A pirin-like protein from *Pseudomonas stutzeri* and its quercetinase activity

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**Abstract**

A pirin-like protein from a marine denitrifying bacterium, *Pseudomonas stutzeri* Zobell has been heterologously expressed in E. coli and purified to homogeneity with metal-affinity and gel filtration chromatographies. The recombinant pirin-like protein has exhibited quercetinase activities upon the incorporation of a divalent metal ion, while its biological role remains unclear. In the case of Cu^{2+} the holo-protein demonstrated the highest activities and spectroscopic properties typical of type II Cu protein. A 3D-structural model constructed using the crystal structure of human pirin as template indicated that the metal binding site is constructed with 3His1Glu located in the consensus sequences in the N-terminal domain.

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1. Introduction

Flavonoids are one class of secondary metabolites of plants. Dietary plant flavonoids have been proposed to contribute to cancer prevention, neuro-protection, and cardiovascular health though their anti-oxidant, anti-inflammatory, pro-apoptotic, and anti-proliferative activities [1,2]. Terrestrial plants secrete flavonoids such as flavone, flavonol, flavanone, flavonol, flavan and isoflavon from their roots to send signals towards symbiotic bacteria or to avoid attacks from other bacteria [3,4]. Recent studies have shown that marine plants also utilize flavonoids for protection [5,6].

Quercetin (3,5,7,3′,4′-pentahydroxy-flavone) is produced in germinating seedlings by UV or blue light, and is widely contained in plants as glycosides, rutin and quercitrin [7,8]. Quercetin is a polyphenolic compound together with anthocyanin and catechin etc. These polyphenolic compounds have received special attentions for their anti-inflammatory actions arising from scavenging capacity of free radicals [9,10].

Quercetin is converted into the corresponding depside (phenolic ester 2-protocatechuylphloroglucinol carboxylic acid) and carbon monoxide by quercetin 2,3-dioxygenase (2,3QD) or quercetinase (Scheme 1) [11]. Transformations of flavonols such as quercetin can be divided into three or four processes: microbial bioconversion, anaerobic or aerobic prokaryotic catabolism, and aerobic eukaryotic microbial catabolism [12]. Quercetinase has been found in various molds [13–17] and some bacteria such as *Bacillus subtilis* [18]. Beside catabolic roles ascribable to eukaryotic quercetinase, detoxification role of this enzyme against deleterious effects has also been postulated inside bacteria.

Amino acid sequence and higher order structure of quercetinases indicated that this enzyme has a strong resemblance with pirin [12,19], which concerns in apoptosis and cellular stress in eukaryotic organisms and in seed germination and transcription of light- and ABA (abscisic acid)-regulated gene in plants. Both quercetinase and pirin belong to cupin superfamily which has one (monocupin) or two (bicupin) domains comprised of characteristic β-stranded motifs and intervening loops with the consensus sequences containing His and Glu residues [11]. It has been clarified that metal ions such as Cu^{2+}, Fe^{2+}, and Ni^{2+} are required to exert quercetinase activities, whilst it is not necessarily clear whether pirins play a role as sensor for metal ions [20]. While pirins and quercetinases have strong structural resemblances, their requirements for metal ions are different. This fact urged us to study quercetinase activity of pirin, although the bindings of divalent metal ions to pirins have already been reported [21].

Metabolisms of flavonoids, especially of quercetin, by microorganisms in terrestrial plant rhizosphere, have been studied in some detail [12,22]. However, few studies have been performed on defense of microorganisms from flavonoids secreted by aquatic plants. We have selected *Pseudomonas stutzeri* Zobell, formerly *Pseudomonas perfrectomarina* (CCUG 16156=ATCC 14405) for the study on decomposition of flavonoids. *P. stutzeri* is a Gram-negative bacterium adapted for a variety of circumstances, and biodegradation...
2. Methods

2.1. Chemicals

Quercetin, fisetin, kaemferol, and myricetin were purchased from Wako chemicals (Japan). Galangin and taxifolin were obtained from Sigma, morin from TCI (Japan), and luteolin from LKT laboratories (U.S.A.). Flavonoid stock solutions were prepared in dimethyl sulfoxide. All other chemicals were of analytical grade.

