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# Isolation and Identification of 3-Methylcrotonyl Coenzyme A Carboxylase cDNAs and Pyruvate Carboxylase, and Their Expression in Red Seabream (Pagrus major) Organs 

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#### Abstract

We determined complementary DNA sequences of biotin-containing (MCCC1) and non-biotincontaining (MCCC2) subunits of 3-methylcrotonyl coenzyme A carboxylase (MCCase) and pyruvate carboxylase (PCase) using reverse transcriptase polymerase chain reaction of RNA extracted from seabream skeletal muscle and liver. We determined the complete coding sequences of MCCC1 and PC and a partial coding sequence of the major part of MCCC2. Molecular sizes of MCCC1, MCCC2, and PC were 4300, 2400, and 6500 nucleotides, respectively, according to Northern blot analysis. The length of $M C C C 1$ from cDNA sequencing was 4249 nucleotides, indicating the full-length messenger RNA sequence was obtained. Northern blot analyses showed that PC was expressed in muscle, heart, liver, and ovary, but not in spleen. MCCCI and MCCC2 were expressed at high levels in muscle and ovary, but only trace levels in heart, spleen, and liver. MCCase appears to be particularly important in muscle and ovary, which are active in protein metabolism, while PCase is important in organs active in glycolysis, such as liver.


Key words: red seabream, MCCase, PCCase, PCase, biotin, mRNA.

## Introduction

Pyruvate carboxylase (PCase, EC 6.4.1.1), propionyl-coenzyme A (CoA) carboxylase (PCCase, EC 6.4.1.3), and 3-methylcrotonyl-CoA carboxylase (MCCase, EC 6.4.14) are known as mitochondrial biotin-dependent carboxylases that play a pivotal role in organic acid metabolic pathways. PCase has been found widely in diverse organisms, and its structure and functions are well characterized. PCase cat-

[^0]alyzes the ATP-dependent carboxylation of pyruvate to form oxaloacetate in gluconeogenesis, lipogenesis, synthesis of certain amino acids, and formation of neurotransmitters (Jitrapakdee et al., 1996; Wallace et al., 1998).

PCase was first discovered in chicken (Utter and Keech, 1963), and many studies on PCase have been performed in vertebrates, invertebrates (Crabtree et al., 1972), and bacteria (Goss et al., 1981). The gene or complementary DNA encoding PCase has also been reported in many organisms including zebrafish (Yoder and Litman, 2000) but not yet in any commercial fishery species. MCCase has been implicated as a component enzyme of the "mevalonate shunt"
(Popjak, 1971), it catalyzes the carboxylation of 3-meth-ylcrotonyl-CoA to form 3-methylglutaconyl-CoA, and it is involved in the catabolism of leucine and isovalerate (Moss and Lane, 1971). MCCase has also been isolated from soybean (Song et al., 1994), tomato (Wang et al., 1994), Arabidopsis thaliana (Weaver et al., 1995), and mammals (Lau et al., 1980).

The full-length cDNAs encoding MCCase have been obtained from soybean and tomato for the biotin, containing subunit (Song et al., 1994, Wang et al., 1994), and from Arabidopsis for the non-biotin-containing subunit (McKean et al., 2000). Both the cDNA and the genomic sequences encoding MCCase in human have been demonstrated for the biotin-containing subunit (Obata et al., 2001) and for the non-biotin-containing subunit (Holzinger et al., 2001), but comparable sequences have not been fully reported in fish.

These biotin-dependent enzymes are also important in nutritional and physiologic aspects, and numerous studies have been carried out on biotin-dependent carboxylase deficiency in humans and other animals. Biotin deficiency in humans is characterized by decreased ability of the carboxylation reactions that affect the metabolism of carbohydrates, fats, and proteins, resulting in retarded growth, as well as other developmental and morphologic abnormalities (Wolf et al., 1981; Mock et al., 1985). In other animals biotin deficiency can cause poor growth, dermal lesions such as parakeratosis, alopecia, and achromotrichia, and fetal and other abnormalities, including biological changes such as depression of the activities of biotindependent enzymes (Whitehead, 1985). Biotin deficiency is also teratogenic in some species of mammals such as in mice (Watanabe, 1983) and human (Zempleni and Mock, 2000).

In the commercial fishery industry, both biotin dietary supplementation and its relation to biotin-dependent enzyme levels have been studied. The symptoms of biotin deficiency are as poor growth, low food consumption, increased mortality, degeneration of the gill lamellae, and skin disease (BASF, 1998) Biotin requirements for optimal growth of fish have been studied for rainbow trout (Woodward and Frigg, 1989), for mirror carp (Gunther and Meer-Burgdorff, 1990), and also for trout and salmon (National Research Council, 1973). However, the molecular basis of physiology and nutrition related to the biotindependent enzymes has not been established yet in fish.

We report here cloning and analysis of complete cDNA sequences for the biotin-containing subunits of MCCase
(MCCC1) and PCase (PC), and identification of PCase, MCCase, and PCase proteins in a member of the Sparidae, red seabream (Pagrus major), which is an important commercial fish. We also identified the cDNA encoding the non-biotin-containing subunit of MCCase (MCCC2) and studied the coexpression of genes encoding PCase and subunits of MCCase in various organs from red seabream.

## Materials and Methods

## Protein Extraction and Isolation of Biotinylated Proteins

Proteins were extracted from skeletal muscle of 2 to 3 -yearold red seabream (Pagrus major). Muscle tissue was ground in 5 volumes of CDB containing 200 mM Tris- $\mathrm{HCl}(\mathrm{pH}$ 8.5), $450 \mathrm{mM} \mathrm{KOAc}, 25 \mathrm{mM} \mathrm{Mg}(\mathrm{OAc})_{2}$, and $2 \%$ PTE (Abe and Davies, 1995) with 0.1 mM phenylmethylsulfonyl fluoride (PMSF) using a mortar and pestle at ice-cold temperature, then transferred to a Teflon homogenizer to continue homogenization at 5 strokes per second for 15 to 20 minutes. The homogenate was filtered through Miracloth, then centrifuged for 15 minutes at $27,000 \mathrm{~g}$ (Kubota model 6800 -rotor, type RA300). The supernatant was filtered through a membrane filter (ADVANTEC) using a vacuum pump, then applied to an avidin affinity column prepared by conjugating egg white avidin to an NHSactivated Sepharose column (Hitrap-NHS activated) from Amersham Biotech according to the manufacturer instructions, using a peristatic pump at $0.5 \mathrm{ml} / \mathrm{min}$ flow rate. The effluent was applied again to the same column at least 3 times. After the application the column was washed with 5 volumes of $\mathrm{CDB}, 5$ volumes of a washing buffer containing 5 mM Hepes-KOH ( pH 7.5), 10 mM $\mathrm{Mg}(\mathrm{OAc})_{2}, 2 \mathrm{mM}$ EGTA, and 0.1 mM PMSF, and 5 volumes of 1 mM biotin solution in the washing buffer. After the column was washed proteins were eluted with 5 ml of 6 M guanidine thiocyanate into 10 fractions. Each fraction was precipitated with 4 volumes of acetone at $-20^{\circ} \mathrm{C}$ overnight and analyzed for proteins using sodium dodecylsulfate polyacrylamide gel electrophoresis (SDS-PAGE).

