Construction of a self-cloning system in the unicellular green alga Pseudochoricystis ellipsoidea

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Abstract

Background: Microalgae have received considerable interest as a source of biofuel production. The unicellular green alga Pseudochoricystis ellipsoidea (non-validated scientific name) strain Obi appears to be suitable for large-scale cultivation in outdoor open ponds for biodiesel production because it accumulates lipids to more than 30 % of dry cell weight under nitrogen-depleted conditions. It also grows rapidly under acidic conditions at which most protozoan grazers of microalgae may not be tolerant. The lipid productivity of this alga could be improved using genetic engineering techniques; however, genetically modified organisms are the subject of regulation by specific laws. Therefore, the aim of this study was to develop a self-cloning-based positive selection system for the breeding of P. ellipsoidea.

Results: In this study, uracil auxotrophic mutants were isolated after the mutagenesis of P. ellipsoidea using either ultraviolet light or a transcription activator-like effector nuclease (TALEN) system. The cDNA of the uridine monophosphate synthase gene (PeUMPS) of P. ellipsoidea was cloned downstream of the promoter of either a beta-tubulin gene (PeTUBULIN1) or the gene for the small subunit of ribulose 1,5-bisphosphate carboxylase/oxygenase (PeRBCS) to construct the pUT1 or pUT2 plasmid, respectively. These constructs were introduced into uracil auxotroph strains, and genetically complementary transformants were isolated successfully on minimal agar plates. Use of Noble agar as the solidifying agent was essential to avoid the development of false-positive colonies. It took more than 6 weeks for the formation of colonies of pUT1 transformants, whereas pUT2 transformants formed colonies in 2 weeks. Real-time PCR revealed that there were more PeUMPS transcripts in pUT2 transformants than in pUT1 transformants. Uracil synthesis (Ura+) transformants were also obtained using a gene cassette consisting solely of PeUMPS flanked by the PeRBCS promoter and terminator.

Conclusions: A self-cloning-based positive selection system for the genetic transformation of P. ellipsoidea was developed. Self-cloned P. ellipsoidea strains will require less-stringent containment measures for large-scale outdoor cultivation.

Keywords: Pseudochoricystis ellipsoidea, Green algae, Microalgae, Self-cloning, Uracil auxotroph, Genetic transformation, Molecular breeding.

Background

Microalgae have received considerable interest as a source of biofuel production. In particular, the biodiesel produced by microalgal biomass may be a viable alternative to petroleum-based and land-plant-based fuels because microalgae have several advantages over terrestrial plants for biofuel production, such as low land requirements and high oil yields [1–3]. Although the production of crude oil from algal biomass has already been achieved in various pilot-scale facilities, the current production costs of algal oil are not competitive with those of fossil oil [4]. The processes used to produce crude algal oil include cultivation and recovery of algal biomass, oil extraction from the algal biomass, and conversion of fatty acid moieties of the oil to liquid biofuels using either methyl esterification or hydrogenation. Currently, efforts are underway to reduce the capital and operating costs of these processes [5, 6]. However, significant cost reduction could also be achieved via algae breeding, particularly using genetic engineering techniques [7, 8].
Genetically modified algal strains are subject to laws concerning biosafety and also raise ambiguous public concerns; therefore, molecular breeding using techniques based on self-cloning and/or natural occurrence (cloning DNA from a donor into a recipient, between which the natural exchange of DNA is possible) was proposed as a desirable method to construct recombinant organisms for deliberate release into the environment. In Japan, recombinant microorganisms including microalgae, obtained using self-cloning or natural occurrence are exempt from the restrictions imposed by the Cartagena domestic law [9, 10].

In plants, concepts named cisgenesis and intragenesis, which are analogous to those of self-cloning and natural occurrence, have been proposed. In cisgenesis and intragenesis, fragments of non-recombinant and recombinant DNA, respectively, from a sexually compatible donor organism is introduced into a recipient. Since both cases involve the exchange of DNA between interbreeding groups, the regulatory measures imposed on cisgenesis and intragenesis could be similar to those for conventional breeding. The regulation of intragenic and cisgenic organisms is currently under discussion in many countries, including in the European Union and the United States, and the regulation of these organisms might be relaxed soon [11]. Amid such trends, the development of self-cloning technology is important for algae breeding using outdoor open pond cultivation, which is currently considered the most viable option for the large-scale cultivation of microalgae [12].

Pseudochoricystis ellipsoidea (non-validated scientific name) strain Obi is a unicellular green alga classified within the family of Trebouxiaceae, which accumulates lipids to an amount of more than 30 % of the dry cell weight under nitrogen-depleted conditions [13]. During nitrogen depletion, the amount of neutral lipids increases, whereas chloroplast-specific glycolipids decrease [14]. A peculiar characteristic of this alga is its ability to grow rapidly under acidic conditions below pH 3.5 without reducing the lipid productivity (Kurano et al., in preparation). Since most protozoan grazers of microalgae might not tolerate that pH, P. ellipsoidea might be suitable for large-scale cultivation in outdoor open ponds for biodiesel production. Thus, we developed a method for the genetic transformation of this alga [15] and are using this method to genetically improve the oil productivity of P. ellipsoidea.