2.2. Construction of the expression system of the pirin-like protein in E. coli

The 5′-UTR region containing Shine-Dalgalno sequence and the open reading frame coding for the pirin-like protein with 6xHis-tag at C-terminus was amplified by PCR using two primers (P1: 5′-ggaattcggagcgaagatggcccaacgggaaattc-3′, containing the EcoRI recognition site (underline), Shine-Dalgalno sequence (bold letters), and coding sequence for the N-terminal region of the pirin-like protein; P2: 5′-cgagcgaagatggcccaacgggaaattc-3′, containing the BamHI site (underline) and the C-terminal region of the protein with 6xHis-tag (italic letters)) and a genomic DNA fragment of P. stutzeri Zobell as template.

The amplified gene fragment was digested with EcoRI and BamHI, and inserted into pUC18 vector to yield an expression plasmid pUC-2,3QD. To avoid co-expression of LacZ-fused pirin-like protein, a stop codon was introduced at an upstream of the SD sequence by PCR mutagenesis using two primers (5′-gaattcggagcgaagatggcccaacgggaaattc-3′, 5′-cttcctcaggaatctggaatctggaagatggcccaacgggaaattc-3′). The result plasmid was designated as pUC-2,3QD*.

2.3. Purification of the recombinant pirin-like protein

E. coli BL21 cells transformed with pUC-2,3QD* were cultured in LB medium supplemented with 50 μg/ml ampicillin at 37°C for overnight with shaking (150 rpm). The recombinant protein was induced with 0.2 mM IPTG. The cells were collected by centrifugation and disrupted by sonication in 20 mM Tris-H2SO4, pH 8.0. After centrifugation, the supernatant of the crude extract was applied onto a His-Accept resin column (Nacalai tesque, Japan) for Ni-affinity chromatography. After the column was washed with the same buffer containing 300 mM NaCl and 20 mM imidazole, proteins were eluted with the buffer containing 300 mM NaCl and 300 mM imidazole. The gel filtration using Superdex 200 (GE healthcare, U.S.A.) has been performed to obtain pure protein.

2.4. Determinations of protein concentration and amino acid sequence

Protein concentration was determined with the BCA protein assay kit (Pierce, U.S.A.) and bovine serum albumin as a standard protein. The N-terminal amino acid sequence of the recombinant protein was analyzed by a Shimadzu PPSQ-33A protein sequencer.

2.5. Construction of the 3D structure model

The 3D model structure for the P. stutzeri pirin-like protein was generated by SWISS-MODEL (http://swissmodel.expasy.org/) using human pirin (PDB ID: 4HLT) as a template.

2.6. Enzyme assays

Quercetinase activity of the pirin-like protein has been determined from the absorption change at 380 nm (ε = 18,700 M⁻¹ cm⁻¹) for the reaction mixture containing 50 μM quercetin, 100 mM NaCl and 5%(V/V) DMSO in 50 mM Tris–HCl buffer at 25°C. One unit is defined as the amount of the enzyme catalyzing the decomposition of 1 μmol of quercetin (flavonol) per min. Production of carbon monoxide as co-product has been ascertained from the reaction with PdCl₂ soaked on filter paper to give black solid precipitates of elemental palladium [17]. Kinetic parameters, Kₘ and Vₘₐₓ values were determined based on triplicate data by using Igor Pro ver. 5.03. The pH dependence study of quercetinase activity has been performed using 50 mM Britton–Robinson buffer. Thermal stability of the holo-protein has been studied by incubations at ambient temperatures for 30 min. In addition to quercetin, enzymatic activities for fisetin, galangin, kaemferol, morin, myricetin, taxifolin, and luteolin are also studied.

2.7. Spectroscopic characterizations of the CU-pirin-like protein

Absorption spectra have been measured on a Shimadzu UV-2600 spectrometer or on a JASCO V-560 spectrometer, to both of which a temperature controller has been attached, using quartz cells with 1 mm or 10 mm path-length. Circular dichroism (CD) spectra have been measured on a JASCO J-720 spectropolarimeter. Electron paramagnetic resonance (EPR) spectrum has been measured on a JEOL JES-RE1X X-band spectrometer at 77 K.

