam-sterilized twice at 80°C for 45 min. Chemical properties of the soil were as follows: pH (H₂O), 4.78; Organic carbon, 10.9%; total nitrogen, 0.88%; Truog-P, 5.02 mg P kg soil⁻¹; cation exchange capacity, 43.2 cmol (+) kg soil⁻¹. The soil was fertilized with ammonium sulfate, potassium sulfate, and superphosphate at rates of 1.00 g nitrogen (N), 0.83 g potassium (K), and 0.15 g P kg⁻¹ soil, respectively, and pH was adjusted to pH 5.1 with calcium carbonate at a rate of 4.19 g kg⁻¹ soil. The AM fungus Rhizophagus clarus (T.H. Nicolson & N.C. Schenck) C. Walker & A. Schüßler strain CK001 was propagated with Welsh onion (Allium fistulosum L. cv. Motokura), sorghum (Sorghum bicolor (L.) Moench. cv. New sorghum 2 gou), and white clover (Trifolium repens L. cv. California ladino) for 3 months, and the soil that includes spores, extraradical hyphae, and the roots was used as inoculum. The two-compartment pots were prepared as described by Tawaraya et al. (2006). A nylon bag (top 140 mm, bottom 80 mm, height 90 mm) was prepared with a 30 µm nylon net (NY30HD, Sefar Inc., Heiden, Switzerland), though which only fungal hyphae are able to pass, but not
plant roots, filled with soil-inoculum layers as follows: 20 g of the sterilized in the bottom, 10 g of the inoculum as the second layer, 20 g of the sterilized soil as the third layer, 10 g of the inoculum as the fourth layer, and then covered with 40 g of the sterilized soil (mycorrhizal compartment). For the uninoculated treatment, no inoculum layers were incorporated in the compartment. The nylon bag was placed at the center of a 500 mL plastic pot (11.5×9.5 cm), and the remaining space was filled with 300 g of the sterilized soil (hyphal compartment). Each six mullite ceramic tubes (50 × 2.5 mm, 1DH-1525, Sakaguchi E.H. VOC Corp., Tokyo, Japan) connected to Teflon tubes (150 × 1.5 mm) were embedded in the mycorrhizal and hyphal compartments at intervals of 2 and 3 cm in a circular pattern, respectively (Tawaraya et al. 2006). There were five replications of each treatment. Twelve seeds of Welsh onion were sown in the mycorrhizal compartment at 1 cm depth, irrigated with deionized water to maintain the water potential at −0.03 Mpa, and grown in a 16 h photoperiod (150 µmol m⁻² s⁻¹) at 27°C/25°C (light/dark) in a growth chamber. The pots were irrigated with deionized water every other day. Each pot was covered until germination, and plants were thinned to six per pot. A plastic syringe (5 mL plastic syringe, TOP Co., Ltd., Tokyo, Japan) was connected to a Teflon tube, and 1 mL of soil solution was collected into the syringe by pulling the piston several times. The solution was collected from all tubes 40, 45, 50, and 55 days after sowing, stored at -30°C, and pooled before concentration. Shoots and roots were harvested 55 days after sowing. Shoots were weighted after drying at 70°C for 72 h and were used for plant P assessment. Roots were washed with tap water, wiped with a paper towel, weighed, and used for enzyme preparation and assessment of mycorrhizal colonization.
Sand culture for preparation of extraradical hyphae

Welsh onion inoculated with *R. clarus* CK001 was grown on sterilized (121°C, 45 min.) sea sand in a glass house. A nutrient solution (40 mg N L\(^{-1}\) (NH\(_4\)NO\(_3\)), 20 mg N L\(^{-1}\) (NaNO\(_3\)), 60 mg K L\(^{-1}\) (K\(_2\)SO\(_4\)), 80 mg Ca L\(^{-1}\) (CaCl\(_2\)), 40 mg Mg L\(^{-1}\) (MgSO\(_4\)), 2 mg Fe L\(^{-1}\) (FeSO\(_4\)), 1 mg Mn L\(^{-1}\) (MnSO\(_4\)), 0.01 mg Cu L\(^{-1}\) (CuSO\(_4\)), 0.005 Mo mg L\(^{-1}\) [(NH\(_4\))\(_6\)Mo\(_7\)O\(_{24}\)], 0.4 mg B L\(^{-1}\) (H\(_3\)BO\(_3\)), 0.2 mg Zn L\(^{-1}\) (ZnCl\(_2\)), and 1 mg P L\(^{-1}\) (NaH\(_2\)PO\(_4\)) (Wagatsuma *et al.* 1988) was applied to plants every other day. Extraradical hyphae were collected 45 days after sowing by wet sieving, wiped with a paper towel, weighed, stored at -30°C, and used for enzyme extraction.

