Title	STRUCTURAL PROPERTIES OF PHYCOERYTHRIN FROM DULSE PALMARIA PALMATA
Author(s)	Miyabe, Yoshikatsu; Furuta, Tomoe; Takeda, Tomoyuki; Kanno, Gaku; Shimizu, Takeshi; Tanaka, Yoshikazu; Gai, Zuoqi; Yasui, Hajime; Kishimura, Hideki
Citation	Journal of food biochemistry, 41(1), UNSP e12301 https://doi.org/10.1111/jfbc.12301
Issue Date	2017-02
Doc URL	http://hdl.handle.net/2115/68247
Rights	This is the peer reviewed version of the following article: UNSP e12301-Journal of food biochemistry, 2017-02, 41(1) UNSP e12301-, which has been published in final form at DOI: 10.1111/jfbc.12301. This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Self-Archiving.
Туре	article (author version)
File Information	kishimura.pdf



### 1 STRUCTURAL PROPERTIES OF PHYCOERYTHRIN

## **2 FROM DULSE PALMARIA PALMATA**

3

- 4 YOSHIKATSU MIYABE <sup>1</sup>, TOMOE FURUTA <sup>1</sup>, TOMOYUKI TAKEDA <sup>1</sup>, GAKU
- 5 KANNO <sup>1</sup>, TAKESHI SHIMIZU <sup>2</sup>, YOSHIKAZU TANAKA <sup>3, 4</sup>, ZUOQI GAI <sup>3</sup>,
- 6 HAJIME YASUI 5 and HIDEKI KISHIMURA 6,7

7

- 8 <sup>1</sup> Chair of Marine Chemical Resource Development, Graduate School of Fisheries Sciences,
- 9 Hokkaido University, Hakodate, Hokkaido 041-8611, Japan
- 10 <sup>2</sup> Department of Research and Development, Hokkaido Industrial Technology Center,
- Hakodate, Hokkaido 041-0801, Japan
- 12 <sup>3</sup> Laboratory of X-Ray Structural Biology, Faculty of Advanced Life Science, Hokkaido
- 13 University, Sapporo 060-0810, Japan
- <sup>4</sup> Japan Science and Technology Agency, PRESTO, Sapporo 060-0810, Japan
- 15 Laboratory of Humans and the Ocean, Faculty of Fisheries Sciences, Hokkaido University,
- 16 Hakodate, Hokkaido 041-8611, Japan
- 17 Laboratory of Marine Chemical Resource Development, Faculty of Fisheries Sciences,
- 18 Hokkaido University, Hakodate, Hokkaido 041-8611, Japan

19

- 20 <sup>7</sup> Corresponding author.
- 21 TEL/FAX: 81-138-40-5519
- 22 EMAIL: kishi@fish.hokudai.ac.jp

23

24 **Short title:** Structural properties of dulse phycoerythrin

#### **ABSTRACT**

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

We found that the red alga dulse (Palmaria palmata) contains a lot of proteins, which is mainly composed of phycoerythrin (PE), and the protein hydrolysates showed high angiotensin I converting enzyme (ACE) inhibitory activities. Therefore, we investigated the structure of dulse PE to discuss its structure-function relationship. We prepared the chloroplast DNA and analyzed the nucleotide sequences encoding PE by cDNA cloning method. It was clarified that dulse PE has  $\alpha$ - and  $\beta$ -subunits and they are composed by 164 amino acids (MW: 17,638) and 177 amino acids (MW: 18,407), respectively. The dulse PE contained conserved cysteine residues for chromophore attachment site. On the alignment of amino acid sequences of dulse PE with those of other red algal PE, the sequence identities were very high (81-92%). In addition, we purified and crystallized the dulse PE, and its crystal structure was determined at 2.09 Å resolution by molecular replacement method. The revealed 3-D structure of dulse PE which forms an (αβ hexamer was similar to other red algal PEs. On the other hand, it was clarified that the dulse PE proteins are rich in hydrophobic amino acid residues (51.0%), especially aromatic amino acid and proline The data imply that the high ACE inhibitory activity of dulse protein hydrolysates would be caused by the specific amino acid composition and sequence of dulse PE.

43

44

45

#### PRACTICAL APPLICATIONS

- Dulse is an abundant and underused resource, which contains a lot of phycobiliproteins.
- 47 Then, the dulse protein hydrolysates strongly inhibited the activity of angiotensin I converting
- 48 enzyme. Therefore, it has the potential to be an ingredient of functional food.

49

- **KEYWORDS:** Red alga; Dulse; *Palmaria palmata*; ACE inhibitory activity; phycoerythrin;
- 52 Primary structure; 3-D structure

#### INTRODUCTION

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

54

In red algae, phycobiliproteins locate as phycobilisomes on the stromal side of thylakoid membranes in a chloroplast and play a role of light capturing on photosynthesis (Apt et al. 1995; Sekar and Chandramohan 2008). The prominent classes of red algal phycobiliproteins are phycoerythrin (PE) followed by phycocyanin (PC) and allophycocyanin (APC), and they are divided on their spectral properties ( $\lambda$ -max of PE = 490-570 nm,  $\lambda$ -max of PC = 610-625 nm,  $\lambda$ -max of APC = 650-660 nm) (Sun et al. 2009). Phycobiliproteins of red algae commonly contain α- and β-subunits, and each subunit bears covalently binding one or several phycobilin chromophores at the specific cysteine residues (PE: phycoerythrobilin and phycourobilin, PC: phycocyanobilin and phycoerythrobilin, APC: phycocyanobilin) (Apt et al. The above spectroscopic property of each phycobiliprotein is derived from the specific chromophore composition. The  $\alpha$ - and  $\beta$ -subunits of phycobiliprotein combine with each other to form an  $(\alpha\beta)$  heterodimer, and then three  $(\alpha\beta)$ s form  $(\alpha\beta)$  trimer arranging a symmetry disc (Apt et al. 1995). The discs are organized in supramolecular complexes called phycobilisomes. The core of phycobilisomes is composed of APC discs and the rod is composed of PC and PE discs. On the previous proteomic and genomic studies, some marine red algal phycobiliproteins were studied (Roell and Morse 1993; Ducret et al. 1994; Hagopian et al. 2004; Niu et al. 2006; Tajima et al. 2012; Wang et al. 2013; DePriest et al. 2013). However, there is no information about structural properties of dulse phycobiliproteins. Dulse (Palmaria palmata) is a red alga mainly distributed in high-latitude coastal areas, and it is popular in Ireland and Atlantic Canada as a food and a source of minerals. Fitzgerald et al. (2012) and Harnedy et al. (2015) also reported the dulse protein hydrolysates show the inhibitory effects for renin and dipeptidyl peptidase IV, respectively. In Japan,

dulse is also distributed around the coast of Hokkaido Prefecture and at Pacific coast of Aomori Prefecture. However, dulse is rarely eaten in Japan. In addition, dulse is even removed from Kombu (*Laminaria* sp.) farming areas in Hokkaido, because it inhibits the growth of young Kombu in winter season. Therefore, we have begun exploring the health benefits of dulse to advance its use as a functional food material. In the previous study, we found that dulse contains a lot of proteins, which are mainly composed of PE followed by PC and APC (Furuta *et al.*, 2016). Then, the dulse protein hydrolysates strongly inhibited the activity of angiotensin I converting enzyme (ACE). Moreover, it was suggested that the ACE inhibitory peptides are mainly derived from the dulse PE by thermolysin hydrolysis. Therefore, in this study, we investigated the primary and 3-D structures of dulse PE to discuss its structure-function relationship.