3. Results and discussion

3.1. Amino acid sequence and model structure of the pirin-like protein

We have cloned the gene coding for the pirin-like protein from...
the genome of *P. stutzeri* Zobell (ATCC 14405) by PCR based on the sequence obtained with genome walking prior to the construction of the expression system (data not shown). The sequence of cloned genome fragment is identical to the reported one (GenBank accession number of EH796871).

The amino acid sequence of the pirin-like protein from *P. stutzeri* Zobell (Fig. 1) shares 21–34% identities with the quercetinase sequences from *A. japonicus* (Acc: Q75126), *P. olsonii* (Acc: AV24349), and *B. subtilis* (Acc: P42106), and from *E. coli* (Acc: P46852), human (Homo sapiens, Acc: AA38390), and *P. stutzeri* Zobell. The white letters on the black background are the common conserved residues of quercetinases and pirins. The conserved residues of quercetinases or pirins are shown by the light-gray and the dark-gray shadows, respectively. The coordination residues of metal ions are marked by asterisk and box.

### Domain 1

**A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | DTSSLTVEDADVPRYPVTYIHSNARLYLYCQTVLYTTGVPGSYPFAPLMGFSADAKVPHKLQTVTVAHYNGC-S |

**P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | MIALSAFLILASLGALSLILYLYLSNPAPPRPYVYNAPPVGDQVYRTGPDPSSDNPAFNPPSSSSGHPQHGYKNDTSF-D |

**B. subtilis**

| Domain | Sequence |
|--------|----------|
| 1      | MRTLCNSFLEPIMYLLGGEERLYDGEQDVAMNGSTEGDDILVSLGKGD---APFVCTWGDHSGILWYK-S |

**E. coli**

| Domain | Sequence |
|--------|----------|
| 1      | MLYILKXANLXNHLGWDTNYTPAMYPXVNYLSNPEWYSPHIAAPP | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | FQPLWAGESTCCTTVLSGGSYSPSFTDIIPDTMELPGGFLDLTFYCHNAATDTHYFIESSSEDSSTIETGSDSTTISITSDVYALE |

**P. olsonii**

| Domain | Sequence |
|--------|----------|
| 2      | FQPLWAGESTCCTTVLSGGSYSPSFTDIIPDTMELPGGFLDLTFYCHNAATDTHYFIESSSEDSSTIETGSDSTTISITSDVYALE |

**B. subtilis**

| Domain | Sequence |
|--------|----------|
| 2      | LEYDISGDSCDVDSSSVTVQCDPPLIFPSTPDPFQPVGFVPLSNNNVS---EVQVOHPPA98 | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | VEIQCLUXGNYCUPDNLOLQDFELIGAGTHFQGLPLPFHGLGSXWNSTN | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HP | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

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| 2      | HX | **A. japonicus**

| Domain | Sequence |
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| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 2      | HX | **A. japonicus**

### Domain 2

**A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | GRVVIGIGATATLGS-ΕEPIEGVEYTVLYLSEKSKLTVFSGYSGDLQDN | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | G | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | G | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | G | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | G | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | G | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | G | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | G | **A. japonicus**

| Domain | Sequence |
|--------|----------|
| 1      | G | **P. olsonii**

| Domain | Sequence |
|--------|----------|
| 1      | G | **A. japonicus**

### Requirement of metal ions for quercetinase activities

The pirin-like protein as isolated did not exhibit activities to quercetin and other flavonols, fisetin, galangin, kaempferol, morin, myricetin, taxifolin, and luteolin. Therefore, divalent metal ions were reacted with the apo-proteins. We could observe flavonol dioxygenase activities (quercetinase activities as a narrow sense) of the holo-proteins from the absorption changes for dioxygenase activities (quercetinase activities as a narrow sense) of the holo-proteins.