We were also interested in developing a self-cloning-based positive selection system for the breeding of P. ellipsoidea for two reasons. First, as discussed above, self-cloning is subject to less-stringent regulation [16, 17], and a consensus regarding the deregulation of the deliberate release of self-cloned microalgae is likely to be built among international experts in the near future. If this occurs, a climate could be developed to foster the cultivation of self-cloned microalgae in outdoor open ponds. Second, genetic engineering includes the controlled over-expression and the targeted inactivation of genes, which both require the use of selectable marker genes to acquire desired clones. Since only one marker is currently available in P. ellipsoidea (G418 resistance), auxotrophic markers will serve as additional selectable markers.

In the current study, we report (i) the isolation of uracil-requiring mutants of P. ellipsoidea that are defective in uridine monophosphate synthetase (UMPS), (ii) the synthesis of the full-length cDNA for UMPS, and (iii) the complementation of P. ellipsoidea uracil auxotrophs by introducing UMPS cDNA. We encountered several problems that were resolved. This information will be useful for individuals interested in constructing self-cloning systems in other microalgae.

Results and discussion
Isolation of uracil auxotrophs
During the ultraviolet light (UV)-mediated mutagenesis of P. ellipsoidea cells, the survival of UV-irradiated cells decreased over UV-C irradiation time and reached 0.1 % after 4 min (see Additional file 1). The mutated cells were screened for the uracil-requiring (Ura+) phenotype on MA5 minimal agar plates containing both 5-fluoroorotic acid (5-FOA) and uracil. Thirty-six 5-FOA-resistant colonies were isolated after a UV-C dose yielded a survival rate of ~50 %, and five of these were uracil auxotrophs. Using genomic DNAs isolated from two of the five uracil auxotrophs named strains M1 and M2, DNA fragments encompassing the coding region (CDR) of the gene for UMPS (PeUMPS) were PCR-amplified, and sequenced. Eukaryotic UMPS is a bifunctional enzyme that consists of orotate phosphoribosyltransferase (OPRTase) and orotidine-5′-phosphate decarboxylase (ODCase), which catalyze the last two steps of the de novo UMP biosynthetic pathway [18]. The DNA sequencing analysis showed that a 5-base insertion and a 27-base deletion were present in PeUMPS in strains M1 and M2, respectively, both in the OPRTase domain. The spontaneous reverse version frequency to Ura+ was 7.1 × 10−6 for strain M1, and less than 10−7 for strain M2. Because of the instability of the M1 mutation, strain M1 was not characterized further. Whole-genome sequencing of strain M2 covering 99.4 % of the whole genomic sequence of P. ellipsoidea strain Obi (Harayama et al., in preparation) revealed the presence of a one-base substitution in an intron of a gene of unknown function in addition to the PeUMPS mutation described above.

Two uracil auxotrophs named strains M3 and M4 were also isolated from a P. ellipsoidea strain Obi derivative carrying two DNA fragments encoding transcription activator-like effector nuclease (TALEN) right and left
arms that were designed for the mutagenesis of PeUMPS. The M3 and M4 strains were identified after selecting 5-FOA-resistant colonies grown on MA5 minimal agar plates containing uracil and 5-FOA. The M3 and M4 strains had a 29-base insertion and a 192-base deletion, respectively, in the ODCase domain of PeUMPS. However, these mutations were outside the TALEN-target sequence, suggesting that they arose independently from TALEN activity. The frequency of reversion of the two mutants to Ura+ was less than $10^{-7}$.

The M3 and M4 strains that carry the TALEN right and left sequences would not be appropriate hosts for self-cloning. Nevertheless, we used these strains together with strain M2 in following complementation analyses because the mutation loci in the M3 and M4 strains were within the OPRTase domain which was different from the mutation locus within the ODCase domain in strain M2.

**Cloning and functional analysis of PeUMPS cDNA**

A 1.5-kb PeUMPS cDNA was cloned into pUC119 to construct pUC_cPeUMPS (Fig. 1). The cloned PeUMPS cDNA sequence was 100% identical to that predicted from the genomic sequence of *P. ellipsoidea* strain Obi. The pTV-cPeUMPS plasmid (Fig. 1) was constructed to express PeUMPS cDNA in two uracil auxotrophic mutants of *Escherichia coli*; JD24489 (an OPRTase mutant) and JW1273 (an ODCase mutant). Both *E. coli* mutants did not grow on glucose M9 minimal medium, whereas clones that possessed pTV-cPeUMPS grew efficiently, suggesting that the PeUMPS cDNA was expressed functionally in *E. coli*.