#### MATERIALS AND METHODS

92 Materials

Dulse (*P. palmata*) was collected in the coast of Usujiri, Hokkaido, Japan in February. A portion of the thalli was steeped into RNAlater solution (Applied Biosystems, CA, USA) and stored at -80 °C until use.

Restriction enzymes, *Hin*d III and *Ssp* I, were purchased from TaKaRa Bio (Shiga, Japan). RNase A was purchased from Nacalai Tesque (Kyoto, Japan). ACE from rabbit lung was purchased from Sigma Chemical Co. (Mo, USA). Hyppuryl-L-histidyl-L-leucine (Hip-His-Leu), thermolysin (EC 3.4.24.27) from *Bucillus thermoproteolyticus*, pepsin (EC 3.4.23.1) from porcine stomach, and trypsin (EC 3.4.21.4) from bovine pancreas were purchased from Wako Pure Chemical (Osaka, Japan). All other regents were purchased from Wako Pure Chemical (Osaka, Japan).

Preparation of dulse protein hydrolysates

The frozen samples were lyophilized and ground into a fine powder by Wonder Blender WB-1 (OSAKA CHEMICAL Co., Osaka, Japan). Proteins were extracted from the powder by adding 20 v/w of distilled water at 4 °C for 7 h. The extracts were centrifuged (H-200, Kokusan, Tokyo, Japan) at 4 °C, 15,000 x g for 10 min, and then the supernatants were used as "dulse proteins". Some of the dulse proteins were hydrolyzed by 1.0 wt% of thermolysin at 70 °C for 3 h, and the reaction was terminated by heat treatment at 100 °C for 5 min. Subsequently, the solution was centrifuged at 4 °C, 15,000 x g for 10 min. The supernatants were dried by lyophilization into the "thermolysin hydrolysates". Other dulse proteins were

adjusted to pH 2.0, and the proteins were digested by 1.0 wt% of pepsin at 37 °C for 3 h. After the reaction, the pepsin digests were adjusted to pH 8.0. Subsequently, the solutions were centrifuged at 4 °C, 15,000 x g for 10 min. The supernatants were dried by lyophilization into the "pepsin hydrolysates". Some of the pepsin hydrolysates were digested by 1.0 wt% of trypsin at 37 °C for 3 h. After that, the digested solutions were boiled for 5 min to inactivate the enzymes, and then centrifuged at 4 °C, 15,000 x g for 10 min. The supernatants were dried by lyophilization into the "pepsin-trypsin hydrolysates".

122

123

115

116

117

118

119

120

121

#### **ACE Inhibitory Assay**

124

ACE inhibitory assay was carried out according to the method of Cheung and Cushman 125 126 (1973) with some modifications. Fifteen microliters of sample solution (5.0 mg/mL) was added to 30 Find the CE t (B72° C/for 5 anish the mixture was

127

128

131

132

133

134

135

136

Thirty microliters of Hip-His-Leu solution (12.5 mM in 0.1 M sodium borate buffer

129 containing 400 mM NaCl at pH 8.3) was added to the mixture. After incubation at 37 °C for

130 1 h, the reaction was stopped by adding 75

□L of 1.0 M

was extracted with 450 µL of ethyl acetate. Four hundred microliters of the upper layer was evaporated, and then the hippuric acid was dissolved in 1.5 mL of distilled water. absorbance at 228 nm of the solution was measured by a spectrophotometer. The inhibition was calculated from the equation [1- (As-Asb) / (Ac-Acb)] x 100, where Ac is the absorbance of the buffer, Acb is the absorbance when the stop solution was added to the buffer before the reaction, As is the absorbance of the sample, and Asb is the absorbance when the stop solution was added to the sample before the reaction.

138

137

#### Isolation of Dulse Chloroplast DNA

141

142

143

144

145

146

147

148

149

150

151

152

153

140

Thawed dulse sample was dissected with scissors and 150 mg of it was put into a microcentrifuge tube. The sample was homogenized in 1.5 mL of TRIzol reagent (Invitrogen, CA, USA) using disperser. Then, 300 µL of chloroform was added to the homogenate, and the solution was mixed. The mixture was centrifuged at 4 °C, 15,000 x g for 20 min, and the supernatant was pooled in a micro tube. Next, equal volume of 2-propanol was added in the tube to precipitate chloroplast DNA, and the solution was centrifuged at 4 °C, 15,000 x g for 20 min. The precipitate was dissolved in 100 µL of TE buffer, and the remaining RNA in it was removed by RNase A treatment (10 µg, 37 °C, 30 After the reaction, 200 µL of sterilized ultrapure water and 300 µL of min). phenol-chloroform-isoamyl alcohol (25:24:1, v/v/v) were added, and the mixed solution was centrifuged at 4 °C, 15,000 x g for 15 min. Following similar treatment with chloroform-isoamyl alcohol (24:1, v/v), chloroplast DNA was collected by ethanol precipitation. The dried precipitate was dissolved in 100 µL of TE buffer.

155

154

#### Degenerate PCR

157

158

159

160

161

162

163

164

156

Forward primer (PE-F1: ATGCT (A/C/G/T) AA (C/T) GC (A/C/G/T) TTTTC (A/C/G/T) (A/C) G) and reverse primer (PE-R1: CC (A/C/G/T) GC (A/G/T) AT (A/C/G/T) CCCCA (C/T) TC (A/G) TC) for degenerate PCR were designed on the basis of well-conserved regions of red algal PE genes (*rpeB* and *rpeA*) (Fig. 1a). TaKaRa EX *Taq* Hot Start Version (TaKaRa Bio, Shiga, Japan) was used on the amplification. The PCR program for TaKaRa EX *Taq* HS was 40 cycles of 98 °C for 10 sec, 47 °C for 30 sec, 72 °C for 2 min, and 10 min at 72 °C. The PCR products were separated by low melting agarose gel electrophoresis and

165 were purified from the gel using Wizard SV Gel and PCR Clean-Up System (Promega, WI, 166 USA). 167 168 Inverse PCR 169 170 The remaining 5'- and 3'-regions of dulse PE genes were determined by inverse PCR method. 171 Dulse chloroplast DNA was digested with restriction enzymes, Ssp I and Hind III. The 172 digested DNA fragments were cleaned by Mini Elute Spin Columns (QIAGEN, Dusseldorf, 173 Germany), and ligated with T4 DNA ligase (TaKaRa Bio, Shiga, Japan) at 16 °C for 18 h. For amplifications, specific forward (PE-IF1: CATTACTGATGGTAACAAACGC, PE-IF2: 174 GAGACGTTGATCATTATATGCG) and reverse (PE-IR1: TCACTGCCACCAACGTAAGC, 175 176 PE-IR2: CTCCACCTTCTTTTACAACAGC) primers were designed using the sequence data 177 determined by degenerate PCR (Fig. 1b). TaKaRa EX Taq Hot Start Version (TaKaRa Bio, 178 Shiga, Japan) was used on the amplification, and the PCR program was 40 cycles of 98 °C for 179 10 sec, 50 °C for 30 sec, 72 °C for 2 min, and 10 min at 72 °C. 180 181 Cloning and Sequencing 182 183 PCR products were subcloned to pDrive Cloning Vector using QIAGEN PCR Cloning Kit (QIAGEN, Dusseldorf, Germany) for sequencing. The nucleotide sequences of cDNAs 184 185 were determined with BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, 186 CA, USA) using ABI PRISM 310 Genetic Analyzer (Applied Biosystems, CA, USA). 187 Nucleotide and deduced amino acid sequences of dulse PE gene were aligned using 188 CLUSTAL W program (Thompson, et al. 1994). Molecular weight and isoelectric point of

dulse PE were calculated from deduced amino acid sequences by using Compute pI/Mw tool

190 (Bjellqvist et al. 1993; Bjellqvist et al. 1994; Hoogland et al. 2000).