In vitro monoxenic culture for preparation of extraradical hyphae

*Agrobacterium rhizogenes*-transformed roots (hairy roots) of flax (*Linum usitatissimum* L.) was purchased from Glomeromycota *in vitro* collection (http://www.mycorrhiza.be/ginco-bel/) were inoculated with surface-sterilized *R. clarus* CK001 spores and grown on modified Strullu–Romand (MSR) medium (Declerck *et al.* 1998) for 60 days in 9 cm Petri dishes according to Chabot *et al.* (1992). After removing the roots with tweezers, the culture medium of 30 Petri dishes was transferred to 1 L of 10 mM sodium citrate (pH 6.0) to solubilize the Gellan Gum medium (Doner and Becard 1991). The extraradical hyphae were collected on a 30 µm nylon net, washed with sterilized water, wiped with a paper towel, weighed, and used for enzyme extraction.

Hydroponic culture for collection of root exudate

Seeds of Welsh onion were sown on moist vermiculite spread. One week after sowing,
30 seedlings were carefully lifted out of the vermiculite, fixed with a sponge in a paper cup filled with 200 mL of the nutrient solution without P, and grown in the growth chamber (light 16 h / dark 8 h, 150 µmol m⁻² s⁻¹, 27°C (light)/ 25°C (dark)). The pH of the nutrient solution was adjusted daily to 5.0 ± 0.05 with 0.5 M H₂SO₄ and 0.5 M NaOH. The nutrient solution was aerated with an air pump and replaced weekly. Thirteen days after transplanting, the seedlings were transferred to 50 ml of sterile deionized water and incubated for 12 h under the same conditions to collect root exudate, and then the solution was stored at –30°C and used for qualitative analysis.

**Enzyme preparation**

Soil solution and root exudates were first concentrated over 200-fold using Amicon® Ultra-15 centrifugal filter units (Millipore, MA, U.S.A.), further to 1800- and 500-fold, respectively, using Microcon® YM-30 centrifugal filter units (Millipore Corporation, MA, U.S.A.), and subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE).

Roots and extraradical hyphae were ground using mortar and pestle at 0°C with an equal volume (wt/vol) of 100 mM borate buffer (pH 8.8) with 10 µL mL⁻¹ protease inhibitor (Plant ProteaseArrest™, Takara Bio Inc., Shiga, Japan). The resulting slurry was transferred to a 1.5 mL tube and centrifuged at 15,000 × g for 30 min at 4°C. The supernatant of root extract was transferred to a new tube and subjected to SDS-PAGE. The supernatant of hyphal extract was concentrated 24 fold using Microcon® YM-30 centrifugal filter units and subjected to SDS-PAGE.

**Qualitative analysis of acid phosphatase**
SDS-PAGE was performed as described by Ezawa and Yoshida (1994). Ten microliter each of the soil solution, root exudate, hyphal extract and root extract were loaded on a 10% SDS-polyacrylamide gel without denaturing and electrophoresed. The gel was gently shaken in washing buffer [50 mL of 100 mM acetate/sodium buffer (pH 5.0) containing 0.5 mL of Triton-X] for 30 min, and ACP activity on the gel was visualized by the azo dye method (Scandalios 1969). The molecular weight of protein was estimated based on mobility relative to those of the protein standards (Takara bio inc. Shiga, Japan) on the gel.

Assessment of plant P and mycorrhizal colonization

Ground shoots were digested in a HNO₃-HClO₄-H₂SO₄ (5:2:1) solution. The P content in the digested solution was determined colorimetrically by vanadomolybdate-yellow assay (Olsen and Sommers, 1982).

Percent AM colonization was determined using the gridline intersect method (Giovannetti and Mosse 1980) after root staining with 500 mg L⁻¹ aniline blue solution (Tawaraya et al. 1998). One hundred root segments from each replicate bag were observed under a microscope (magnification, × 100).

ANOVA with Tukey-HSD test (p < 0.05) was performed with Kaleida Graph 4.0j (HULINKS Inc., Tokyo, Japan).

RESULTS

Qualitative characterization of acid phosphatase released from mycorrhiza

AM colonization was observed in the inoculated treatment 55 days after sowing, but not in the uninoculated treatment. Shoot P concentration, P content, and dry weight
were higher in the inoculated treatment than in the uninoculated treatment (Table 1).

Activity staining of the gel revealed that ACP activity at 187 kDa was observed exclusively in the presence of the fungus in the solutions obtained from the mycorrhizal and hyphal compartments in the soil culture (Fig. 1a). This activity was also detected in the extracts of extraradical hyphae grown both in the sand culture and the in vitro monoxenic culture, but neither in the exudate of non-mycorrhizal roots grown in the hydroponic culture nor in the root extracts irrespective of mycorrhizal status (Figs. 1b and c). ACP activity at 41 kDa was observed in all samples irrespective of mycorrhizal status, except for the root extracts.