**Genetic complementation of uracil auxotrophic *P. ellipsoidea* mutants using PeUMPS cDNA**

Plasmid pUT1 containing PeUMPS cDNA flanked by the promoter of *PeTUBULIN1* and the terminator of an actin gene (*PeACTIN1*) (Fig. 1) was introduced into the uracil auxotrophic strain M4, and prototrophic transformants were selected on MA5 minimal medium solidified using Bacto agar (BD Difco). However, prototrophic transformants could not be selected because of the background growth of strain M4 on MA5 minimal agar plates even when M4 cells grown in MA5 medium supplemented with uracil were washed extensively before plating (data not shown). To reduce the background cell growth, both pG418-T1A [15] (Fig. 1) and pUT1 were co-introduced into strain M4, and the resulting G418-resistant transformants were selected on MA5 minimal agar plates.

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**Fig. 1** Construction of the plasmids used in this study. The genes and loci are abbreviated as follows: cPeUMPS, PeUMPS cDNA; lacP, the promoter and operator regions of the *E. coli* beta-galactosidase gene (*lacZ*); nptII, the gene for neomycin phosphotransferase II to confer G418 resistance; *TUBP*, the promoter region of *PeTUBULIN1*; *ACTT*, the terminator region of *PeACT1*; *RBCS*, the promoter region of *PeRBCS*; *RBCST*, the terminator region of *PeRBCS*. **Kasai et al. Biotechnology for Biofuels** (2015) 8:94
resistant transformants were screened on MA5 minimal agar plates containing G418 and uracil.

Six G418-resistant colonies were obtained, and the existence of pUT1 sequences in the transformants grown on MA5 minimal agar plates containing uracil was examined using PCR with two forward primers (one that hybridized with the 5'-ends of the PeTUBULIN1 promoter, and another that hybridized with the 5'-ends of the PeUMPS cDNA) and three reverse primers that hybridized the 3'-end of the PeUMPS cDNA, the middle part of the PeACTIN1 terminator, and the 3'-end of the PeACTIN1 terminator (Fig. 2). In the genomes of three out of the six transformants (named strains M4-1A, M4-1B, and M4-1C), the DNA region covering the PeTUBULIN1 promoter, PeUMPS cDNA, and the PeACTIN1 terminator was integrated. Conversely, the genome of the transformant named strain M4-1D contained a region covering the PeTUBULIN1 promoter, PeUMPS cDNA, and a 0.4-kb 5'-end of the PeACTIN1 terminator (Fig. 2).

The PCR products amplified from the M4-1C genome using the primers Ptub_F and Tact_R were a mixture of four or more DNA fragments (Fig. 2b, lane 2_C), suggesting that the DNA region, the PeTUBULIN1 promoter–PeUMPS cDNA–the PeACTIN1 terminator, integrated in the genome of the transgenic M4-1C strain was unstable when grown in the presence of uracil.

When these four PeUMPS-cDNA-positive transgenic strains were streaked onto uracil-free MA5 agar plates, colonies developed only after incubation for 6–8 weeks at 25 °C. Since strain M4 formed no colony on uracil-free MA5 agar plates after 8 weeks, the introduced PeUMPS cDNA complemented the PeUMPS mutation in strain M4. But the effect was very weak. To shorten the time required for colony formation, MA5 minimal agar plates supplemented with precursors of the UMP biosynthetic pathway (L-glutamate, orotate, and phosphoribosyl pyrophosphate) were prepared, and the growth of the four transgenic strains on the plates was examined. The transgenic strains M4-1C and M4-1D formed colonies in 3 weeks on MA5 minimal agar plates containing orotate; however, the growth improvement by orotate was not significant in the M4-1A and M4-1B strains, which still formed colonies in only 8 weeks. Neither phosphoribosyl pyrophosphate nor L-glutamate improved the growth of any transgenic strains.

To evaluate the correlation between the expression of PeUMPS cDNA and the growth speed, the expression of PeUMPS cDNA in the four transgenic strains was analyzed using real-time PCR and normalized against 18S rRNA levels (Fig. 3a). The PCR primers Umps_QRT_F and Umps_QRT_R were used to detect both the expression of transgenic PeUMPS cDNA and the expression of
defective PeUMPS in the chromosome of strain M4. Strain M4-1C, which grew fastest (its colonies were visible in 6 weeks), exhibited a 1.5-fold higher expression than did strain M4. It suggested that the expression of the transgenic PeUMPS cDNA in strain M4-1C was 1.5 –1-fold of the expression of the original PeUMPS. Similarly, the expression of the transgenic PeUMPS cDNA in strain M4-1D was approximately one quarter of the expression of the original PeUMPS. Strain M4-1D grew faster than strains M4-1A and M4-1B in the presence of orotate. In contrast, no transgene expression was detectable in strains M4-1A and M4-1B, which grew very slowly even in the presence of orotate. These results suggest that the slow growth of the transgenic strains on MA5 minimal agar plates was due to the low expression of the transgenic PeUMPS cDNA.