191

Crystallization, X-ray diffraction data collection, and structure determination

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

192

Frozen dulse samples (-30 °C) were taken into a flask, and 4 volumes (v/w) of distilled water was added in it. The dulse phycobiliproteins were extracted at 4 °C for 12 h, and the extracts were filtered. Then, the filtrates were centrifuged at 4 °C, 15,000 x g for 15 min. The extracted dulse proteins were dialyzed against distilled water at 4 °C for 24 h. The dulse PE was purified from the protein extracts by a preparative electrofocusing using Rotofor system (Bio-Rad, CA, USA) (Fig. 4a and 4b). Crystallization was carried out by hanging-drop vapor diffusion method. Crystals of dulse PE were grown from a buffer containing 0.1 M sodium acetate (pH 4.8) and 12% PEG4000 (Fig. 4b). X-ray diffraction dataset of dulse PE was collected on the beamline BL17A at Photon Factory (Tsukuba, Japan) under cryogenic condition (100 K). Crystals were mounted on the X-ray diffractometer after soaked into a crystallization buffer containing 20% PEG400 as a cryoprotectant. The diffraction data were indexed, integrated, scaled, and merged using the XDS program (Kabsch 2010). The data statistics are shown in Table 1. Crystal structures were determined by the molecular replacement method with the program MOLREP (Vagin and Teplyakov 1997) using the structure of PE from *Polysiphonia urceolata* (PDB ID 1LIA) as a search model. To monitor the refinement, a random 5% subset was set aside for the calculation of the R<sub>free</sub> factor. Structure refinement was carried out with phenix.refine (Adams et al. 2010). stereochemical quality of the structure was analyzed with the program MOLPROBITY (Chen et al. 2010). The refinement statistics are summarized in Table 1. The atomic coordinates of dulse PE has been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 5B13).

#### RESULTS AND DISCUSSION

215

214

216 Inhibition of ACE activity of dulse protein hydrolysates

217

218

219

220

221

222

223

224

225

226

227

228

229

230

231

232

233

234

235

236

237

238

In the previous study, we found that dulse contains a lot of proteins, which are mainly composed of PE (Furuta et al. 2016). The extracted dulse proteins showed slight ACE inhibitory activity, but the activity was extremely enhanced by thermolysin hydrolysis. In addition, nine ACE inhibitory peptides (YRD, AGGEY, VYRT, VDHY, IKGHY, LKNPG, LDY, LRY, FEQDWAS) were isolated from the hydrolysates by reversed-phase high-performance liquid chromatography (HPLC), and the sequences of YRD, AGGEY, VYRT, VDHY, LKNPG, LDY and LRY were detected in the primary structures of PE α- and β-subunits (Furuta et al. 2016). From these results, it was suggested that the ACE inhibitory peptides are mainly derived from the dulse PE by thermolysin hydrolysis. Therefore, in this study, we prepared the dulse protein hydrolysates by thermolysin, pepsin, and pepsin-trypsin digestion, and we compared with their ACE inhibitory activity. As shown in Fig. 2, the thermolysin hydrolysates inhibited 88% of ACE activity, and pepsin and pepsin-trypsin hydrolysates also suppressed 72% and 75% of them, respectively. We calculated the peptide sequences derived from the deduced amino acid sequences of dulse PE  $\alpha$ - and  $\beta$ - subunits by using PEPTIDEMASS (Wilkins et al. 1997). As a result, it was predicted that 76 peptides (α-subunit: 38 peptides, av. length=3, av. mass=346; β- subunit: 38 peptides, av. length=4, av. mass=396) are derived from dulse PE  $\alpha$ - and  $\beta$ - subunits by pepsin-trypsin hydrolysis. From the result, ACE inhibitory peptides are also produced from dulse proteins, especially PE, by proteolytic hydrolysis in our digestive tract. In future, we would like to analyze the structural properties of ACE inhibitory peptides in the pepsin-trypsin hydrolysates to compare with those of thermolysin hydrolysates.

Then, in the next stage, we investigated the primary and 3-D structures of dulse PE to discuss its structure-function relationship.

241

239

240

Nucleotide sequences of dulse phycoerythrin genes

243

244

245

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

262

263

242

In this study, we obtained 1,560 bp of nucleotide sequences on the analysis of the dulse PE gene, and the gene structure encoding dulse PE (*rpeA* and *rpeB*, GenBank accession number AB625450) (Fig. 3) was clarified. This is the first report for the PE gene of Palmariales.

As shown in Fig. 3, the dulse PE gene was constituted of  $\alpha$ - and  $\beta$ -subunit genes and A/T-rich spacer. AT contents of the spacer in dulse PE gene were 79% (60 bp/76 bp). Bernard et al. (1992) reported that rpeB gene of Rhodella violacea is split by intervening sequence and the sequence has a feature of group II intron that is typical in eukaryotic organisms, however the dulse PE gene has no introns. The dulse rpeB was present in prior to the rpeA (Fig. 3). The positions of rpeA and rpeB were the same as those of other red algae, for example Gracilaria tenuistipitata (Hagopian et al. 2004), Chondrus crispus (GenBank accession number HF562234), Pyropia yezoensis (Wang et al. 2013), P. haitanensis (Wang et al. 2013) and P. purpurea (GenBank accession number U38804). The nucleotide sequences of dulse PE gene also showed considerably high identities (about 80%) with those of other red algae (Table 2). The GC contents in dulse PE gene were about 40% (rpeA: 40.2%, rpeB: 40.5%), and these numerical values showed very high similarity to those of P. yezoensis (rpeA: 42.6%, rpeB: 40.6%), P. haitanensis (rpeA: 41.2%, rpeB: 41.4%) and P. purpurea (rpeA: 41.8%, rpeB: 42.0%), whereas it was a little higher than those of G. tenuistipitata (rpeA: 37.0%, rpeB: 38.8%) and C. crispus (rpeA: 37.2%, rpeB: 39.1%) (Table 2).