## SDS-PAGE and Detection of Biotin-Containing Proteins

The protein samples obtained from the avidin affinity column chromatography were dissolved in a sample buffer
containing 2\% LDS and 0.01 M Tris- HCl (pH 6.8), 20\% (v/ v) glycerol, $1 \%(\mathrm{v} / \mathrm{v}) 2$-mercaptoethanol, and heated for 5 minutes at $95^{\circ} \mathrm{C}$. These samples were electrophoresed in an 8.5\% acrylamide gel, and transferred onto a PVDF membrane (Immobilon-P, Millipore) at $2 \mathrm{~mA} \mathrm{~cm}{ }^{-2}$ for 1 hour using an electroblotting apparatus (BE300, Biocraft) for detection of biotin-containing proteins. To detect biotincontaining proteins, the blotted membrane was blocked in $5 \%$ dry milk in TBS for 1 hour, washed with TBS $(10 \mathrm{mM}$ Tris $\mathrm{HCl}[\mathrm{pH} 7.6], 0.8 \% \mathrm{NaCl}$ ) containing $0.5 \%$ Tween 20) 5 times, then hybridized with a streptavidin-alkaline phosphatase conjugate for 30 minutes. After washing the membrane 5 times with TBS, the binding of alkaline phosphatase conjugate was detected in 100 mM diethanolamine ( pH 9.5 ) with BCIP and NBT as substrates.

## Determination of Partial Amino Acid Sequences of Avidin Binding Proteins Digested by V8 Protease

The avidin-affinity column fractions containing avidinbinding proteins were pooled by separated by (SDS-PAGE). Bands of avidin-binding proteins stained with Brilliant Blue R-250 were cut out and loaded on a stacking gel, and the protein was digested with Staphylococcus aureus V8 protease (EC 3.4.21.19) by the method described by Cleveland et al. (1977). The digested polypeptides were separated on $15 \%$ acrylamide gels using SDS-PAGE and electroblotted onto a PVDF membrane (Immobilon-P, Millipore). The transferred peptides were detected by staining with Brilliant Blue R-250, and peptide bands were excised and sequenced using a protein sequencer (model 476A, PerkinElmer, Applied Biosystems Division). These streptavidin-binding proteins are referred to as $S B P$ s hereafter.

## Cloning and cDNA Sequence Analysis

Partial nucleotide sequences were obtained by RT-PCR using primers derived from the homologous nucleotide sequences corresponding to the genes coding for MCCase and PCase of human (BAA99407), mouse (NP_076133), and Arabidopsis (Q42523). The PCR products obtained were subcloned into pMOSblue vector, and the positive colonies were isolated and purified using a FlexiPrep kit (Amersham Pharmacia Biotech). DNA sequencing was performed using a BigDye Terminator Cycle Sequencing Ready Reaction Kit and ABI PRISM 377 DNA Sequencer (PE Applied Biosystems). The $3^{\prime}$ - and 5 '-cDNA extremities were determined by the rapid amplification of cDNA ends (RACE) method.

The poly $(A+)$ RNA was purified from total RNA (100 $\mu \mathrm{g}$ ) using the mRNA Purification kit (Amersham Pharmacia Biotech). The blunt-ended double-stranded cDNA was synthesized by incubating $1 \mu \mathrm{~g}$ of poly(A+) RNA with 10 pmol gene-specific primers under a ReverTra Ace reverse transcriptase (TOYOBO Co., Ltd.) for 1 hour at $42^{\circ} \mathrm{C}$. This was followed by the simultaneous inclusion of DNA polymerase I $\left(9 \mathrm{U} \cdot \mu \mathrm{g}^{-1}\right)$, Escherichia coli DNA ligase ( $6 \mathrm{U} \cdot \mu \mathrm{g}^{-1}$, Takara), and T4 DNA polymerase ( $9 \mathrm{U} \cdot \mu \mathrm{g}^{-1}$ ) in the presence of RNase $\mathrm{H}\left(1 \mathrm{U} \cdot \mathrm{Ll}^{-1}\right)$ in a reaction buffer composed of 33.4 mM Tris- $\mathrm{HCl}(\mathrm{pH} 8.5), 4.6 \mathrm{mM} \mathrm{MgCl} 2_{2}, 10 \mathrm{mM}$ $\left(\mathrm{NH}_{4}\right)_{2} \mathrm{SO}_{4}, 6.6 \mathrm{mM}$ 2-mercaptoethanol, 0.06 mM EDTA, $0.005 \%$ bovine serum albumin (BSA) 100 mM KCl , and $0.15 \mathrm{mM} \beta-\mathrm{NAD}$, and incubated at $16^{\circ} \mathrm{C}$ for 90 minutes. The double-stranded cDNA obtained was ligated with 12.5 pmol of an adapter consisting of complementary oligo DNA (5'-gtaatacgactcactatagggcacgcgtggtcgacggcccgggctg gt-3') and ( $3^{\prime}$-gggcccgacca-5'), using $4 \mathrm{U} \cdot \mu \mathrm{g}^{-1} \mathrm{~T} 4$ DNA ligase at $16^{\circ} \mathrm{C}$ overnight. (T4 DNA ligase, DNA polymerase I, and RNase $H$ were obtained from TOYOBO).

RACE-PCR was performed in $25-\mu$ l-volume reactions amplified by the adapter primer ( 5 '-gtaatacgactcactatagggc3') with gene-specific primers using a XL-PCR kit (PE Applied Biosystem). The target fragments from the RACEPCR were subcloned and sequenced as above. Nucleotide sequences were analyzed using the basic local alignment search tool (BLAST) algorithm program of the National Center for Biotechnology Information, and deduced amino acid sequences were analyzed with the CLUSTAL W multiple alignment algorithm (DNA Data Bank of Japan) and the GENETYX-WIN (Version 6.1.0) tree-generation program.