We next attempted to place the PeUMPS cDNA under the control of a stronger promoter. Whole transcriptome shotgun sequencing (RNA-seq) was applied to P. ellipsoidea strain Obi, which revealed that mRNA of the gene for ribulose-1,5-bisphosphate carboxylase/oxygenase small subunit (PeRBCS) was the highest among all nuclear genes and was almost 20 times higher than was that of PeTUBLIN1 (Ide et al., unpublished data). Thus, the PeUMPS cDNA fragment was inserted downstream of the PeRBCS first intron in plasmid pHA1101 to construct plasmid pUT2 (Fig. 1). The pUT2 and pG418-T1A plasmids were then used to co-transform strain M4 using particle bombardment. Among the 106 G418-resistant colonies that developed, the PeUMPS cDNA sequence was identified in eight colonies using PCR. To determine the pUT2 sequences that integrated in the genome of these eight transgenic strains, PCR was performed using three primer sets to detect the 5′-end of the PeRBCS promoter, the middle part of the PeUMPS cDNA, and the 3′-end of the PeRBCS terminator, respectively (Fig. 4). All three regions were detected in five transgenic strains (M4-2A, M4-2B, M4-2C, M4-2D, and M4-2E), whereas the remaining three transgenic strains contained the PeUMPS cDNA and PeRBCS terminator regions, but not the PeRBCS promoter region (data not shown). The PCR products amplified from the genome of strain M4-2A contained multiple fragments, suggesting that a population of strain M4-2A harbored partial deletions in the integrated expression cassette of the PeRBCS promoter—PeUMPS cDNA—PeRBCS terminator. Thus, the expression cassette in strain M4-2A was unstable to a certain extent in strain M4-2A. Using genomic DNA isolated from strain M4-2C, fragments 2 and 4 were not amplified efficiently. We interpreted this result as suggesting that strain M4-2C did not carry the full-length expression cassette, but instead carries two or more incomplete expression cassettes: at least one covering the PeRBCS promoter—PeUMPS cDNA region, and one covering the 3′-end PeUMPS cDNA—the PeRBCS terminator region.

Strains M4-2A, M4-2B, M4-2D, and M4-2E grew on MA5 minimal agar plates and formed colonies within 7 days, similar to the wild-type P. ellipsoidea strain Obi. However, strain M4-2C grew on MA5 minimal agar plates and formed colonies after only 6 weeks. Strain M4-2C grew slower than other transgenic strains probably because the strain only carried incomplete forms of the gene expression cassette, as described above.

The expression of PeUMPS cDNA in strains M4-2A, M4-2B, and M4-2E was determined using real-time PCR and normalized to 18S rRNA (Fig. 3b). The expression of the transgene was three- to seven-fold higher than was the expression of the chromosomal copy of PeUMPS.

Southern blotting of genomic DNAs extracted from strains M4-1A, M4-1B, M4-1C, M4-1D, M4-2A, M4-2B, M4-2D, and M4-2E was performed, and the results are shown in Additional file 2. Southern blotting analysis of
genomic DNAs digested with EcoRI should yield two DNA fragments of 8 and 3 kb (see Additional file 2) originated from the original \textit{UMPS} gene, that were hybridized to the \textit{UMPS} cDNA probe. These two bands were detected in all DNA samples, including that from strain M4. In each of the transgenic strains, additional band(s) were detected, and the number of the additional bands indicated the copy number of \textit{UMPS} cDNAs that integrated into the genome. Thus, the copy numbers were expected to be 2, 4, 4, and 4 for strains M4-1A, M4-1B, M4-1C, and M4-1D, respectively, and 3, 1, 4, and 7 for strains M4-1A, M4-1B, M4-1D, and M4-1E, respectively, based on the Southern blots obtained using EcoRI-digested genomic DNA. Similar results were obtained from Southern blotting performed using XbaI-digested genomic DNA (see Additional file 2). Since strain M4-2B, which carried a single copy of \textit{PeUMPS} cDNA under the control of the \textit{PeRBCS} promoter, expressed the transgene at a much higher than did the transgenic strains carrying multiple copies of \textit{PeUMPS} cDNA under the control of the \textit{PeTUBULIN1} promoter, we concluded that the transgene was expressed at significantly higher levels from the \textit{PeRBCS} promoter compared with the \textit{PeTUBULIN1} promoter.

Analysis of the growth of the transgenic strains in MA5 minimal liquid medium showed that transgenic strains containing the pUT1 sequence started to grow with a lag phase of 2–8 days and slower growth rates compared with the wild-type strain (\textit{P. ellipsoidea} strain Obi) and strain M4 in the presence of 1 mM uracil (Fig. 5a). In contrast, two of the transgenic strains carrying the pUT2 sequence (4 M-2B and 4 M-2D) grew in MA5 minimal liquid culture at growth rates equivalent to wild-type \textit{P. ellipsoidea} strain Obi, whereas the other two transgenic strains (4 M-2A and 4 M-2E) grew slower than the wild-type strain, despite high levels of transgene expression (Fig. 5b). Although the reason for the slower growth of strains 4 M-2A and 4 M-2E is unclear, it is possible that the insertion of a copy of \textit{PeUMPS} cDNA or the G428-resistance gene into the genomes of these strains disrupted gene(s) that are essential to the rapid growth of \textit{P. ellipsoidea}.