The consensus sequences at -10 (5'-TATAAT-3') and -35 (5'-TTGACA-3')

promoter elements for RNA polymerase were searched in the dulse PE genes. As a result, putative motifs were found at upstream regions of *rpeB* (-10: TATATT or TGTAAT, -35: TAAACA or GAAACA) (single and double underlines in Fig. 3). We also sought out the Shine-Dalgarno sequence (5'-AGGAGGT-3') acting as a binding site with 16S rRNA, and then the homologous structures were detected in the upstream of each gene (*rpeB*: AGGAGA, *rpeA*: AGGAGA,) (dotted underlines in Fig. 2).

Primary structure of dulse phycoerythrin

The deduced amino acid sequences of dulse PE  $\alpha$ - and  $\beta$ -subunits are shown in Fig. 3. The PE  $\alpha$ -subunit consists of 164 amino acids (495 bp), and its molecular weight and isoelectric point were calculated at 17,638 and 5.40, respectively. Red algal PE commonly has two kinds of chromophores, phycoerythrobilin and phycourobilin. Generally, red algal PE  $\alpha$ -subunit binds to two phycoerythrobilin with two Cys residues (Lundell *et al.* 1984; Ficner *et al.* 1992), and the dulse PE  $\alpha$ -subunit also retained Cys residues at the corresponding positions ( $\alpha$ Cys82 and  $\alpha$ Cys139 in Fig. 3 and Fig. 4a). The dulse PE  $\beta$ -subunit consists of 177 amino acids (534 bp), and its molecular weight and isoelectric point were calculated at 18,407 and 5.42, respectively. It is already known that one phycourobilin and two phycoerythrobilins bind to four Cys residues in  $\beta$ -subunit apo-protein through thioether linkage (Lundell, *et al.* 1984; Ficner *et al.* 1992). In the dulse PE  $\beta$ -subunit, corresponding Cys residues binding with phycourobilin ( $\beta$ Cys50 and  $\beta$ Cys61 in Fig. 3 and Fig. 4b) and with phycoerythrobilins ( $\beta$ Cys82 and  $\beta$ Cys158) were all conserved.

3-D structures of dulse phycoerythrin

We purified and crystallized the dulse PE (Fig.5a), and its crystal structure was determined by molecular replacement method (Fig.5b and Table 1). The revealed 3-D structure of purified dulse PE in this study formed an  $(\alpha\beta \Box hexamer, which was similar to other red algal PEs$ (Chang et al. 1996; Contreras-Martel et al. 2001; Ritter et al. 1999). The root mean square deviations (r.m.s.d) with other PEs are as follows, Polysiphonia urceolata PE: 0.70 Å, Griffithsia monilis PE: 0.55 Å, Gracilaria chilensis PE: 0.60 Å. As observed for other homologous phycobiliproteins such as PE, PC and APC, the backbone conformations of αand β-subunits of dulse PE have nine α-helices (X, Y, A, B, E, F', F, G, and H) as a dominant secondary structure element (Fig. 5b) (Lundell et al. 1984; Ficner et al. 1992; Liu et al. 1999; Jiang et al. 2001). Each subunit had a structure quite similar to those of other PEs. The r.m.s.d. was 0.39 Å, 0.33 Å, and 0.37 Å for  $\alpha$ -subunit, and 0.56 Å, 0.48Å, and 0.55Å for β-subunit of *P. urceolata* PE, *G. monilis* PE and *G. chilensis* PE, respectively. The electron density clearly showed the presence of chromophores covalently linked to Cys residue through thioether bond. Phycoerythrobilins were linked covalently with each of aCys82, αCys139, βCys82, and βCys158, whereas a phycourobilin was linked doubly to βCys50 and BCys61. The presence of chromophores at these sites is highly conserved among PEs of which structures have been reported (Camara-Artigas et al. 2012; Chang et al. 1996; Contreras-Martel et al. 2001; Lundell et al. 1984; Ritter et al. 1999). observations together, we concluded that dulse PE has structural characteristics common to other PEs.

309

310

308

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

#### Structure-function relationship of dulse phycoerythrin

311

312

313

ACE is a key enzyme in the regulation of peripheral blood pressure catalyzing the production of angiotensin II and the destruction of bradykinin (Cheung *et al.* 1980). The specific

inhibitors of the enzyme therefore have been considered with effective antihypertensive drugs. In addition to the drugs, ACE inhibitory peptides from daily food are also useful for maintaining blood pressure at a healthy level. Although the potency of peptide is lower than drug, it does not have side effect (Balti et al. 2015). Up to now, many researchers have identified various ACE inhibitory peptides from the enzymatic hydrolysates of food (Amado et al. 2014; Ghassem et al. 2014; Balti et al. 2015; García-Moreno et al. 2015). Besides, Cheung et al. (1980) obtained the interesting results by using several synthetic peptides for a substrate of ACE, that is to say, the ACE inhibitory activity of peptide is closely related to the C-terminal dipeptide residues in it. Specifically, in case of tryptophan, tyrosine, or proline residue is located at the N-terminal side of dipeptide and aromatic amino acid or proline residue is at the C-terminus, its inhibitory potency is the most. Indeed, it has been well known that the peptides are usually composed of hydrophobic and aromatic amino acids (Amado et al. 2014; Ghassem et al. 2014; Balti et al. 2015; García-Moreno et al. 2015). Therefore, we calculated the contents of hydrophobic and aromatic amino acid residues in dulse PE by using the primary structures in this study (Fig. 3). As a result, it was clarified that the dulse PE are rich in hydrophobic amino acids (51.0%), especially the contents of aromatic amino acids and proline (10.0-10.9%) are relatively high. On the other hand, crystal structure analysis clearly showed that dulse PE shares significant similarity in their tertiary structure with other PEs. Therefore, we concluded that the cause of high ACE inhibitory activity of dulse PE hydrolysates would be the specific amino acid compositions and sequences, independent of the tertiary structure.

314

315

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

#### **ACKNOWLEDGMENTS**

We thank Dr. Koji Mikami, Faculty of Fisheries Sciences, Hokkaido University, for the technical assistance of inverse PCR. We thank Dr. Hiroki Saeki, Faculty of Fisheries Sciences, Hokkaido University, for the technical assistance of preparative electrofocusing using Rotofor system. We also thank Mr. Yuki Kato, Hokkaido Industrial Technology Center, for the operative support of DNA sequencer.

This work was supported in part by the Regional Innovation Cluster Program (Global Type), Ministry of Education, Culture, Sports, Science and Technology, Japan and the Grant-in-Aid for High Technology Research Program from the Ministry of Education, Culture, Sports, Science, and Technology of Japan. This work was supported in part by Grants-in-Aid for Scientific Research (24000011, 20374225, and 16H00748 to YT) and Platform for Drug Discovery, Informatics, and Structural Life Science from the Ministry of Education, Culture, Sports, Science and Technology, Japan, and JST, PRESTO (YT).