## Primers for Cloning

Gene-specific primers for cloning cDNA fragments, along with the adapter primer, are summarized in Table 1. A forward primer, $\mathrm{MCCC} 1-\mathrm{Fa}$, and a reverse primer, $\mathrm{MCCC} 1-\mathrm{Ra}$, were derived from conserved $M C C C 1$ sequences in human (AB029826) and fugu genome databases (CAAB01000411). A forward primer, $\mathrm{MCCCl}-\mathrm{Fb}$, and a reverse primer, MCCC1 Rb , were designed from the partial MCCC1 sequence obtained in the present study (AB081831), for amplifying the $3^{\prime}-$ and $5^{\prime}-\mathrm{cDNA}$ ends, respectively.

To obtain a cDNA fragment for MCCC2, a forward primer, MCCC2-Fa, and a reverse primer, MCCC2-Ra, were designed using conserved $M C C C 2$ sequences for human (AB029826) and Arabidopsis (AF059510) MCCC2

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Table 1. Gene Specific Primers for Cloning cDNA Fragments for MCCC1, MCCC2, and PC
Name of primer

MCCC1-Fa
MCCCI-Ra
$\mathrm{MCCCl}-\mathrm{Fb}$
MCCC1-Rb
MCCC2-Fa
MCCC2-Ra
MCCC2-Rb
$\mathrm{PC}-\mathrm{Fa}$
$\mathrm{PC}-\mathrm{Fb}$
PC-Ra
PC-Fc
PC-Rb
PC-Rc
Tag primer Adapter primer

Sequence 5' - $3^{\prime}$
Source
tgcaggtggaacatcctgt
gccactttgtctccagcttt
cgtgtacaacaaagacggaag
cgagcttggcgatcattggg
cgcaagcagggaaccat
cacatgccgtagtttcc
gctctccatttagcaagaaaggcg
caacttcagcaacctcttcag
gaggttgtacgcaagatggg
agtgtaggttggtgtactgg
tgccctggagaaggtgtttg
tcctggagtctcttccatgg
agagatggctatggagcgag
ctgaaggttccagaatcgat
gtaatacgactcactataggge

AB029826, CAAB01000411
AB029826, CAAB01000411
AB081831
AB081831
AB029826, AF059510
AB029826, AF059510
AB082535
AF295372
AF295372
AF295372
AB081832
AB081832
AB081832
cDNAs. For amplifying the 5 '-cDNA end, a reverse primer, MCCC2-Rb, was designed using the partial sequence for MCCC2 obtained in the present study (AB082535).

To obtain a cDNA fragment for $P C$, forward primers $\mathrm{PC}-\mathrm{Fa}$ and $\mathrm{PC}-\mathrm{Fb}$ and a reverse primer, $\mathrm{PC}-\mathrm{Ra}$, were designed using the zebrafish sequence for PCase (AF295372). A forward primer, $\mathrm{PC}-\mathrm{Fc}$, for the $3^{\prime}-\mathrm{cDNA}$ end and reverse primers, $P C-R b$ and $P C-R c$, for the 5 '-cDNA end were designed from the partial $P C$ sequence obtained in the present study (AB081832). The adapter and tag primers for amplifying the 5'- and $3^{\prime}$-cDNA ends, respectively, are also shown in Table 1.

## Northern Blotting Analyses

Total RNA was isolated from heart, spleen, brain, muscle, liver, and ovary tissues from the same fish, using a RNA Extraction Kit (Amersham Pharmacia Biotech). Each total RNA ( $20-\mu \mathrm{g}$ ) fraction was denatured at 65 C for 5 minutes in $1 \times$ MOPS buffer ( 20 mM MOPS, 5 mM sodium acetate, 1 mM EDTA, pH 7.0 ), 2.2 M formaldehyde, and $50 \%(\mathrm{v} / \mathrm{v})$ formamide. The denatured RNA was separated on a $1.0 \%$ agarose gel containing 0.66 M formaldehyde, then transferred to a Hybond XL nylon membrane (Amersham Pharmacia Biotech), and the membrane was baked at $80^{\circ} \mathrm{C}$ for 2 hours. The membranes with immobilized RNA were pre hybridized for 30 minutes at $42^{\circ} \mathrm{C}$ in a plastic bag in hybridization buffer containing $5 \times \mathrm{SSC}$
( $750 \mathrm{mM} \mathrm{NaCl}, 75 \mathrm{mM} \mathrm{Na} 3$-citrate), $5 \times$ Denhardt's reagent ( $0.1 \%[\mathrm{w} / \mathrm{v}]$ BSA, $0.1 \%[\mathrm{w} / \mathrm{v}]$ Ficoll-400, 0.1\% [w/ v] polyvinylpyrrolidone), $5 \%$ dextran sulfate, $0.5 \% \mathrm{SDS}$, $100 \mu \mathrm{~g} \cdot \mathrm{ml}^{-1}$ salmon sperm DNA, and $50 \%(\mathrm{v} / \mathrm{v})$ formamide. Then the randomly primed ${ }^{32} \mathrm{P}$-labeled probe (specific activity, $1.0 \times 10^{6} \mathrm{cpm} \cdot \mu \mathrm{g}^{-1}$ ) was added to the bag, and hybridization was carried out at $42^{\circ} \mathrm{C}$ overnight with constant agitation.

The membranes were washed at $60^{\circ} \mathrm{C}$ twice with $2 \times$ SSC and $0.1 \%$ SDS for 5 minutes, followed by $1 \times$ SSC and $0.1 \%$ SDS at $60^{\circ} \mathrm{C}$ for 15 minutes, and finally washed at $60^{\circ} \mathrm{C}$ twice with $0.1 \times \mathrm{SSC}$ and $0.1 \% \mathrm{SDS}$ for 10 minutes. The membrane was exposed to Imaging Plate BAS III (Fuji Fihn, Co. Ltd.) for 12 hours and analyzed using Bio Image Analyzer (BAS-2000, Fuji Film, Co. Ltd.). The membrane was hybridized with $\alpha_{-}^{32} \mathrm{P}$-labeled cDNA probes of MCCC1, MCCC2, and PC obtained in the present study. The same membrane was consecutively reprobed with $\alpha_{-}^{32} \mathrm{P}$-labeled cDNA probes for $M C C C 1, M C C C 2$, and $P C$. The hybridized probe was stripped after each probing. For preparing probes a Rediprime II random prime labeling system was used (Amersham Pharmacia Biotech). The same specific activity ( $1.0 \times 10^{6} \mathrm{cpm} \cdot \mu \mathrm{g}^{-1}$ ) of randomly primed $\alpha-{ }^{32} \mathrm{P}$-labeled probe was hybridized with each probe at $42^{\circ} \mathrm{C}$ overnight in $5 \times \mathrm{SSC}\left(750 \mathrm{mM} \mathrm{NaCl}, 75 \mathrm{mM} \mathrm{Na}_{3}-\right.$ citrate) containing hybridization buffer.