In the studies described above, the \textit{PeUMPS} cDNA was introduced into strain M4 using a co-transformation technique with pG418-T1A plasmid to first select G418-resistant transformants. However, since we demonstrated the practical utility of pUT2 plasmid for complementing the defect in \textit{PeLUMPS}, we next attempted to select Ura$^+$ transformants of the uracil auxotroph strains M2, M3, and M4 using this plasmid. As described above, the background growth of these auxotrophs on MA5 minimal agar plates hampered the effective selection of Ura$^+$ transformants. Nevertheless, we found that the use of Noble agar (BD Difco) instead of Bacto agar eliminated background growth of the uracil auxotrophic
strains on MA5 minimal agar plates completely. Thus, plasmid pUT2 was introduced in strains M2, M3, and M4, and Ura⁺ transformants were selected directly on MA5 minimal agar plates solidified using Noble agar (BD Difco). Colonies of Ura⁺ transformants of strains M2, M3, and M4 appeared in 10 days, whereas no colony appeared on the agar plates containing bombarded microcarriers without DNA coating. The transformation rates differed among the uracil auxotrophic strains (Table 1); that of strain M4 was significantly lower than were those of strains M2 and M3.

To establish a method for genetic transformation without introducing foreign nucleic acid sequences (self-cloning), a gene cassette consisting of the PeRBCS promoter—PeUMPS cDNA—the PeRBCS terminator was excised from the pUT2 plasmid using restriction-enzyme digestion, and purified after gel electrophoresis. The linearized gene cassette DNA was then introduced into strain M2 using particle bombardment, and Ura⁺ transformants were selected on MA5. Ura⁺ colonies appeared in 10 days at a frequency of 57.3 ± 6.9 transformants/μg DNA. Although this value was half the transformation frequency obtained using pUT2, it was sufficient for the practical use of the gene cassette for the genetic transformation of Ura⁻ strains of _P. ellipsoidea_.

**Conclusions**

In this study, a self-cloning-based positive selection system was developed in three distinct steps to obtain transgenic derivatives of _P. ellipsoidea_ strain Obi. In the first step, four uracil auxotrophic mutants named strains M1 to M4 were isolated by selecting 5-FOA-resistant derivatives. Strain M2, which was isolated after weak UV mutagenesis, was most convenient as a recipient in genetic transformation because it contained only two mutations: a 27-base deletion in _PeUMPS_, and a base substitution in an intron of a gene of unknown function. In the second step, UMPS cDNA was cloned and sandwiched between a promoter sequence and a terminator sequence. The _PeRBCS_ promoter was suitable for the transgenic expression of _PeUMPS_, whereas the _PeTUBULIN1_ promoter did not support sufficient transgenic expression of _PeUMPS_. In the third step, the preparation of minimal agar plates was critical for the efficient screening of Ura⁺ transformants. The information obtained in this study will be useful for researchers interested in constructing self-cloning systems in other microalgae.

**Methods**

**Algal strain and culture conditions**

The algal and bacterial strains used in this study are listed in Table 2. _P. ellipsoidea_ strain Obi (wild-type) and its uracil auxotrophic mutants were cultured in a 120-ml test tube containing 50 ml of MA5 medium [15] or MA5 medium supplemented with 1 mM uracil under continuous illumination with daylight fluorescent tubes (40 W FL40S • FR • P, Panasonic) at 100 μmol photons m⁻² s⁻¹ in a plant-growth chamber (type #CLE-303, Tomy). The media were bubbled with 1 % (v/v) CO₂ at 25 °C. For agar-plate cultivation, the media were solidified using 1.5 % agar (Bacto agar or Noble agar, BD Difco), and the resulting agar plates were incubated in the plant-growth chamber.