## 350 **REFERENCES**

- 352 APT, K.E., COLLIER, J.L. and GROSSMAN, A.R. 1995. Evolution of the phycobiliproteins.
- 353 J. Mol. Biol. 248, 79-96.
- 354 ADAMS, P.D., AFONINE, P.V., BUNKOCZI, G., CHEN, V.B., DAVIS, I.W., ECHOLS, N.,
- 355 HEADD, J.J., HUNG, L.W., KAPRAL, G.J., GROSSE-KUNSTLEVE, R.W., MCCOY,
- 356 A.J., MORIARTY, N.W., OEFFNER, R., READ, R.J., RICHARDSON, D.C.,
- 357 RICHARDSON, J.S., TERWILLIGER, T.C. and ZWART, P.H. 2010. PHENIX:
- a comprehensive Python-based system for macromolecular structuresolution.
- 359 Acta Crystallogr. *D66*, 213-221.
- 360 AMADO, I.R., VAZQUEZ, J.A., GONZALEZ, P., ESTEBAN-FERNANDEZ, D.,
- 361 CARRERA, M. and PINEIRO, C. 2014. Identification of the major ACE-inhibitory
- peptides produced by enzymatic hydrolysis of a protein concentrate from cuttlefish
- 363 wastewater. Marine Drugs *12*, 1390-1405.
- 364 BERNARD, C., THOMAS, J.C., MAZEL, D., MOUSSEAU, A., CASTETS, A.M., DE
- 365 MARSAC, N.T. and DUBACQ, J.P. 1992. Characterization of the genes encoding
- phycoerythrin in the red alga *Rhodella violacea*: evidence for a splitting of the *rpeB* gene
- 367 by an intron. Proc. Natl. Acad. Sci. U.S.A. 89, 9564-9568.
- 368 BJELLQVIST, B., HUGHES, G., PASQUALI, C., PAQUET, N., RAVIER, F., SANCHEZ,
- J.-C., FRUTIGER, S. and HOCHSTRASSER, D.F. 1993. The focusing positions of
- polypeptides in immobilized pH gradients can be predicted from their amino acid
- sequences. Electrophoresis 14, 1023–1031.
- 372 BJELLQVIST, B., BASSE, B., OLSEN, E. and CELIS, J.E. 1994. Reference points for
- comparisons of two-dimensional maps of proteins from different human cell types defined
- in a pH scale where isoelectric points correlate with polypeptide compositions.

375 Electrophoresis 15, 529–539. 376 BALTI, R., BOUGATEF, A., SILA, A., GUILLOCHON, D., DHULSTER, P. and 377 NEDJAR-ARROUME, N. 2015. Nine novel angiotensin I-converting enzyme (ACE) 378 inhibitory peptides from cuttlefish (Sepia officinalis) muscle protein hydrolysates and 379 antihypertensive effect of the potent active peptide in spontaneously hypertensive rats. 380 Food Chem. 170, 519-525. 381 CHEUNG, H.S. and CUSHMAN, D.W. 1973. Inhibition of homogeneous 382 angiotensin-converting enzyme of rabbit lung by synthetic venom peptides of *Bothrops* 383 jararaca. Biochim. Biophys. Acta 293, 451-463. 384 CHEUNG, H.S., WANG, F.L., ONDETTI, M.A., SABO, E.F. and CUSHMAN, D.W. 1980. 385 Binding of peptide substrates and inhibitors of angiotensin-converting enzyme: 386 importance of the COOH-terminal dipeptide sequence. J. Biol. Chem. 25, 401-407. 387 CHANG, W.R., JIANG, T., WAN, Z.L., ZHANG, J.P., YANG, Z.X. and LIANG, D.C. 1996. 388 Crystal structure of R-phycoerythrin from Polysiphonia urceolata at 2.8 Å resolution. J. 389 Mol. Biol. 262, 721-31. 390 CONTRERAS-MARTEL, C., MARTINEZ-OYANEDEL, J., BUNSTER, M., LEGRAND, P., 391 PIRAS, C., VERNEDE, X. and FONTECILLA-CAMPS, J.C. 2001. Crystallization and 392 2.2 Å resolution structure of R-phycoerythrin from Gracilaria chilensis: a case of perfect 393 hemihedral twinning. Acta Crystallogr. D57, 52-60. 394 CHEN, V.B., ARENDALL 3rd. W.B., HEADD, J.J., KEEDY, D.A., IMMORMINO, R.M., 395 KAPRAL, G.J., MURRAY, L.W., RICHARDSON, J.S. and RICHARDSON, D.C. 2010. 396 MolProbity: all-atomstructure validation for macromolecular crystallography. Acta

397

Crystallogr. *D66*, 12-21.

- 398 CAMARA-ARTIGAS, A., BACARIZO, J., ANDUJAR-SANCHEZ, M.,
- ORTIZ-SALMERON, E., MESA-VALLE, C., CUADRI, C., ... and ALLEN, J.P. 2012.
- 400 pH-dependent structural conformations of B-phycoerythrin from Porphyridium cruentum.
- 401 FEBS J. 279, 3680-91.
- 402 DUCRET, A., SIDLER, W., FRANK, G. and ZUBER, H. 1994. The complete amino acid
- sequence of R-phycocyanin-I  $\alpha$  and  $\beta$  subunits from the red alga *Porphyridium cruentum*:
- structural and phylogenetic relationship of the phycocyanins within the phycobiliprotein
- 405 families. Eur. J. Biochem. 221, 563-580.
- 406 DEPRIEST, M.S., BHATTACHARYA, D. and LOPEZ-BAUTISTA, J.M. 2013. The plastid
- genome of the red macroalga *Grateloupia taiwanensis* (Halymeniaceae). PLoS One, 8,
- 408 e68246.
- 409 FICNER, R., LOBECK, K., SCHMIDT, G. and HUBER, R. 1992. Isolation, crystallization,
- structure analysis and refinement of B-phycoerythrin from the red alga *Porphyridium*
- 411 *sordidum* at 2.2 Å resolution. J. Mol. Biol. 228, 935-950.
- 412 FITZGERALD, C., MORA-SOLER, L., GALLAGHER, E., O'CONNOR, P., PRIETO, J.,
- SOLER-VILA, A. and HAYES, M. 2012. Isolation and characterization of bioactive
- pro-peptides with *in vitro* renin inhibitory activities from the macroalga *Palmaria*
- 415 *palmata*. J. Agric. Food Chem. 60, 7421-7427.
- 416 FURUTA, T., MIYABE, Y., YASUI, H., KINOSHITA, Y. and KISHIMURA, H. 2016.
- 417 Angiotensin I converting enzyme inhibitory peptides derived from phycobiliproteins of
- dulse *Palmaria palmata*. Marine Drugs, 14, 32; doi:10.3390/md14020032.
- 419 GHASSEM, M., BABJI, A.S., SAID, M., MAHMOODANI, F. and AEIHARA, K. 2014.
- 420 Angiotensin I-converting enzyme inhibitory peptides from snakehead fish sarcoplasmic
- protein hydrolysate. J. Food Biochem. 38, 140-149.

