Identification of enzymes responsible for the reduction of geraniol to citronellol

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Abstract: The reduction of geraniol to citronellol is the first step for the synthesis of natural phytol in the production of tocopherols and natural vitamin K. Baker’s yeast (Saccharomyces cerevisiae) was used in the bioreduction described above as a whole-cell biocatalyst. However, the enzyme responsible for the reduction of geraniol to citronellol is not yet known. Four old yellow enzyme (OYE) genes were cloned from yeast and plants, and expressed in Escherichia coli for a high level of recombinant proteins. The recombinant protein displayed a catalytic activity of converting geraniol to citronellol as a sole product verified by GC-MS analyses. The recombinant OYE2 intact cells were found to show 3.7 and 1.9-fold higher activity than that of yeast cells and the recombinant crude extracts, respectively. Compared to the recombinant fusion enzyme, the entrokinase-cleaved enzyme displayed nearly identical activity for geraniol reduction. To our knowledge, this is the first enzyme identified to catalyze the formation of citronellol from geraniol by reducing the allylic alcohol double bond, which is normally known as inactivating group for the old yellow enzymes.

Keywords: allylic alcohol, citronellol, geraniol, old yellow enzyme, reduction

Introduction

The reduction of geraniol to citronellol is the first step for the synthesis of natural phytol in the production of tocopherols and natural vitamin K. Baker’s yeast (Saccharomyces cerevisiae) was used in the bioreduction described above as a whole-cell biocatalyst. 1,2 However, the enzyme responsible for the reduction of geraniol to citronellol is not yet known.3 Geraniol is considered to be a “non-activated” alkene, in contrast to the activated alkenes bearing an electron-withdrawing group, such as ketone, aldehyde, carboxylic acid, ester, anhydride, lactone, imide, nitro. Recently, enzymes from the old yellow enzyme (OYE) family have been discovered to catalyze the asymmetric reduction of activated alkenes, creating up to two chiral centers and, thus, showing a great potential in the industrial reduction of many commercially useful substrates. 3–9 The OYE s are characterized by high stereospecificity, strict regioselectivity, and rather broad substrate specificity. Fungi, particularly various yeasts, and bacteria were prominently used in whole-cell biotransformations of activated alkenes; whereas plant cells have only a minor part in the field. However, whole-cell biotransformations frequently resulted in poor product yields due to undesired side reactions such as carbonyl group reduction, ester hydrolysis, acyloan reactions, and acetyl cleavage, in addition to substrate competition and limited enzyme amount in the original cells. To avoid these problems, isolated and/or cloned enzymes have gained growing interest in biocatalytic applications. Successful attempts in the chemo- and stereoselective reduction of activated alkenes were reported by using cloned yeast OYE1 from Saccharomyces carlsbergensis, 3,9 OYE2 and OYE3 from S. cerevisiae, 8–10 CmOYE from Candida macedoniensis; 9 and bacterial OYE homologues YqjM from Bacillus subtilis, NCR from Zymomonas mobilis; 8,10 and plant OYE homologues LeOPR1 and LeOPR3 from Lycopersicon esculentum (tomato). 10,11 The plant 12-oxophytodienoate reductase (OPR) is involved in the reduction of (9S,13S)-12-oxophytodienoate to the corresponding cyclo-pentenoic acid OPC-8:0 for jasmonate biosynthesis. 12 By using the cloned OPR1 and OPR3 from tomato, a striking change of stereoselectivity and a remarkably broad substrate spectrum were achieved for asymmetric reduction. 10,11 Although the above OYE enzymes have been biochemically well characterized, they have never been investigated for the capacity of reducing the allylic alcohol double bond in geraniol, because alcohols are not typically favored as activating groups for OYEs. 3 In our effort to broaden the applicability of plant OPR, we came across its catalytic activity of converting geraniol to citronellol as a sole product (Scheme).

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Results and Discussion

Two novel OPR homologues designated as EaOPR and leOPR, together with the known HbOPR from rubber tree and OYE2 from S. cerevisiae were cloned and functionally expressed by pET32α+ in E. coli BL21 (DE3). The complete cDNAs for EaOPR [GenBank: JN828575] and leOPR [GenBank: JN828576] were recovered from the perennial herbs Epimedium acuminatum (Berberidaceae) and Isodon eriocalyx (Lamiaceae) through degenerate primers and subsequent rapid amplification of cDNA ends (RACE). Based on SDS-PAGE analyses, higher level of soluble protein expression was obtained with 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) at 20°C for 23 h (Figure S1 in Supplementary Material). Crude extracts of the induced cells were assayed for the C=C bond reductase activity using methyl crotonate, methyl trans-cinnamate, 3-methyl-2-cyclohexanone, and geraniol as a substrate. Significant enzymatic production of the predicted product was observed when geraniol was incubated in the presence of each crude enzyme; and citronellol was detected as a sole product by GC-MS with authentic compound. Incubating with 3-methyl-2-cyclohexanone, only the recombinant HbOPR and OYE2 led to formation of 3-methyl-2-cyclohexanone as detected by GC-MS (Figure S2 in Supplementary Material). This activity was previously observed for the recombinant OYE2. All the recombinant enzymes were inactive to methyl crotonate and methyl trans-cinnamate based on GC-MS analyses (data not shown). The purified recombinant HbOPR was used for further characterization of enzyme properties. After purification under native condition on Ni-NTA His•Bind Superflow, the analysis of the elute on SDS-PAGE led to the detection of one band corresponding exactly to the predicted size of the recombinant protein of approximately 60 kDa (Figure 1A). When incubated with geraniol and cofactor NADH or NADPH, the purified HbOPR made citronellol as a primary product. The product peak at 9.31 min (retention time) generated the dominant mass spectra (9.31 min of retention time) matching citronellol (I) and geraniol (2), and the reaction product formed by the recombinant HbOPR and the inactive enzyme as control, with geraniol as substrate. GC profiles (B) and the corresponding mass spectra (C) illustrate the product peak (9.31 min of retention time) matching citronellol (1) with respect to retention time and mass fragmentation pattern.