Radioactivity on the probed membrane was visualized and quantified by BAS-2000. The amounts of total RNA
loaded were equalized according to densitometric estimation of relative amounts in agarose gels.

## Results

## Isolation of Biotin-Containing Proteins and Their Identification as PCase, MCCase, and PCCase

Figure 1 shows Brilliant Blue-stained proteins obtained by avidin-affinity column chromatograpy in an SDS-PAGE gel and detection of SBP using streptavidin-conjugated alkaline phosphatase with BCIP and NBT as substrates as described in "Materials and Methods." Six major proteins, of 209 $121,76,72,56$, and 43 kDa , were visible in the stained gel (lane 2). Among them, 3 protein bands, of 121,76 , and 72 kDa , reacted with streptavidin-conjugated alkaline phosphatase; therefore, they are annotated as $121-\mathrm{kDa}$ SBP, 76kDa SBP, and $72-\mathrm{kDa}$ SBP (lane 3).

Table 2 shows the results of analyses of partial amino acid sequencing and the homology search in databases for the peptide fragments obtained by V8 proteinase digestion. A fragment of $121-\mathrm{kDa}$ SBP was $100 \%$ identical to zebrafish, human, and mouse PCases; a fragment of $76-\mathrm{kDa} \mathrm{SBP}$ was $100 \%$ identical to MCCase from human and mouse; and a fragment of $72-\mathrm{kDa}$ SBP was $90 \%$ identical to human and mouse PCCase. According to the molecular mass, the $56-\mathrm{kDa}$ protein band was considered to be a mixture of non-biotin-containing subunits of MCCase and PCCase (Oei and Robinson, 1985). The 209-kDa and 43-kDa proteins were not identified in this study.

## Cloning of cDNAs Encoding the Biotin-Containing Subunit of MCCase (MCCC1)

A partial nucleotide sequence of 949 nucleotides of red seabream cDNA encoding 316 amino acids (AB081831) was obtained by RT-PCR with a forward primer, MCCC1Fa, and a reverse primer, MCCC1-Ra. 5' RACE was performed to obtain the 5 ' end of cDNA sequence using a gene-specific reverse primer, $\mathrm{MCCC1}-\mathrm{Rb}$, and the adapter primer. Sequences assembled from these RT-PCR products revealed a cDNA of 2140 nucleotides. 3' RACE was then performed using the tag primer and a gene-specific primer, $\mathrm{MCCC} 1-\mathrm{Fb}$, and furnished a 2109 nucleotide-long 3' sequence containing the stop codon and the canonical poly (A) signal ( $5^{\prime}$-aataaa- $3^{\prime}$ ), which was also polyadenylated at the 3' end. The compiled sequence for red seabream


Figure 1. Isolation and detection of biotin-containing proteins from fish skeletal muscle tissue. Avidin-binding proteins obtained from skeletal muscle tissue of red seabream were separated by SDS-PAGE, electroblotted onto PVDF membrane, and either stained with Brilliant Blue R-250 (lane 2) or processed for detection of biotin-containing proteins using streptavidin-conjugated alkaline phosphatase with NBT and BCIP as substrates (lane 3). Brilliant Blue-stained molecular mass markers are shown in lane 1 . Major protein bands in lane 2 are annotated at $43,56,72,78,121$, and 209 kDa . The SBP bands in lane 3 are protein bands that reacted with the streptavidin conjugate.

MCCC1 was 4249 nucleotides long, and it was deposited in the database (AB100429). This full-length cDNA sequence contained a canonical translation initiation motif, [a/ g]nnatgg (Kozak, 1999), at 185 to 191, with the initiating methionine beginning at 188 and a stop codon occurring at 2339. This defined an open reading frame of 2151 nucleotides that putatively encoded a polypeptide of 717 amino acid residues. An in-frame stop codon at base 12 occurs upstream of the putative initiating methionine, consistent with the methionine codon at 188 being identified as the true initiating methionine and the recovered 5 ' sequences as the 5 'untranslated region (UTR). The partial amino acid sequences of V8 fragments (Table 2: E-XQVFGDMHG) of the $76-\mathrm{kDa}$ biotin-containing protein, along with the cleavage site E for V 8 proteinase, were also located in the encoded amino acid sequence at 925 to 952 .

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5 3 2 ~ S h u n n o s u k e ~ A b e ~ e t ~ a l . ~
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Table 2. Identification of SBP by Partial Protein Sequencing ${ }^{\text {a }}$

| SBP (kDA) | V8 fragment (kDa) | Sequence | Identification | Identity (\%) | Protein ID | Species |
| :--- | :--- | :--- | :--- | :--- | :--- | ---: |
| 121 | 15 | NEIPGGQYTN | PCase | 100 | AF295372 | Zebrafish |
|  |  |  |  |  | L09192 | Mouse |
| 76 | 25 | XQVFGDMHG | MCCase | 100 | BC011617 | Human |
| 72 | 27 | AAPIGYPVMI | PCCase | 90 | BAA99407 | Human |
|  |  |  |  | LXM_049016 | Human |  |

${ }^{\text {a }}$ Molecular masses of SBP were calculated from the results shown in Figure 1.

Figure 2 shows a comparison of the amino acid sequences between the encoded polypeptide and MCCases from different organisms. All of the hallmarks for MCCase were present: CPSase L-chain domain (37-149), CPSase Lchain D2 domain (152-267), biotin carboxylase $c$ domain (372-479), biotin lipoyl biotin requiring enzyme domain (640-692), biotin-binding motif (AMKME), and the binding site (K); D-Ala D-Ala ligase domain (135-327), which is a component of D-Ala-D-Ala ligase enzymes (EC:6.3.2.4).