- 422 GARCIA-MORENO, P.J., ESPEJO-CARPIO, F.J., GUADIX, A. and GUADIX, E.M. 2015.
- Production and identification of angiotensin I-converting enzyme (ACE) inhibitory
- peptides from Mediterranean fish discards. J. Func. Foods 18, 95-105.
- 425 HOOGLAND, C., SANCHEZ, J.-C., TONELLA, L., BINZ, P.-A., BAIROCH, A.,
- 426 HOCHSTRASSER, D.F. and APPEL, R.D. 2000. The 1999 SWISS-2DPAGE database
- 427 update. Nucleic Acids Res. 28, 286–288.
- 428 HAGOPIAN, J.C., REIS, M., KITAJIMA, J.P., BHATTACHARYA, D. and DE OLIVEIRA,
- M.C. 2004. Comparative analysis of the complete plastid genome sequence of the red alga
- 430 Gracilaria tenuistipitata var. liui provides insights into the evolution of rhodoplasts and
- their relationship to other plastids. J. Mol. Evol. *59*, 464-477.
- 432 HARNEDY, P.A., O'KEEFFE, M.B. and FITZGERALD, R.J. 2015. Purification and
- identification of dipeptidyl peptidase (DPP) IV inhibitory peptides from the macroalga
- 434 *Palmaria palmata*. Food Chem. 172, 400-406.
- JIANG, T., ZHANG, J.P., CHANG, W.R. and LIANG, D.C. 2001. Crystal structure of
- 436 R-phycocyanin and possible energy transfer pathway in the phycobilisomes. Biophys. J.
- 437 *81*, 1171-1179.
- 438 KABSCH, W. 2010. Xds. Acta Crystallogr. *D66*, 125-132.
- 439 LUNDELL, D.J., GLAZER, A.N, DELANGE, R.J. and BROWN, D.M. 1984. Bilin
- attachment sites in the  $\alpha$  and  $\beta$ -subunits of B-phycoerythrin: amino acid sequence
- 441 studies. J. Biol. Chem. 259, 5472-5480.
- 442 LIU, J.Y., JIANG, T., ZHANG, J.P. and LIANG, D.C. 1999. Crystal structure of
- allophycocyanin from red algae *Porphyra yezoensis* at 2.2- Å resolution. J. Biol. Chem.
- 444 *274*, 16945-16952.
- NIU, J.F., WANG, G.C. and TSENG, C.K. 2006. Method for large-scale isolation and
- purification of R-phycoerythrin from red alga *Polysiphonia urceolata* Grev. Protein Expr.

- 447 Purif. 49, 23-33.
- 448 ROELL, M.K. and MORSE, D.E. 1993. Organization, expression and nucleotide sequence of
- the operon encoding R-phycoerythrin

Polysiphosido units from the red

- 450 *boldii*. Plant Mol. Biol. 21, 47-58.
- 451 RITTER, S., HILLER, R. G., WRENCH, P. M., WELTE, W. and DIEDERICHS, K. 1999.
- 452 Crystal structure of a phycourobilin-containing phycoerythrin at 1.90- Å resolution. J.
- 453 Struc. Biol. 126, 86–97.
- 454 SEKAR, S. and CHANDRAMOHAN, M. 2008. Phycobiliproteins as a commodity: trends in
- applied research, patents and commercialization. J. Appl. Phycol. 20, 113-136.
- 456 SUN, L., WANG, S., GONG, X., ZHAO, M., FU, X. and WANG, L. 2009. Isolation,
- purification and characteristics of R-phycoerythrin from a marine macroalga
- 458 *Heterosiphonia japonica*. Protein Expr. Purif. *64*, 146-154.
- THOMPSON, J.D., HIGGINS, D.G. and GIBSON, T.J. 1994. CLUSTAL W: improving the
- sensitivity of progressive multiple sequence alignment through sequence weighting,
- positions-specific gap penalties and weight matrix choice. Nucleic Acids Res. 22, 4673-
- 462 4680.
- 463 TAJIMA, N., SATO, S., MARUYAMA, F., KUROKAWA, K., OHTA, H., TABATA, S.,
- SEKINE, K., MORIYAMA, T. and SATO, N. 2012. Analysis of the complete chloroplast
- genome of the unicellular red alga *Porphyridium purpureum*. Photosynthetic Res. 22,
- 466 156-159.
- VAGIN, A. and TEPLYAKOV, A. 1997. MOLREP: an automated program for molecular
- replacement. J. Appl. Crystallogr. 30, 1022-1025.
- 469 WILKINS M.R., LINDSKOG I., GASTEIGER E., BAIROCH A., SANCHEZ J.C.,
- 470 HOCHSTRASSER D.F. and APPEL R.D. 1997. Detailed peptide characterization using

471	PEPTIDEMASS - a World-Wide-Web-accessible tool. Electrophoresis, 18, 403-408.
472	WANG, L., MAO, Y.X., KONG, F.N., LI, G.Y., MA, F., ZHANG, B.L., SUN, P.P., BI,
473	G.Q., ZHANG, F.F., XUE, H.F. and CAO, M. 2013. Complete sequence and analysis of
474	plastid genomes of two economically important red algae: Pyropia haitanensis and
475	Pyropia yezoensis. PLoS One, 8, e65902.

476	(Captions to figures)
477	FIG. 1. GENERAL STRUCTURES OF RED ALGAL PHYCOERYTHRIN GENES AND
478	POSITIONS OF PRIMERS USED IN DEGENERATE AND INVERSE PCRS.
479	a: Positions of primers used in degenerate PCR.
480	b: Positions of primers used in inverse PCR.
481	PE represent phycoerythrin. Sequences of each primer are shown in the text.
482	Restriction sites are expressed as Ssp I, Hind III.
483	
484	FIG. 2. ACE INHIBITORY ACTIVITIES BY DULSE PROTEIN HYDROLYSATES.
485	1: ACE inhibitory activity with thermolysin hydrolysates.
486	2: ACE inhibitory activity with pepsin hydrolysates.
487	3: ACE inhibitory activity with pepsin-trypsin hydrolysates.
488	
489	FIG. 3. NUCLEOTIDE AND DEDUCED AMINO ACID SEQUENCES OF DULSE
490	PHYCOERYTHRIN GENE.
491	Asterisks show stop codon. Single and double underlines express putative -10 and
492	-35 consensus sequences, respectively. Dotted underline is putative RNA
493	polymerase-binding motif.
494	
495	FIG. 4. ALIGNMENT OF AMINO ACID SEQUENCES OF RED ALGAL
496	PHYCOERYTHRINS.
497	a: $PE\alpha$ ; Phycoerythrin $\alpha$ -subunit.
498	b: PE $\beta$ ; Phycoerythrin $\beta$ -subunit.
499	Asterisks show characteristic amino acid residues in the molecule.  P. palmata
500	(GenBank accession number: AB625450, in this study); G. tenuistipitata (AY673996), C.