Original pET32α+ displayed a very low activity as background (Table S2 and Figure S3 in Supplementary Material). In addition, the results of recombinant OYE2 were compared with those obtained by employing baker’s yeast intact cells. For the estimation we used equal amount of S. cerevisiae yeast cells, the recombinant OYE2 intact cells, and the recombinant OYE2 crude extracts from the same amount of OYE2 cells. Results demonstrated that the recombinant OYE2 intact cells were best performer, showing 3.7 and 1.9-fold higher activity than that of yeast cells and crude extracts, respectively (Figure 2C, also see Table S2 and Figure S4 in Supplementary Material). Similar results were observed when the reaction mixtures were incubated at 28°C (Table S2).

To further clarify the role of the fusion tag in the recombinant proteins, enzymatic cleavage was carried out with entrokinase upon the recombinant HbOPR fusion protein. A complete cleavage was observed for the protein incubating with 40 IU of entrokinase at RT for 2 h, yielding the fusion tag and the ‘net’ HbOPR of approximately 44 kDa (Figure 3A). Compared to the fusion HbOPR, the ‘net’ HbOPR enzyme possessed nearly identical activity for geraniol reduction (columns a and b, Figure 3B), reducing geraniol to citronellol as a primary product. By subtracting the relative activity of the...
induced BL21 (DE3) cell lysate from that of the induced original pET32a+/BL21 (DE3) cell lysate, we can estimate a ‘net’ fusion tag contribution to the reduction activity. By the same way we’ll know if the entrokinase is active to geraniol. As shown in Figure 3B (columns c–e), neither fusion tag nor entrokinase was capable of forming citronellol from geraniol (Table S2 in Supplementary Material).

Conclusions

In summary, we have first identified the enzyme responsible for the reduction of geraniol to citronellol. The recombinant HbOPR from rubber tree and the recombinant OYE2 from baker’s yeast were found to be best performer and can be further prospected for their industrial biocatalysis applications. Despite of high protein level achieved by fusion expression of pET32a+, other expression vectors and *E. coli* strains may also be tried to harvest more enzymes. With the strict regioselectivity, the identified geraniol reductase should be further investigated for its stereospecificity and substrate specificity. Finally, we suggest the recombinant whole cells as biocatalysts in an economical way, considering the universal existence of NAD(P)H in living cells, the predominance of the enzymatic product citronellol, and almost no undesired side products.

Experimental Section

Gene Cloning. Young leaves of *Epimedium acuminatum* and *Isodon eriocalyx* were collected from Kunming Botanic Garden, Chinese Academy of Sciences. Young shoots of *Hevea brasiliensis* were collected from Hekou County of southeast Yunnan province. The full length cDNAs of *EaOPR* and *IeOPR* were obtained by a homology-based cloning strategy using Superscript™ III First-strand Synthesis System and GeneRacer™ Kit (Invitrogen), and deposited in the GenBank/EMBL database under accession numbers JN828575 (for *EaOPR*) and JN828576 (for *IeOPR*), respectively. The open reading frame (ORF) of *HbOPR* was cloned by RT-PCR according to the sequence [GenBank: DQ004685.1]; and the ORF of *OYE2* [GenBank: NM_001179310.1] was amplified from the genomic DNA of *Saccharomyces cerevisiae*. All primers were listed in Table S1 in Supplementary Material.