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decompositions of substrates and also from the formation of CO by
the Pd²⁺ reduction. Relative activities for every substrate were
changed in the following order: Cu²⁺ > Fe³⁺ > Zn²⁺ > Mn²⁺, Co²⁺, Ni²⁺. The specific activity of the Cu²⁺-acted pirin-like protein
against quercetin was 1.2 U/mg. The fact that Zn²⁺ also exhibited
oxygenase activities, although considerably low, indicates that the
central metal ion does not perform a redox change but constructs
the active center to activate flavonoids and O₂. According to the
studies on quercetinase [11,19], Cu²⁺ and Fe³⁺ are most fre-
frequently required divalent metal ions. Recently, other metal ions
such as Ni²⁺, Co²⁺, and Mn²⁺ requiring quercetinases have also
been reported [20]. Different activities shown by divalent metal
ions would be due to differences in their Lewis acid character and
steric requirement. Cu²⁺ ion favors the five- to six-coordinated
structures with one or two elongated axial bonds according to
small molecule studies. Otherwise, the tetrahedrally hindered
four-coordinated structure is favored by Cu²⁺ ion [15,24–26].

To explore how many metal ions are required for quercetinase
activities, titrations for Cu²⁺ and Fe³⁺ have been performed. Figs. 2A
and B unequivocally show that exactly one Cu²⁺ ion (Fig. 2B) is
bound to a pirin-like protein. This experimental fact coincides
with the prediction from the 3D-structure model which shows a single potential metal binding site in the N-terminal do-
main of the pirin-like protein (Supplementary Fig. 1).

### Table 1

| Substrates          | Relative activity (%) | CO formation |
|---------------------|-----------------------|--------------|
| Quercetin           | 100                   | +            |
| Myricetin           | 600                   | +            |
| Fisetin             | 28                    | +            |
| Kaempferol          | 0                     | –            |
| Galangin            | 0                     | –            |
| Morin               | 0                     | –            |
| Taxifolin           | 0                     | –            |
| Luteolin            | 0                     | –            |

*Measurement conditions: protein concentration, 50 nM; 50 μM flavonol in 50 mM Tris–HCl buffer (pH 7.5) supplemented with 0.1 M NaCl and 5% (V/V) DMSO.

### 3.4. Enzymatic activities

The relative oxidation activities of the Cu-pirin-like protein
against quercetin, myricetin, fisetin, kaempferol, galangin, morin,
taxifolin, and luteolin are tabulated in Table 1 together with si-
multaneous qualitative analyzes of CO formation. The present Cu-
pirin-like protein exhibited narrow substrate specificities, the high
oxidation activities to quercetin and myricetin differing from other
quercetinases. The kinetic parameters, Kᵣ and V_max values for
quercetin and myricetin are 13 μM and 1.2 U/mg, respectively,
and 9.4 μM and 5.3 U/mg, respectively. The Kᵣ values are analogous to
the values reported for quercetinases, while the V_max values are
considerably low (45–180 U/mg for fungal quercetinases) [12],
presumably because quercetinase activity is not intrinsic to the
present pirin-like protein. Expecting an increase in activity by
enlarging the entrance of cavity to accommodate the bulky sub-
strates, we performed a mutation at the non-coordinating Phe56
for Ala. However, enzymatic activities were significantly decreased
presumably because the mutant molecule became unstable (data
not shown).

Considerably broad dependencies of activity in the pH range of
3.5–11.5, highest at pH ca. 7.0–7.5, are shown in Fig. 3A. The
optimum pH value of the Cu-pirin-like protein is 1.0–1.5 pH units
higher than those of the fungal quercetinases [12], Fig. 3B indicates
that the present Cu-pirin-like protein exhibits the highest activity
at ca. 40°C but loses activity at 90°C.

### 3.5. Spectroscopic characterizations

Absorption spectrum of the pirin-like protein as isolated is
shown in Fig. 4A as broken line. Only one absorption band is ob-
served at 280 nm (ε = 35,000 M⁻¹ cm⁻¹), indicating the protein
is in the apo-form (vide supra). With the action of excess Cu²⁺ ions
on apo-protein, the absorption bands appeared at ca. 650 nm (not
shown), the wavelength that Cu²⁺ ions in water never give the
absorption band. The prolonged incubation of a stoichiometric
amount of Cu²⁺ ion with the apo-protein molecule also gave the
same spectrum (full line in Fig. 4A). The 650 nm band is assigned
to the d–d band from the Cu²⁺ ion bound to the pirin-like protein.
The intensity (ε = ~100 M⁻¹ cm⁻¹) is in the range reported for
Cu-proteins containing a type II Cu (ε < ~500 M⁻¹ cm⁻¹) [28].
Absorption spectra of other Cu-quercetinases have not been pub-
lished yet, and accordingly, comparison is not possible.