The overall homologies of the red seabream protein to the biotin-containing subunits of MCCases were $73 \%$ to mouse (Mus musculus), $71 \%$ to human (Homo sapiens), $57 \%$ to mosquito (Anopheles gambiae str.), $56 \%$ to fruit fly (Drosophila melanogaster), and $47 \%$ to Arabidopsis (Arabidopsis thaliana), while low overall homologies $(40 \%)$ to the biotin containing subunit of PCCases from human (XM_049016) and mouse (NM_144844) were obtained. There was also $49 \%$ overall homology between the red seabream protein and a biotin-containing car-bamoyl-phosphate synthase L chain from a nematode (Caenorhabditis elegans (NP_501777).

## Cloning of cDNA Encoding the Non-BiotinContaining Subunit of MCCase (MCCC2)

We also cloned a cDNA fragment encoding most of the coding sequence of the non-biotin-containing subunit of MCCase, MCCC2 (MCCB), by RT-PCR. A partial nucleotide sequence of 595 nucleotides of red seabream cDNA encoding 198 amino acids was obtained by RT-PCR with forward primer MCCC2-Fa and a reverse primer MCCC2Ra , which were derived from conserved MCCC2 sequences in human (AB029826) and Arabidopsis (AF059510). 5' RACE PCR using a gene-specific reverse primer, MCCC2Rb , and the adapter primer was also performed to recover a complete 5 ' sequence including the start codon, and the
compiled sequence of 1361 bp was obtained and deposited in the GenBank database (AB099519). The initiating methionine codon was found at nucleotides 48 to 50 , and the mRNA encoded 438 amino acid residues. The encoded amino acid sequence is highly homologous to the non-biotin-containing subunit of MCCases from human, mouse, insects, bacteria, and plant, and these are compared in Figure 3. This red seabream protein contained a complete domain characteristic of MCCC2, AccD for acetylCoA carboxylase $\beta$ subunit (COG0777) at 81 to 255 . In addition, other essential domains for MCCC2 were found: the $\alpha$ and $\beta$ subunits of carboxyltransferase domain (COG4799, 40-438); and the carboxyltransferase domain (pfam01039, 61-437). When the red seabream protein was compared with the non-biotin-containing subunits of PCCases from human and mouse, there were conserved amino acid residues but less homology to those for MCCC2. Overall homologies to human, fly, mosquito, Agrobacterium, and Arabidopsis were $80 \%, 69 \%, 68 \%, 72 \%$,

Figure 2. Comparison of amino acid sequences of $M C C C I$ between red seabream and other organisms. The deduced amino acid sequence derived from the coding sequence of the red seabream cDNA (AB097824) was compared with the MCCCI amino acid sequence of mouse (NP_076133), human (BAA99407), mosquito (EAA14349), fruit fly (NP_651896), nematode (NP_501777), and Arabidopsis (Q42523). The motif representing biotin binding is boxed. Genus and species abbreviations: red seabream (Pgm), human (Hsap), mouse (Mmus), mosquito (Agam), fruit fly (Dmel), nematode (Cele), and Arabidopsis (Athal). Dashes represent gaps inserted by the program to optimize the alignment. Domains identified are also shown as shaded areas: D-Ala D-Ala ligase biotinrequiring enzymes, carbamoyl-phosphate synthase (CPSase), phosphoribosylglycinamide synthetase (GARS). Asterisks and dots below the peptide sequence rows indicate completely matched and similar amino acid residues, respectively.
pfam00289, CPSase I chain





 Del Cele Atal


pfam02786.6, CPSase I D2,








$\nabla$ ATP binding site $\quad C_{2} b i n d i n g$ site
Pgm 179 RUAER














pfam02785.6, Biotin carb C






 *. * .. . **.**.*.***.

PgT 456 :LLHESSSGHPEHEANVLSELEQHYADLFIPKAPSGEITW-----QPA-LG---LVLQER--RHTQEFIQTSIDPES-P-FGSSSGNRSNIQFNRN


 DMEl 446 :THFTIDASHPEROLANVIGEIDEQFDITFPPIIISPQQVSQ----A-A-IA---LVLNEL----QAAFRNGNKDQD-P-FVATPNARLNYSLVRR



Pgm 542:MIIQ-------VGZKKVDWWYNKD-GSYT--MQIGEEVHHVTGDVEMEDGASFLHCS-VNGVKSRPKLVILDSTVHIFSTEGSAEVSVPVPKYLAA Maus $549: M I L R------S G K S D I V I A V I Y N R D-G S Y D--M Q I D N K S F R V L G D S S E T G C T Y L K S S-T N G V A R K S K F I L J D N I V H L F S M E G S I E V G I P V P K Y L S P ~$
 Agam 527:IKIK-------VGAMEHIVITEIDND-GSYS--VRVDDGEWMKVIVRRKPYANRFI工EINIEGHSCFNAVISGFNISIFDE-----------PRFLVI Dmel $528: Y D L K-------A N E K V Y S V A V K F D K E-D M Q--I Q V D N G W Q D A K V E R V Q D G S R I K I R A N I N S N V I T Y N A S I D G I S V S L F S E S G K D F E V G Q P K F L S A ~$ Cele 531:--D--GKDVS---IRFDSDSRLTVSYDCNSYETI-INDIESTIDGSFKFTIEANGRRWSIVKNLG-NSLM---VN-GVGQNEYEIPQ-IH-----ET Atal 556:NECEGIGSNLISLGVRYQEDGGYLIEE-GNDSPSLELR-VIRAGKCD--FRVEAAGLSMNVSLAAYLKDGYKHIHIWHGSEHHQFKQKVGIEFSEDEEG
pfam00364 biotin lipoyl $₹$ btotin binding. site








Figure 2.

| Pgm |  |
| :---: | :---: |
| Hsap |  |
| Agt | --MK--LTSSINRDP-AF 13 |
| Son |  |
| Agam | 1:TRELAKLHCRTDSPVRCFPRRRRTKKASSMLSKA-RSILS-LTR-VHHGSVANVLGV-RSVHISEANVLPTEVNRQSAEF 76 |
| Dmel | MIR-INW-LERSSSVIJRSQVRLTHVGDANVLHSEVDKQSAEY |

## AOCD











Agam 157: L N




 Agam 237:AR V1.