Expression in *E. coli*. Each ORF of the four OYE genes was cloned into the expression vector pET32a+ (Novagen) which were subsequently transformed into *E. coli* BL21 (DE3) for a fusion expression, using the original pET32a+ as nega-
Enzyme Assays and GC-MS Analyses. Crude extracts of the induced BL21 (DE3) cells were assayed for enzyme activity using 1 mM of the corresponding substrate geraniol, methyl crotonate, methyl trans-cinnamate, and 3-methyl-2-cyclohexenone (all purchased from Sigma), incubating for 8 h at 37°C. No additional enzyme cofactor NADH or NADPH was needed since the crude extracts contain both. After extraction with ethyl acetate, the upper phase was collected by sonication on ice and then centrifuged at 12,000 rpm for 10 min at 4°C and the supernatant, containing the completely cleaved recombinant HbOPR enzyme was used for crude enzyme assay or purification. His-tagged proteins were purified to near homogeneity according to the Ni-NTA His•Bind Superflow protocol (Novagen) with a wash buffer (20 mM Tris-HCl pH 7.5, 0.5 M NaCl, 5 mM imidazol). The cell lysate obtained by sonication on ice was then centrifuged at 12,000 rpm for 10 min at 4°C and the supernatant, containing the completely cleaved (2 h) protein was assayed for enzyme activity as described above. Meanwhile, crude extracts from the same amount of induced OYE2/pET32a+/BL21 (DE3) cells harboring the original PET32a+ as controls. For estimation of whole-cell transformation in comparison to crude enzyme activity, we used an equal amount of S. cerevisiae yeast cells, the induced OYE2/pET32a+/E. coli BL21 (DE3) intact cells, and the crude extracts from the same amount of induced OYE2/pET32a+/E. coli BL21 (DE3) cells. For fusion tag cleavage, the induced HbOPR/pET32a+/BL21 (DE3) cells were cleaved with 40 IU Entrokinase (Shanghai, Sangon) at RT (24–27°C) for 0.5, 1, 1.5, 2, 5 h, and subsequently analyzed by SDS-PAGE. The completely cleaved (2 h) protein was assayed for enzyme activity as described above. Meanwhile, crude extracts (kept for 2 h before enzyme assay) from the induced HbOPR cells and the induced cells harboring the original pET32a+, together with the induced BL21 (DE3) cell lysate prepared with or without entrokinase were analyzed in parallel to serve as controls. For cofactor determination, the purified recombinant HbOPR enzyme was incubated at 37°C for 4 h in the reaction mixture containing 1 mM geraniol and 3 mM NADH or NADPH.

GC-MS [Agilent HP6890/5973, column: 0.25 mm × 30 m, 0.25 µm (HP-5MS)] was conducted under electron-impact (EI) mode (70 eV). The flow rate of helium carrier gas was set at 1.0 ml/min. Samples (2 µL) were injected at 60°C. After holding the samples for 5 min at 60°C, the column temperature was increased at 10°C/min to 260°C and hold for 10 min. The MS date was collected from 35 to 450 m/z.

Electronic Supplementary Material
Supplementary material is available in the online version of this article at http://dx.doi.org/10.1007/s13659-011-0032-6 and is accessible for authorized users.

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References
[1] Gramatica, P.; Manitto, P.; Ranzi, B. M.; Delbianco, A.; Francavilla, M. Experientia 1982, 38, 775–776.
[2] Gramatica, P.; Manitto, P.; Monti, D.; Speranza, G. Tetrahedron 1987, 43, 4481–4486.
[3] Toogood, H. S.; Gardiner, J. M.; Scrutton, N. S. ChemCatChem 2010, 2, 892–914.
[4] Steuerman, R.; Hauer, B.; Hall, M.; Faber, K.Curr. Opin. Chem. Biol. 2007, 11, 203–213.
[5] Swiderska, M. A.; Stewart, J. J. Mol. Catal. B Enzym. 2006, 42, 52–54.
[6] Hall, M.; Stueckler, C.; Hauer, B.; Stuermer, R.; Friedrich, T.; Breuer, M.; Kroull, W.; Faber, K. Eur. J. Org. Chem. 2008, 9, 1511–1516.
[7] Brenna, E.; Fronza, G.; Fugganti, C.; Monti, D.; Parmeggiani, F. J. Mol. Catal. B Enzym. 2011, 73, 17–21.
[8] Wada, M.; Yoshizumi, A.; Noda, Y.; Katoaka, M.; Shimizu, S.; Takagi, H.; Nakamori, S. Appl. Environ. Microbiol. 2003, 69, 933–937.
[9] Kataoka, M.; Kotaka, A.; Thiwthong, R.; Wada, M.; Nakamori, S.; Shimizu, S. J. Biotechnol. 2004, 114, 1–9.
[10] Hall, M.; Stueckler, C.; Ehamner, H.; Pointner, E.; Oberdorfer, G.; Kruber, H.; Hauer, B.; Stuermer, R.; Kroull, W.; Machereoux, P.; Faber, K. Adv. Synth. Catal. 2008, 350, 411–418.
[11] Hall, M.; Stueckler, C.; Kroull, W.; Machereoux, P.; Faber, K. Angew. Chem. Int. Ed. 2007, 46, 3934–3937.
[12] Schaller, F.; Biesgen, C.; Müssig, C.; Altmann, T.; Weiler, E. W. Planta 2000, 210, 979–984.
[13] Schlieben, N. H.; Niefind, K.; Müller, J.; Riebel, B.; Hummel, W.; Schomburg, D. J. Mol. Biol. 2005, 349, 801–813.