501	crispus (HF562234), P. yezoensis (D89878), P. haitanensis (DQ449070), P. purpurea (U38804).
502	
503	FIG. 5. DULSE PHYCOERYTHRIN CRYSTAL AND 3-D STRUCTURE OF DULSE
504	PHYCOERYTHRIN.
505	a: Crystallization of purified dulse phycoerythrin.
506	Purified PE: purified dulse phycoerythrin. PE crystal: dulse phycoerythrin crystal.
507	b: 3-D structure of dulse phycoerythrin.
508	PE (αβ mer: Ribbon representation of dulse phycoerythrin
509	$(\alpha\beta)$ mæthe α- and β-subunits are colored red and blue, respectively. For clarity,
510	one subunit of $\alpha$ - and $\beta$ -subunit is colored orange and green, respectively. The bound CYC
511	and PUB are also shown as yellow and green sticks, respectively. PEα: Ribbon representation
512	of dulse phycoerythrin $\alpha$ -subunit. The model is colored according to the sequence from blue
513	at the N-terminus to red at the C-terminus. Bound CYC chromophores are shown as yellow
514	sticks. The cysteine resides linked with the chromophres are also shown. PEβ: Ribbon
515	representation of dulse phycoerythrin $\beta$ -subunit colored according to the sequence from blue
516	at the N-terminus to red at the C-terminus. Bound CYC and PUB chromophores are shown as
517	vellow and green sticks

 TABLE 1
 DATA COLLECTION AND REFINEMENT STATISTICS

Data collection						
Beamline	Photon Factory BL17A					
Space group	C2					
Cell dimensions						
a, b, c (Å)	187.5, 111.9, 112.7					
α, β, γ (°)	90.0, 91.9, 90.0					
Wavelength (Å)	0.98					
Resolution (Å) $^a$	50-2.09 (2.22-2.09)					
No. of total/unique reflections	519,606/135,827 (81,130/21,390)					
$R_{\mathrm{sym}}$ (%) <sup>a, b</sup>	11.6 (69.9)					
Completeness (%) <sup>a</sup>	99.5 (97.6)					
Multiplicity <sup>a</sup>	3.8 (3.8)					
Average $I/\sigma(I)^a$	11.21 (2.13)					
Refinement						
Resolution (Å)	50–2.09					
$ m R_{work}/R_{free}$	0.198/0.237					
No. of atoms						
Protein	15,114					
Ligand	1,290					
Solvent	1,812					
r.m.s.d.						
Bond lengths (Å)	0.003					
Bond angles (°)	1.318					
Ramachandran plot						
Favored (%)	97.6					
Allowed (%)	2.4					
Outlier (%)	0					

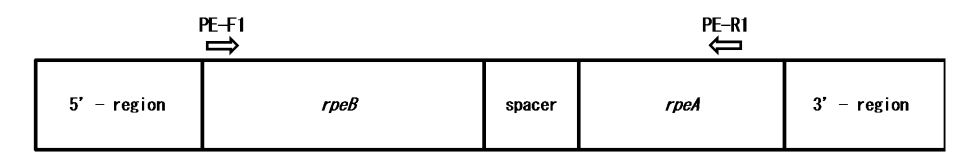
 $<sup>\</sup>it a$  Values in parentheses correspond to the highest resolution shell.

 $b~R_{\rm merge} = \Sigma_{\rm h}~\Sigma_{\rm i}~|I_{\rm h,i}-<\!\!I_{\rm h}\!\!>\!\!|/\Sigma_{\rm h}\Sigma_{\rm i}~|I_{\rm h,i}|,$  where  $<\!\!I_{\rm h}\!\!>$  is the mean intensity of a set of equivalent reflections.

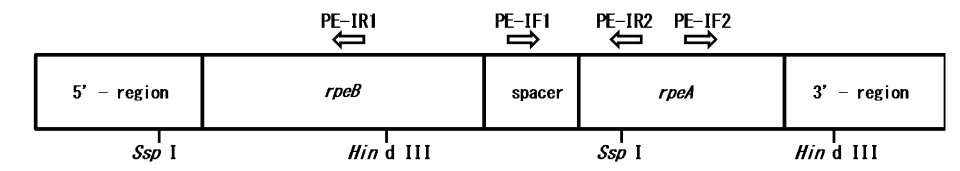
**TABLE 2.** GC CONTENT, NUCLEOTIDE IDENTITY, AND AMINO ACID IDENTITY ON RED ALGAL PHYCOERYTHRINS

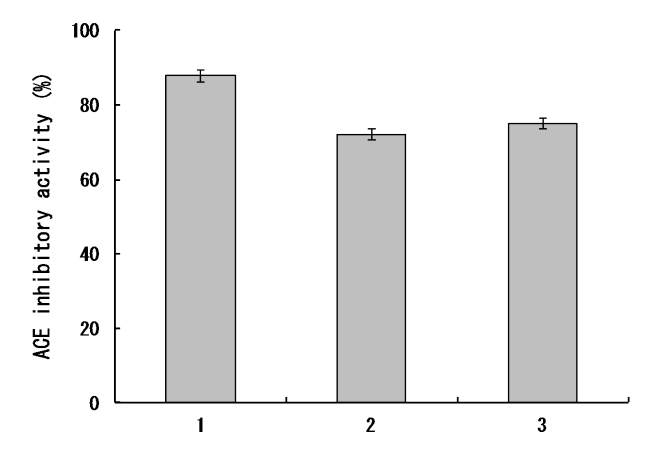
Organism		Gene name	GC content (%)	Nucleotide identity to <i>P.palmata</i>	Amino acid identity to <i>P.palmata</i>	Accession No.	
D. I	DE	α-subunit	40.2	(%)	(%)		
Palmaria palmata	naria palmata PE		40.2			AB625450	
		β-subnuit	40.5		_		
Gracilaria tenuistipitata	PE	α-subunit	37.0	79	87	AY673996	
		β-subnuit	38.8	78	81		
Chondrus crispus	PE	α-subunit	37.2	82	85	HF562234	
		β-subnuit	39.1	80	85	HF302234	
Porphyra yezoensis	PE	α-subunit	42.6	82	89	D89878	
		β-subnuit	40.6	82	92		
Porphyra haitanensis	PE	α-subunit	41.2	83	90	<b>Y Y Y Y Y Y Y Y Y Y</b>	
		β-subnuit	41.4	83	92	HM008261	
Porphyra purpurea	PE	α-subunit	41.8	41.8 82 90			
		β-subnuit	42.0	83	92	NC_000925.1	