The CD spectrum of the Cu-pirin-like protein is shown in
Supplementary Fig. 2 as full solid line, indicating that protein
conformation did not change practically after the incorporation of Cu\(^{2+}\) ion into protein molecule. The CD spectrum in the visible region afforded some CD bands at around 650 nm (shown elsewhere after ascertaining reproducibility).

Fig. 4B shows the EPR spectrum of the holo-protein. It appears that a single tetragonal Cu\(^{2+}\) species is present with the spin Hamiltonian parameters of \(g_\parallel = 2.26\), \(g_\perp = 2.06\), \(A_\parallel = 167\) mT (17.6 \(\times\) 10\(^{-3}\) cm\(^{-1}\)) typical for type II Cu with the binding of 2N2O to 4 N atoms [29]. The presence or absence of water molecules or a hydroxide ion in an equatorial or the axial coordination position(s) is unclear from the EPR parameters. Nevertheless, the EPR parameters and the correlation of the \(g_\parallel\) vs. \(A_\parallel\) values are indicative for the binding of the conserved His59, His61, His103 and Glu105 to Cu\(^{2+}\) ion. Type II Cu EPR signals have been reported for the quercetinase from \(A.\) japonicus: \(g_\parallel = 2.33\) and \(A_\parallel = 15 \times 10^{-3}\) cm\(^{-1}\) in MES buffer, pH 6.0 (minor species: \(g_\parallel = 2.29\) and \(A_\parallel = 13 \times 10^{-3}\) cm\(^{-1}\)) (Major and minor species have been considered to have the six-coordinated and distorted five-coordinated structures, respectively) [30]. In the present study, the Tris-HCl buffer, pH 8 was used to ensure the bindings of ligand groups to Cu\(^{2+}\). Therefore, the Cu\(^{2+}\)-EPR signal with a highly planar character might have been obtained due to the binding of a hydroxide ion in the place of a water molecule, and minor species is practically negligible.

4. Conclusions

We constructed an expression system of a pirin-like protein from \(P.\) stutzeri Zobell in \(E.\) coli. The recombinant protein was expressed as the mixture of the target pirin-like protein and LacZ fusion at its N-terminus, but the pure recombinant protein molecules could be obtained by introducing the stop codon at the upstream region of the SD sequence. One divalent metal ion was incorporated into an apo-protein molecule and exhibited quercetinase activities, highest for Cu\(^{2+}\) ion. Furthermore, we performed detailed kinetic analysis for the pirin derived from a microbe for the first time and revealed that the present pirin-like protein has unique specificities considerably different from quercetinases reported hitherto. Absorption, CD, and EPR spectra of the Cu protein indicated that the coordination of all or some of His59, His61, His103 and Glu105 located in the consensus sequences for cupin

**Fig. 3.** (A) \(pH\) dependence and (B) optimum temperature of the quercetinase activity of the Cu-pirin-like protein from \(P.\) stutzeri Zobell. Quercetinase activity was determined at different \(pH\) with 50 mM Britton-Robinson buffer (Fig. 3A), and 50 mM Tris-HCl buffer, pH 7.5 (Fig. 3B).

**Fig. 4.** Absorption (A) and EPR (B) spectra of the Cu-pirin-like protein from \(P.\) stutzeri Zobell. Absorption spectrum of the apo protein is shown with broken line in Fig. 4A. Measurement conditions: protein concentration, 21 \(\mu\)M, in 50 mM Tris-HCl buffer (pH 7.5). The absorption spectrum was measured at room temperature using a 10 mm path length quartz cell and the EPR spectrum measured at 77 K with 9.2 GHz microwave.
superfamily. The biological role of the present pirin-like protein still remains unclear, but the fact that a divalent metal ion such as Cu²⁺ is required for quercetinase activities will be closely concerned with the involvement of this protein in a transcription process for regulation. In the next stage we will focus our study to explore this uninsured evolutional question, especially higher structural homology of the present pirin-like protein with the human pirin.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.bbrep.2015.08.001.

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