**Isolation of uracil auxotroph mutants**

Two methods were used to isolate uracil-requiring mutants: UV mutagenesis and TALEN mutagenesis. For UV
mutagenesis, *P. ellipsoidea* strain Obi cells cultured at an OD_{750} of 2.2 were collected by centrifugation at 5000×g for 5 min, and then re-suspended in 50 mM potassium phosphate buffer (pH 7.4) to an OD_{750} of 1 (approximately 2 × 10^7 cells ml⁻¹). Two milliliters of the cell suspension was transferred into an empty Petri dish, stirred using a stirring bar, and irradiated with an ultraviolet (UV) lamp (15 W GL15, Mitsubishi Electric) for 1–15 min at a distance of 6 cm. Subsequently, the irradiated cells were stored in the dark for 18 h at 4 °C to prevent photoreactivation, and then each cell suspension was added to 25 ml of MA5 medium containing 1 mM Na_2CO_3 and 1 mM uracil and incubated photosynthetically in the plant-growth chamber until it reached an OD_{750} of more than 5 to segregate mutated genes. Cells in these cultures were diluted serially, and then spread on MA5 minimal agar plates containing 1 mM uracil and 4.3 mM 5-FOA (Wako Pure Chemical Industries). The agar plates were incubated for 2 weeks in the plant-growth chamber, and the colonies that grew on the agar plates were picked using sterile toothpicks and streaked free DNase Set (Qiagen) according to the manufacturer's instructions. The RNA concentration was determined from the absorbance at 260 nm, and the RNA integrity was examined using agarose gel electrophoresis. First-strand cDNA was synthesized using a Transcriptor High Fidelity cDNA Synthesis kit (Roche) with anchored-oligo(dT)18 primers according to the manufacturer's instructions. The RNA concentration was determined from the absorbance at 260 nm, and the RNA integrity was examined using agarose gel electrophoresis. First-strand cDNA was synthesized using a Transcriptor High Fidelity cDNA Synthesis kit (Roche) with anchored-oligo(dT)18 primers according to the manufacturer's instructions. The RNA concentration was determined from the absorbance at 260 nm, and the RNA integrity was examined using agarose gel electrophoresis. First-strand cDNA was synthesized using a Transcriptor High Fidelity cDNA Synthesis kit (Roche) with anchored-oligo(dT)18 primers according to the manufacturer's instructions. The RNA concentration was determined from the absorbance at 260 nm, and the RNA integrity was examined using agarose gel electrophoresis.

**Cloning of the cDNA encoding UMPS**

Standard molecular procedures were performed as described by Sambrook et al. [22]. Total RNA was extracted from *P. ellipsoidea* strain Obi cells grown to an OD_{750} of 1.2 using an RNeasy Mini kit (Qiagen), and any remaining genomic DNA was digested using an RNase-free DNase Set (Qiagen) according to the manufacturer's instructions. The RNA concentration was determined from the absorbance at 260 nm, and the RNA integrity was examined using agarose gel electrophoresis. First-strand cDNA was synthesized using a Transcriptor High Fidelity cDNA Synthesis kit (Roche) with anchored-oligo(dT)18 primers according to the manufacturer's instructions. A fragment of PeUMPS cDNA was amplified from a cDNA library using PCR with the primers Umps_{F} (5′-CGGGATCCGGCTGCACTTTATTTG TGC-3′) and Umps_{R} (5′-GGAATTCCCAACCTTCATTGAGAGGCAAT-3′) and the entire sequence in the two mutants was determined using dideoxy chain termination via a commercial service provided by Macrogen Japan Corp.

| Strains | Genotype | Source or reference |
|---------|----------|---------------------|
| Obi     | Wild-type | [13]                |
| M2      | Carry a 27-base deletion within the OPRTase domain of UMPS | This study |
| M3      | Carry a 29-base insertion within the ODCase domain of UMPS | This study |
| M4      | Carry a 192-base deletion within the ODCase domain of UMPS | This study |
| TAL-1   | Carry two chromosomally integrated DNA fragments encoding TALEN left and right arms | This study |
| E. coli |          |                     |
| DH5a    | F′ endA1, glnV44, thi-1, recA1, relA, gyrA96, deoR, nupG, Δ(88dlacZΔM15), Δ(lacZYA-argF)U169, hsdR17 (K2- mK) | TOYOBO |
| JD24489 | W3110 (typeA) lacIq, lacZΔM15, galK2, galK22, ΔpyrEkan | National Institute of Genetics |
| JW1273  | Δ(araD-araB)567, ΔlacZΔM15 (rmb-3), ΔpyrF789:kan, rph-1, Δrha-dhaB568, hsdR514 (K2- mK) | National Institute of Genetics |
cDNA was verified using double-stranded sequencing as described above. To construct a PeUMPS cDNA expression vector in E. coli, PCR was performed using the primers Umps_CDRF (5′-CACCCATGGCAACTTCAACTCCCTCAGT-3′; an NcoI site is underlined) and Umps_CDRR′ (5′-CCAAGCTTCTATGCGGCTTCAT-3′; a HindIII site is underlined). The amplified fragments were then digested using NcoI and HindIII and cloned into the Ncol and HindIII sites of pTV118 (TaKaRa) to create pTV_cPeUMPS. To confirm the functionality of the cloned PeUMPS cDNA, pTV_cPeUMPS was introduced into the uracil auxotrophic E. coli strains JYD4489 [W3110 (type A), lacF, lacZΔM15, galK2, galK22, ΔpyrE::kan] and JW1273 [Δ araD Δ araB], ΔlacZ Δ787::c: Δ araB-3, Δ pyrF789::kan, rph-1, Δ (rhaD- rhaB)], which were purchased from the National Institute of Genetics of Japan (Mishima). The growth of the E. coli transformants was examined on glucose M9 minimal medium [22].