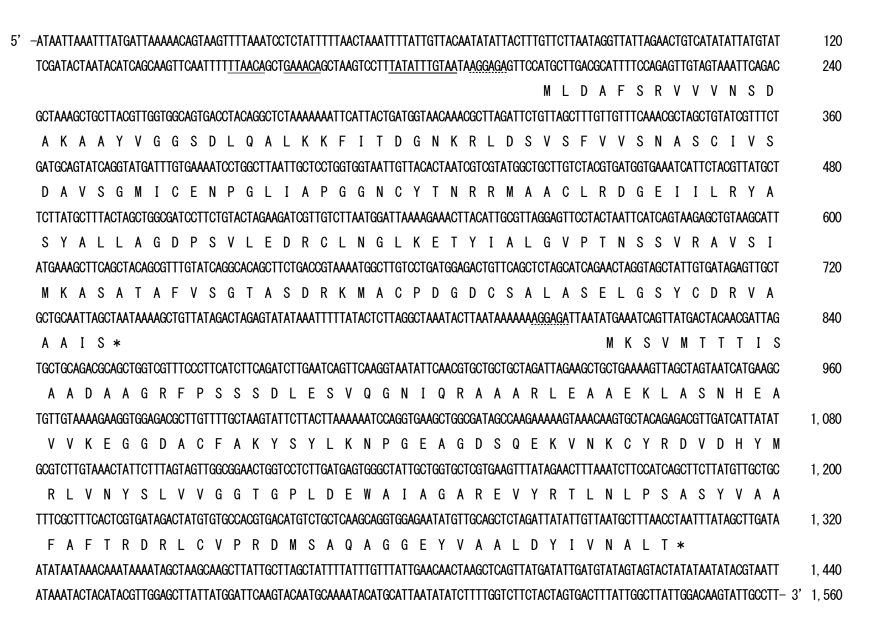
# a: degenerate PCR



## b: invers PCR







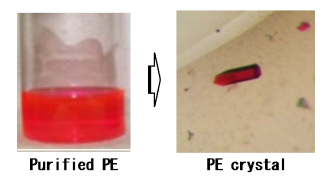
a: PEα						
	1	10	20	30	40	50
P. palmata	MKSVMTT	TISAADAAG	RFPSSSDLE	SVQGNIQRAA	ARLEAAEKLA	SNHEA
G. tenuistipitata	₩KSVITT	VISAADAAG	RFPSSSDLE	SIQGNIQRAS	ARLEAAEKLA	DNHDA
C. crispus	MKSVITT	IISAADAAG	RFLTSSDLE	SVQGNIQRAG	ARLEAAEKLA	NNHEA
P. yezoensis	MKSVITT	TIGAADAAG	RFPSSSDLE	SVQGNIQRAA	ARLEAAEKLA	SNHEA
P. haitanensis	MKSVITT	TISAADAAG	RFPSSSDLE	SVQGNIQRAA	ARLEAAEKLA	SNHEA
P. purpurea	MKSVITT	TISAADAAG	RFPSSSDLE	SVQGNIQRAA	ARLEAAEKLA	SNHEA
	**	*	* *			
		60	70	80	90	100
P. palmata	VVKEGGD	ACFAKYSYL	KNPGEAGDS	QEK <b>VNK</b> CYRD	VDHY <b>M</b> rlvny	SLVVG
G. tenuistipitata	VVKEAGD	ACFGKYSYL	KNAGEAGEN	QEKVNKCYRD	I DHYMRLVNY	SLVVG
C. crispus	VVKEAGD	ACFAKYSFL	KNSGEAGDS	QEKVNKCYRD	IDHYMRLINY.	ALIVG
P. yezoensis	VVKEAGD	ACFAKYSYL	KNPGEAGDS	QEKVNKCYRD	VDHYMRLVNY	CLVVG
P. haitanensis	VVKEAGD	ACFAKYSYL	KNPGEAGDS	QEKVNKCYRD	VDHYMRLVNY	CLVVG
P. purpurea	VVKEAGD	ACFAKYSYL	KNPGEAGDS	QEKVNKCYRD	VDHY <b>m</b> rlvny	CLVVG
				*	*	
		110	120	130	140	150
P. palmata	GTGPLDE	WAIAGAREV	YRTLNLPSA	SYVAAFAFTR	DRLCVPRD <b>M</b> S	AQAGG
G. tenuistipitata	GTGPLDE	WAIAGAREV	YRTL <b>N</b> LPTS.	AYVAAFAFTR	DRLCVPRD <b>M</b> S	AQAGV
C. crispus	GTGPFDE	WGIAGAREV	YRALNLPSA:	SYLAAFVFTR	DRLCVPRD <b>M</b> S	AQAGL
P. yezoensis	GTGPVDE	WGIAGAREV	YRTLNLPTS	AYVASFAFAR	DRLCVPRD <b>M</b> S	AQAGV
P. haitanensis	GTGPVDE	WGIAGAREV	YRTL <b>N</b> LPTS.	AYVASFAFAR	DRLCVPRD <b>M</b> S	AQAGV
P. purpurea	GTGPVDE	WGIAGAREV	YRTLNLPTS	AYVASFAFAR	DRLCVPRDMS	AQAGV
	*				*	
		160				
P. palmata	EYVAALD	YIVNALT				
G. tenuistipitata	EYTTALD	YIINSLS				
C. crispus	EYGAALD	YVINSLS				
P. yezoensis	EYAGNLD	YLINALS				
P. haitanensis	EYAGNLD	YIINSLC				
P. purpurea	EYAGNLD	YIINSLC				

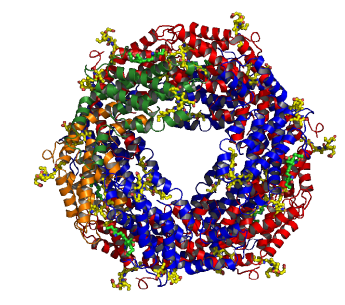
b: PEB 10 20 30 50 1 40 MLDAFSRVVVNSDAKAAYVGGSDLQALKKFITDGNKRLDSVSFVVSNASC P. palmata G. tenuistipitata MLDAFSRVVIDSDTKAAYVGGSNLQALKTFISEGNQRLDAVNSIVSNASC C. crispus MLDAFSRVVVNSDAKAAYVGGSDLQALKTFIADGNKRLDAVNSIVSNASC MLDAFSRVVVNSDAKAAYVGGSDLQALKKFIADGNKRLDSVNAIVSNASC P. yezoensis P. haitanensis MLDAFSRVVVNSDAKAAYVGGSDLQALKKFIADGNKRLDSVNAIVSNASC P. purpurea MLDAFSRVVVNSDAKAAYVGGSDLQALKKFIADGNKRLDSVNAIVSNASC \* 60 70 80 90 100 IVSDAVSGNICENPGLIAPGGNCYTNRRMAACLRDGEIILRYASYALLAG P. palmata IVSDAVSGNICENPGLTSPGGNCYTNRRMAACLRDGEIILRYISYALLAG G. tenuistipitata C. crispus IVSDAVSGMICENPGLIAPGGNCYTNRRMAACLRDGEIILRYISYALLAG IVSDAVSGMICENPGLIAPGGNCYTNRRMAACLRDGEIILRYVSYALLAG P. yezoensis P. haitanensis IVSDAVSGMICENPGLIAPGGNCYTNRRMAACLRDGEIILRYVSYALLAG P. purpurea IVSDAVSGMICENPGLIAPGGNCYTNRRMAACLRDGEIILRYVSYALLAG \* 110 120 130 140 150 P. palmata DPSVLEDRCLNGLKETYIALGVPTNSSVRAVSIMKASATAFVSGTASDRK G. tenuistipitata DPSVLEDRCLNGLKETYIALGVPITSSARAVNINKASVAAFILNTAPGRK C. crispus DASVLEDRCLNGLKETYIALGVPNNSSIRSVVIMKAAAVAFVNNTASQRK P. yezoensis DPSVLEDRCLNGLKETY I ALGVPTNSSVRAVS I NKAAAVAFITNTASQRK P. haitanensis DPSVLEDRCLNGLKETYIALGVPTNSSVRAVSIMKAAAVAFITNTASQRK P. purpurea DPSVLEDRCLNGLKETYIALGVPTNSSVRAVSINKASAVAFITNTASQRK \* 160 170 MACPDGDCSALASELGSYCDRVAAAIS P. palmata G. tenuistipitata **MDTASGDCTALASEVGSYFDRVCAAIS** MATTSGDCSALSAEVASYCDRVGAALS C. crispus P. yezoensis MATADGDCSALASEVASYCDRVAAAIS MATADGDCSALASEVASYCDRVAAAIS P. haitanensis P. purpurea MATADGDCSALASEVASYCDRVAAAIS

k

b

a





PE( $\alpha\beta$ )<sub>6</sub> hexamer