 Son 255:ALETARRAVSRLNHQKETT-_-_LRLSP-VK-_-_-_-_-_-_-_PPKFDISELYGIVGIDLKKPFDVKEVIARIVDDS 314 Agam 317:ALYLARQVVKNLNRPGSASYNE-LAGSSTATMMAREGITEGI-DPEPPQYPATDIYGIVGSNLTKTFDVREVIARIVDGS 394 Imel 282:ALYLARQIVSNLNLSATNSYNDQIMHSSQNNFQTATPPS-AVEF--P-RYTAEFLYGIVGPNLTKSEDVVREVIARTVDGS 357

Egm 344:KFDEFKAFYGDILVIGESRIFGYPVGIIGNGGVFSESAKKGIHFIELCCORNIDLIEIGNITGMMVGREYEAGGIAKDG 423 Hsap 343:RFTEFKAFYGDILVIGFARIFGYPVGIVGNNGVLFSESAKKGTHFVQLCCQRNIPLUELQNITGEMVGREYEAEGIAKDG 422 Agt 313:QEDEEKALYGTTLVCGEAHTHGYPVGIIANNGILESESALKGAHFIEICCQRGIPLVELQNITGEMVGKAYEAGGIAKDG 392 SOn 315:DFDEFKANYGATLVCGFARIHGYVVGIVANNGILESESAQKGAHFIEUCCQRKIPLLELQNITGHMVGKKYEHEGIAKHG 394 Agam 395:RFIEFKKFYGETIVCGYARLYGQLVGIVGNGGLESESALKGAHFIQLCAQKRIPL工FLQNITGEMVGRDAEAGGIAKNG 474 Imel 358:RFIEEKKLYGETLVCGEAKLYGHIVGIVGNNGVLFSESALKGAHFIQLCAQRKIPLVFLONITGEMVGRDAEANGIAKNG 437
 Hsap 423:AKMVAAVACAQVPKITLIIGGSYGAGNYGMCGRAYSPRFLYIWPNARISVMGGFQAANVLATITKDQRAREGKQFSSADE 502 Agt 393:AKLVIPAVASAKVPKFIVIIGGSFGAGNYGMCGRAYSPRELhMWPNARISVMGGEQAASVLAQLRRDGIEADCRRHWSKEGE 472 Son 395:AKMVTAVSCANVPKFIVIIGGSYGAGNYGMCGRAFEPTMMMMWPNARISVMGGEQAAGVLAIVRRDGI ARKGEEWSAEDE 474 Agam 475:AKMTPAVACANVPKLILIIGGSYGAGMYGMCGRAYSPRELYMWPNSRISVMGGSQAAGVLAQITEEQYRRTGREWIEEIG 554 Dmel 438:AKMVTAVACANVPKFTVIIGGSYGAGNYGMCGRAYSPRFLYMWPNSRISVMGGIQAANVMAQITEDQRKRAGKEFSEEEA 517
**.*.**..*.***

Hsap 503:AALKEPIIKKFEEEGNPYYSSARVWDDGIIDPADTRIVLGLSESAAINAPIEKIDFGIFRM Agt 473: EAFKOPIREKYEREGHPYYASARLWDDGIIDPKDTRLVLGIGLSAALNAPIEPTRFGIFRM Son 475:KAFKAPIIAQYDKEGHPYHASARLWDDGIIDPAOTRDVVGIALSAAINAPIEDTRFGVFRM Agam 555:NRIKAPIVQQEEAEGSPYYSTARLWDDGIIDPVDTRRVLGLSLQAAINHPVGETREGVFRM Dmel 518:QKLKAPIVEMFEAEGSPYYSTARLWDDGIIDPANTRQILGLSLKAALNNAGQEIKFGVFRM

Figure 3. Alignment of MCCC2 from red seabream and from other organisms. The deduced amino acid sequence derived from the partial coding sequence of the red seabream protein obtained in this study (AB099519) compared with the MCCC2 amino acid sequence of human (NP_071415), fruit fly (NP_652724), mosquito (EAA07889), and Arabidopsis (AF059511). MCCC2 from 2 bacteria, Shewanella oneidensis MR-1 (NP_717504) and Agrobac-
terium tumefaciens str (NP_533975), were also included for this comparison. AccD, the characteristic domain for acetyl-CoA carboxylase $\beta$ subunit (COG0777), is shaded. Asterisks and dots below the peptide sequence rows indicate completely matched and similar amino acid residues, respectively. Genus and species abbreviations are as in Figure 2, except Agt is Agrobacterium; Son, Shewanella.
and $61 \%$, respectively, yet overall homology was distinctly low ( $40 \%$ ) to the non-biotin-containing subunit of PCCase in human (NM_000532) and mouse (NM_025835), which is a biotin-requiring enzyme family closely related to MCCases.

## Cloning of cDNA Encoding PCase

A cDNA fragment was obtained by RT-PCR using forward primer PC-Fa and reverse primer PC-Ra, which was based on the zebrafish sequence for PCase (AF295372). It was subcloned, sequenced, and shown to be 880 nucleotides long. The 3 ' region was obtained by RT-PCR using a genespecific primer based on this sequence, $\mathrm{PC}-\mathrm{Fc}$, and the tag primer. RT-RCR was also performed using a forward primer based on the zebrafish PCase sequence (AF295372), $\mathrm{PC}-\mathrm{Fb}$, and a reverse primer designed from the seabream cDNA obtained above, PC-Rb. We thus identified a 3170nucleotides sequence encoding a polypeptide of 1029 amino acid residues highly homologous to zebrafish PCase, and the compiled sequence was deposited in the Genbank database (AB081832).

To obtain the complete 5 ' sequence for red seabream $P C, 5$, RACE was performed using a gene-specific primer based on the cDNA sequence, $\mathrm{PC}-\mathrm{Rc}$, and the adapter primer. We thus obtained the complete coding sequence of red seabream PCase and deposited it in the GenBank database (AB086371). This sequence was 4010 bp long with an open reading frame from 389 to 3928 encoding a polypeptide of 1179 amino acid residues. This encoded polypeptide was highly homologous to PCases from zebrafish (AAG37836), human (JC2460), mouse (A47255), chicken (AAM92771), and a nematode (NP_505977), as shown in Figure 4. All of the important conserved domains for PCases were also found in the red seabream protein: CPSase L chain (36-149), CPSase-L 02 (151-364), biotin carboxylase c domain (375-482), HMGL-like domain (569-733), pyravate carboxylase conserved domain (8611069), and biotin-binding domain (1109-1177). The bio-tin-binding motif (AMKME) from 1142 to 1146 with the biotin-binding site (lysine) at 1144, and D-Ala-D-Ala ligase (EC:6.3.2.4) domain were also found (151-326), as in MCCC1. The partial N-terminal sequence of a V8 proteinase fragment of the $121-\mathrm{kDa}$ biotin-containing protein (Table 2) was found from 864 to 873 including the cleaving site E at 863 (including white letters in black box: ENEIPGGQYTN, with the first $E$ being the cleavage site for V8 proteinase).