Construction of the PeUMPS cDNA vectors for expression in P. ellipsoidea

The pUT1 plasmid (NCBI/DDBJ accession number LC016841) carrying the PeUMPS cDNA flanked by the PeTUBULIN1 promoter and the PeACTIN1 terminator was constructed in two steps (see Additional file 3) using PrimeSTAR MX DNA polymerase (Takara) as described previously [24]. In the first step, three DNA fragments were amplified separately using PCR. Specifically, a 0.9-kb promoter region of P. ellipsoidea strain Obi was amplified using primers Ptub_F (5′-CCAAGCTTCTATGCGGCTTCAT-3′; an adapter sequence for the second-step PCR is underlined) and Ptub_R_Umps5′ (5′-GAGTTGAAGTGCAGGTGGACCTGGCCTCATA-3′), a HindIII site is underlined) with pG418-T1A DNA [15] as the template. In the second step, the three fragments amplified in the first step were assembled into a single fragment using Ncol and HindIII and cloned into the Ncol and HindIII sites of pTV118 (TaKaRa) to create pTV_cPeUMPS. To confirm the functionality of the cloned PeUMPS cDNA, pTV_cPeUMPS was introduced into the uracil auxotrophic E. coli strains JYD4489 [W3110 (type A), lacF, lacZΔM15, galK2, galK22, ΔpyrE::kan] and JW1273 [Δ araD Δ araB], ΔlacZ Δ787::c: Δ araB-3, Δ pyrF789::kan, rph-1, Δ (rhaD- rhaB)], which were purchased from the National Institute of Genetics of Japan (Mishima). The growth of the E. coli transformants was examined on glucose M9 minimal medium [22].

Table 3 List of PCR primers used in this study

| Primer name | Sequence (5′ to 3′) | Target |
|-------------|---------------------|--------|
| Umps_F      | CCACAAAACCCATTGCCCTCAACA | 5′-end of UMP5 |
| Umps_R      | TCCGTTGCTCTGCAAGTCTTT | 3′-end of UMP5 |
| Umps_cDNAF  | CCAAGGTTCGATTGAATTGCCACACATCAGT | 5′-end of UMP5 cDNA |
| Umps_cDNAR  | GGACAGCTTCTATGCGGCTTCAT-3′ | 3′-end of UMP5 cDNA |
| Umps_CDRF  | CACCCATGGCAACTTCAACTCCCTCAGT | 5′-end of UMP5 promoter |
| Umps_CDRR  | CACCCATGGCAACTTCAACTCCCTCAGT | 5′-end of UMP5 promoter |
| Ptub_F      | CCAAGCTTCTATGCGGCTTCAT-3′ | TUBULIN1 promoter |
| Ptub_R_Umps5′ | GAGTTGAAGTGCAGGTGGACCTGGCCTCATA-3′ | 5′-end of UMP5 cDNA and 3′-end of TUBULIN promoter |
| Umps3′_Tact_F | GCCACGCTTCTATGCGGCTTCAT-3′ | 3′-end of UMP5 cDNA and 3′-end of ACT1 terminator |
| Tact_R      | TGGCTACGGGTGTTGAGTGGACACTGGAC | 3′-end of ACT1 terminator |
| Tact_R2     | CCCAGCGTCGCTACGCTATCA | A middle part of ACT1 terminator |
| Prbc_F      | CTCATGCGGCAACTTCAACTCCCTCAGT | 5′-end of RBCS promoter |
| Trbc_F      | GGCATGGATCTATGCGGCTTCAT-3′ | 3′-end of RBCS terminator |
| Ptub3′_Umps_CDRF | GGGAAGTCGTTCAAGATGGCAACTTCAACTCCCTCAGG | 3′-end of UMP5 cDNA |
| Umps_CDRR_Tact5′ | TGGCTACGGGTGTTGAGTGGACACTGGAC | 5′-end of UMP5 terminator and 3′-end of UMP5 cDNA |
| G418F       | GATGGCAACTTCAACTCCCTCAGT | nptII |
| G418R       | GCCATGGATCTATGCGGCTTCAT-3′ | nptII |
| Umps_QRT_F  | TCTTCCAATGCGGATGATGG | Position 908–930 of UMP5 cDNA |
| Umps_QRT_R  | GTTACGGGATGATGG | Position 993–1012 of UMP5 cDNA |
| 18S rRNA_QRT_F | GCCCATGGATCTATGCGGCTTCAT-3′ | 5′-end of ACT1 terminator and 3′-end of UMP5 cDNA |
| 18S rRNA_QRT_R | GCCCATGGATCTATGCGGCTTCAT-3′ | 5′-end of ACT1 terminator and 3′-end of UMP5 cDNA |

The functionality of the cloned PeUMPS cDNA was confirmed by examining the growth of E. coli transformants on glucose M9 minimal medium [22].
as the templates and the primers Ptub_F and Tact_R. The resulting DNA fragment was purified using a PCR purification kit (Qiagen), digested using HindIII and KpnI, and cloned into the HindIII and KpnI sites of pBluescript II sk (+).