**DISCUSSION**

**Release of ACP from extraradical hyphae of AM fungus R. clarus**

The detection of the ACP activity of 187 kDa in the in vitro culture provides strong evidence that the corresponding activity in the soil solutions in soil sand culture is of R. clarus CK001 origin, suggesting that the fungus releases ACP from extraradical hyphae into the hyphosphere. This finding is further supported by the absence of the activity in the root exudate and extract. The detection of the ACP activity in the in vitro culture also excludes the possibility that the activity was of other microorganisms present in the rhizosphere/hyphosphere.

We demonstrated for the first time using mullite ceramic tubes in conjunction with SDS-PAGE that AM fungi release ACP from the extraradical hyphae to the hyphosphere. Our results has been suggested that a large part of extracellular ACP is bound to the cell wall in AM fungi, because released ACP fraction collected from the culture medium i.e. polysaccharide gel in the in vitro culture (Koide and Kabir, 2000;
Olsson et al., 2002) or that collected by 1 h incubation of hyphae in a buffer (Joner and Johansen, 2000) showed little activity. The successful detection of released ACP activity in the present study was likely to be achieved by the concentration of the activity up to three orders of magnitude by ultrafiltration.

Although, fungi have several ACP orthologus on genome (for example, at least 7 genes are expressed in R. clarus, e.g. Kikuchi and Ezawa, 2012, unpublished data), molecular characterization has been rarely reported. Additionally, mobility on the PAGE analysis is modified by glycosylation (Weber and Pitt 1997). Thus, estimation of gene is impossible. The identification of the ACP of 187 kDa is necessary using immunological approach and genetic information of known ACPs.

Although ACP of 41 kDa was detected in all soil solution irrespective of mycorrhizal status, it seems unlikely that the activity is solely of the host plant Welsh onion. This is because the activity was detected not only in the hyphal extract in the sand culture but also that in the in vitro culture, in which different host plants, onion for the sand culture and flax for the in vitro culture, were used to grow the fungus, raising two possibilities; the two plants exude ACPs of similar molecular weight that migrate at similar rates on the gel, or onion and the fungus possess ACPs of similar molecular weight, which were in either case indistinguishable in electrophoresis. The ACP of 41 kDa was detected in root exudates, but not root extracts. The reason of incompatible result might be different mobility caused by formation of dimer (Ozawa et al., 1995) or glycosylation (Weber and Pitt 1997). Further biochemical characterization and identification of genes encoding these ACPs would be necessary to test the possibilities.

Our results indicate that the AM fungus R. clarus CK001 releases ACP from its
extraradical hyphae. It has been considered that increase in surface area for $P_i$ uptake by extraradical hyphae is the main mechanism underlying the enhanced $P_i$ uptake in the mycorrhizal plants. The present study can propose a new mechanism that extraradical hyphae of AM fungi hydrolyze soil organic phosphate and improve $P$ acquisition. The hydrolysis of soil organic phosphate with released ACP and uptake of the resultant $P_i$ by extraradical hyphae remain to be clarified.

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Table 1 AM colonization, shoot P concentration, shoot P content, and shoot dry weight of Welsh onion without (-) or with (+) AM fungus *R. clarus* CK001 inoculation 55 days after sowing. S.E. indicates standard error. Values followed by a different letter are significantly different (*P*<0.05).

<table>
<thead>
<tr>
<th>AMF</th>
<th>AM colonization ± S.E. (%)</th>
<th>Shoot P concentration ± S.E. (mg P g⁻¹)</th>
<th>Shoot P content ± S.E. (mg P pot⁻¹)</th>
<th>Shoot dry weight ± S.E. (mg pot⁻¹)</th>
</tr>
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<tbody>
<tr>
<td>-</td>
<td>0 ± 0 b</td>
<td>0.825 ± 0.086 b</td>
<td>0.0473 ± 0.0044 b</td>
<td>58.0 ± 3.7 b</td>
</tr>
<tr>
<td>+</td>
<td>87 ± 2 a</td>
<td>2.422 ± 0.261 a</td>
<td>0.7686 ± 0.0320 a</td>
<td>325.2 ± 24.2 a</td>
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Fig. 1 SDS-PAGE analysis of ACP activity of 1800-fold concentrated soil solution collected from the mycorrhizal compartment (MC) and hyphal compartment (HC) of Welsh onion inoculated with (+M) or without (-M) *R. clarus* CK001 (a), 24-fold concentrated extraradical hyphal extract from *R. clarus* CK001 collected from sand culture (Hyphal ext. sand) or *in vitro* monoxenic culture (Hyphal ext. *in vitro*), 500-fold concentrated root exudates (Root exud.) from Welsh onion grown on P-free nutrient solution (b), and root extract from Welsh onion inoculated with (+M) or without (-M) AM fungus *R. clarus* CK001 (c). Black and white arrows indicate fungal ACP activity.