## Phylogenetic Trees for MCCase Subunits and PCase in Various Organisms

Figure 5 shows phylogenetic trees for proteins MCCC 1 , MCCC2, and PC. There were 4 distinct classes of MCCC 1 and MCCC 2 consistent with evolution of these organisms: i.e., insects, vertebrates, nematode, and plant. The red seabream proteins were close to those in mammals, and fit in well with vertebrate MCCC 1 and MCCC 2 . As expected, the plant orthologs were the most distant from any of the others presented here.

Red seabream PCase was most homologous to zebrafish PCase ( $89 \%$ ), highly homologous to mammalian ( $82 \%$ ) and avian (77\%) PCases, and more distant from a nematode PCase ( $67 \%$ ). There was a deviation between fish and bird PCases from their typical phylogeny, suggesting a novel functional evolution in birds.

## Expression Analysis of Genes Encoding PCase and Biotin-Containing and Non-Biotin-Containing MCCase Subunits

Figure 6 shows transcript abundance of the different genes using total RNA isolated from various organs of red seabream blotted onto a nylon membrane and probed successively with $M C C C 1, M C C C 2$, and $P C$. In all panel lanes represent total RNA from heart, liver, spleen, muscle, and ovary. Molecular markers and the total RNA from heart electrophoresed under the same conditions are also shown.

## Discussion

In the present study we isolated cDNAs encoding 2 important biotin-containing mitochondrial enzymes, MCCase and PCase, in a commercially important fish, red seabream (Pagrus major). For MCCase we determined the complete cDNA sequence for the biotin-containing subunit (MCCC1 or MCCA), which is essential for the catalytic role. The overall homologies of the biotin-containing subunits of MCCases were high to mammals ( $71 \%-73 \%$ ) and other lower animals $(56 \%-57 \%)$, but only $47 \%$ to plant, as expected. In addition, all of the hallmarks for MCCase including the biotin-binding site (K) and motif (AMKME) were present. However, the homology of red seabream MCCC1 was only $47 \%$ to mammalian PCC1, which is a paralogous gene to $M C C C 1$ with a very similar domain structure. These results justify our assignment of the red

$$
\text { CPSase-L Chain pfam } 00289.6
$$

Fgm Drer Hsap Mmus Gaal Cel







CPSase-L 02pfam 02786.6 ATP bindina site








OObindina site





 Biotin carboxylase $c$ domain











 *****

$$
\text { HMG-like domain pfam } 00682.6
$$














750: EASKL工IGALRDRFFDVFIHVHTHDIAGAGVAAMLACAEAGADVVDAVDSMAGMISQPSMEATVACAKGIKIDIGIAI EKVFDYSEYYEVARGLYAPFDCIATMKSG 752 :QASRLITEALRDRFPDIPIHVHIHDIAGAGVAAMLACAQAGADIVDVAVDSWAGMISQPSMGAIVACIKGIKDIGISIDKVFDYSEYYEVARGIYAPFDCIATMKSG 750 :TACIMLVSSLRDRFPDIPLHTHIHDISGAGVAAMLACAQACADVVDVAADSMSGMISQPSMCALVACIRGIPIDIEVPMERVFDYSEYWEGARGLYAAFDCIATMKSG 750 : AACIMLVSSLRDRFEDIFLHTHIHDISGAGVAAMLACAQAGADVVDNAVDSMSGMISQPSMGALVACIKGIPIDIEVPLERVFDYSEYWEGARGLYAAFDCIATMKSG 750 : AAARLLVSSIRDRFPDVPIHVHIHIIIAGAAIATLIAAAANADADVVDVAVDAMSGMISQPSMGALVACARGIPIDIGIAIERVFEYSEYWEGARGL YAAFDCIATMKSG 746:EAAKLIIGALRDKFPDIFIHVHIHDISGAGVAAMECAKAGADVVDAAVDSMSGMISQPSMMGAIVASLQGIKHDIGISILDISKYSAYWESTRQLYAPFECAITIMKSG

Pyruvate carboxylase conserved domain pfam 02436.6 OADA













Biotin hindina domain: pfam $00364 \quad \gamma$ biotin binding site







Figure 4. Comparison of amino acid sequences of PCases between red seabream and other organisms. The deduced amino acid sequence derived from the full coding sequence of the red seabream cDNA (AB097824) was compared with the PCases from zebrafish (AAG37836), human (JC2460), mouse (A47255), chicken (AAM92771), and a nematode (NP_505977). The motif representing biotin binding is boxed. Asterisks and dots below the peptide sequence rows indicate completely matched and similar amino acid residues, respectively. Genus and species abbreviations are as in Figure 2, except Ggal is chicken.
seabream cDNA obtained in the present study ( AB 100429 ) as MCCC1. The fish MCCC1 shown here was the most similar to mouse $M C C C 1$ in terms of both homology (73\%) and length of encoded polypeptide (717 amino acid residues). Because the mRNA was 4300 nucleotides long according to the Northern blot (Figure 6), and which is very close to that predicted from the nucleotide sequence obtained in the present study (AB100429), we presume that the sequence for red seabream $M C C C 1$ obtained in the present study is full length. This is about 200 nucleotides longer than that in human (Obata et al., 2001).

Such a long 3' UTR can often be found in fish gene transcripts, and it is sometimes difficult to obtain a fulllength sequence of the $3^{\prime}$ UTR from fish (Gray et al., 2001). We also obtained most of the coding sequence of $M C C C 2$ containing the start codon. Although 438 amino acid residues were encoded by the cDNA fragment here, the encoded polypeptide was highly homologous to MCCC2 in other organisms and especially to human ( $84 \%$ over 429 amino acids overlap out of 438 amino acids) and $67 \%$ to $71 \%$ homologous in other organisms, including the most distant species, Arabidopsis. Human MCCC2 encodes 563 amino acid residues (Obata et al., 2001). However, when compared with the paralogous PCC2, even from mammals, the homology was only $35 \%$ to $37 \%$ in a 127 amino acid overlap with red seabream MCCC2. In addition, the mRNA hybridized with this red seabream cDNA was about 2400 nucleotides long (Figure 6, B), which is close to the reported size of the full-length $M C C C 2 \mathrm{cDNA}$ in human of 2016 nucleotides (AB050049). The expression pattern was also exactly matched with that MCCC1 (Figure 6, A). These results strongly imply that the partial cDNA sequence obtained here corresponds to red seabream MCCC2.

PCase is composed of only one type of subunit, and the cDNA for the complete coding sequence of PCase in red seabream was obtained in the present study. In this

## A MCCC1 <br> 

## B MCCC2



## C PC



Figure 5. Phylogenetic analysis of MCCC1, MCCC2, and PC in divergent species of organisms. A phylogenetic tree was constructed using UPGMA method in the alignment tool (GENETYX-WIN, Version 6.1.0). Genus and species abbreviations are as in Figures 2-4. Divergence distances calculated by GENETYX-WIN are shown on individual branches. A: The tree derived from amino acid sequence alignments of 2 vertebrate MCCCl sequences from human (BAA99407) and mouse (NP_076133); 2 insects, fruit fly and mosquito; one lower animal, nematode; and one plant, Arabidopsis, and the red seabream MCCCl obtained in this study (Pgm) is shown. B: The tree derived from partial amino acid sequence alignments of the red seabream MCCC2 (Pgm) is shown. Phylogenetic relations to a paralogous gene, $P C C B$, are also shown. Hsap-PCCB and Mmus-PCCB represent $P C C B$ from human and mouse, respectively. $C$ : The tree derived from amino acid sequence alignments in Figure 4 including the red seabream protein ( Pgm ) is shown.


Figure 6. Northern blotting analysis of $M C C C 1, M C C C 2$, and $P C$ in red seabream organs. Total RNA isolated from various organs of red seabream was blotted and successively probed with ${ }^{32} \mathrm{P}$-labeled cDNAs for MCCC1 (A), MCCC2 (B), and PC (C) as isolated and identified in this study. In all panels, the ethidium-bromide-stained gel showing total RNA from heart (lane 1), liver (lane 2), spleen (lane 3 ), muscle (lane 4), and ovary (lane 5) is given. D: M indicates molecular markers (Hind III digest of $\lambda$ DNA) electrophoresed under the same conditions; $R$, ribosomal RNA from red seabream. Randomly primed ${ }^{32}$ P-labeled probe (specific activity, $1.0 \times 10^{6}$ cpm $\mu \mathrm{g}^{-1}$ ) was hybridized for each probe. The mRNAs for MCCC1, MCCC2, and PC in the blots are 4300, 2400, and 6500 nucleotides long, respectively.
encoded polypeptide, all of the hallmarks for PCase were present. PCase mRNA in red seabream was a single transcript of 6500 nucleotides according to the Northern blot (Figure 6, C). The cDNAs encoding PC were 3543 to 4000 nucleotides long in zebrafish (Yoder and Litman, 2000) and virtually identical length in red seabream. In addition, both zebrafish and red seabream sequences lack the canonical poly(A) signal (aataaa). In humans a very similar length (4111-nucleotide) PC was also reported (NM_000920), and this human sequence contains the canonical poly(A) signal and is thought to be full length. However, considering that the mRNA was significantly longer ( 6500 nucleotides) in the Northern blot (Figure 6, C) than that predicted from the cDNA sequence ( 4010 nucleotides) in red seabream, we anticipate that the compiled cDNA sequence for $P C$ may not be complete at its $3^{\prime}$ UTR.

Transcripts for both the biotin and non-biotin-containing subunits of MCCase were abundant in skeletal muscle and ovary, but only trace amounts were present in other organs (Figure 6A and B). In contrast, the more essential pyruvate carboxylase was expressed significantly in all organs except spleen. We do not know why the transcript levels for these nuclear-encoded, mitochondrial enzymes were so low in the spleen. PCase is one of the key enzymes to incorporate pyruvate into mitochondria, and its expression is more ubiquitous (being significantly expressed in liver
and highly in heart, muscle, and ovary) than that of MCCase subunits. In particular, $P C$ transcripts were more abundant than MCCase subunit transcripts in heart, and may play an essential role in metabolizing pyruvate in heart. Extremely strong expression of MCCase in muscle and ovary indicates very active metabolism of amino acids in these organs, including active protein synthesis and degradation (Obata et al., 2001). This might be related to a physiologic factor such as laying eggs, which was occuring in the red seabream we used Northern blot analysis (Jitrapakdee et al., 1996) has shown that in mouse, PC mRNA is not expressed in spleen or skeletal muscle, while it is higher in liver, and except for spleen, this expression pattern in mouse organs is different from that in red seabream. Indeed, little or no expression of these biotin-dependent enzyme transcripts takes place, suggesting that protein turnover in heart, liver, and spleen might not be important in red seabream. However, expression and levels of these biotin-dependent enzymes may be affected by environmental and physiologic factors such as nutrition in mouse (Goldberg and Chang, 1978) and human (Murayama et al., 1997), and such factors should be considered in future studies.

We have previously reported protein bands reacting with a streptavidin-conjugated alkaline phosphatase in different classes of organisms including plants, animals, and several fish species (Abe et al., 1992, 1996; Abe and Davies 1995), and we identified some of them as the biotin-containing subunit of MCCase in a plant, Pisum sativum (AB075695, AB075695), and in human (Obata et al., 2001). We confirmed that the $76-\mathrm{kDa}$ and $121-\mathrm{kDa}$ biotin-containing proteins are a subunit of MCCase and a biotincontaining enzyme, PCase (Table 2), while homology searches for the partial amino acid sequence of the $72-\mathrm{kDa}$ biotin-containing protein suggested that the protein is likely to be PCCase in red seabream (Figure 1, Table 2). Therefore, the protein bands at 56 kDa , which are found in abundant amounts but do not react with streptavidin, are likely to be the non-biotin-containing subunits of MCCase and PCCase (Oei and Robinson, 1985). However, the guanidine- HCl methods we used are not suitable for isolating these enzymes in active form, which would require a biotin gradient. This makes it very simple to isolate biotin-containing enzymes, for analyses such as protein sequencing. The detection method for these biotin-containing enzymes (using a streptavidin conjugate on transferred proteins) is also simple and sensitive, and may be useful for analyzing the amounts of these biotin-containing enzymes under different physiologic conditions in fish and other species.

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