Next, the PelUMPS cDNA was placed between two DNA fragments: one containing the promoter—5’ untranslated region—first intron region of PeRBCS and the other containing the 3’ untranslated region—terminator sequence of PeRBCS. Briefly, the 1.5-kb PelUMPS cDNA was amplified using PCR with the primers Umps_CDRF and Umps_CDRR with pUC-cPelUMPS DNA as the template. The PCR product was digested using NcoI and HindIII, and then cloned into the NcoI and HindIII sites of pHA1011 (NCBI/DDBJ accession number LC016840) between the promoter and terminator regions of PeRBCS to construct pUT2 (NCBI/DDBJ accession number LC016842).

Transformation of P. ellipsoidea cells

Nuclear transformation was performed using particle bombardment as described previously [15], with slight modifications. Uracil auxotrophic strains were grown to an OD_{750} of 0.5 in MA5 medium supplemented with 1 mM uracil, and 1 ml of the culture was spread on an MA5 minimal agar plate containing 1 mM uracil. After drying the surface of the plate, samples were incubated at 25 °C for 3 days under 12-h light/dark cycles at a light intensity of 50 μmol m^{-2} s^{-1}. Then, the plates were placed under the stopping screen of a Biolistic PDS-1000/He particle delivery system (Bio-Rad) at a distance of 6 cm. DNA-coated gold particles of either 0.6 μm (Bio-Rad) or 0.3 μm (Recentec) diameter were prepared according to the instructions given by Bio-Rad (technical note # 2453), and the cells on MA5 minimal agar plates were bombarded twice at helium pressures of 1100 or 1550 psi, respectively, in vacuo with 125 mm Hg. After bombardment, the cells were incubated at 25 °C in the dark for 24 h, and then transferred onto appropriate selection plates.

For the isolation of G418-resistant transformants, cells were suspended in 1 ml of MA5 minimal medium, spread onto MA5 minimal agar plates containing 100 μg ml^{-1} G418 and 1 mM uracil, and incubated in a CO_{2} incubator at 5 % (v/v) CO_{2} and 25 °C under fluorescence tubes at 50 μmol photon m^{-2} s^{-1}. The numbers of G418-resistant colonies that developed on the selection plates were counted according to the instructions given by Bio-Rad for 14 days to calculate the transformation frequencies. Each of the single colonies that developed on the agar plates was streaked onto fresh MA5 minimal agar plates, and cells that grew on the plates were used for genotype analyses. The PCR to detect Umps cDNA in Ura^{+} transformants was performed using the primers Umps_CDRF and Umps_CDRR.

Southern blotting

Three micrograms of genomic DNA was digested using restriction enzyme EcoRI or XbaI, separated on 0.8 % (w/v) agarose gels, and blotted onto a Hybond-N+ (GE Healthcare, UK) using a standard capillary transfer method with 20× SSC as the transfer buffer. The blotted filter was baked at 80 °C for 2 h, and the PeLUMPS cDNA fragment was labeled using a DIG High Prime DNA labeling and detection kit (Roche Applied Sciences). Hybridization and signal detection were performed according to the manufacturer’s instructions.

Real-time PCR

Total RNA was extracted from cells grown in MA5 minimal medium to an OD_{750} of 2.0 using a TRIZol® plus RNA purification kit (Ambion), and remaining DNA was digested using a TURBO DNA-free kit (Ambion) according to the manufacturer’s instructions. First-strand cDNA was synthesized using a PrimeScript^™ RT reagent kit with gDNA Eraser (Perfect Real Time, Takara) and an RT primer mix containing oligo (dT)_{18} and random hexamers. Real-time PCR (95 °C for 30 s, followed by 40 cycles of 95 °C for 5 s, and 60 °C for 30 s) to quantify
PeLUMPS cDNA was performed using the primers Umps_QRT_F (5'-TCTTCGACAAGTTGGGATGATG-3') and Umps_QRT_R (5'-TGTACGGTAGTCCGCCAACGAC-3'); position 908–930 of the 1957-bp LUMPS cDNA and PeTUBULIN1 (5'-GGATCAATGGAGGGCAAGT-3'); position 627–646 of the 1989-bp 18S rRNA) and 18SrRNA_QRT_R (5'-GCCCGAAATCCAACTAGG-3'); position 713–732 of the 1989-bp 18S rRNA). The melting curves for each PCR product were determined by measuring the decrease in fluorescence with increasing temperature from 60 to 95 °C. Real-time PCR without a template was also performed in each experiment as a negative control.

Accession numbers
Sequence data from this study can be found in the DDBJ/NCBI data libraries under the accession numbers LC016838 (PeLUMPS), LC016839 (PeLUMPS_cDNA), LC016841 (pUT1), LC016840 (pHA1011), and LC016842 (pUT2).

Additional files
Additional file 1: UV radiation survival curves. UV radiation survival curves for P. ellipsoidea strain Obi.

Additional file 2: Southern blotting. (A) The restriction sites in the 13-kb genomic DNA region comprising PeLUMPS are shown. (B) The restriction sites in pUT1 and pUT2 are shown. (C) Southern blotting of genomic DNA isolated from eight transgenic strains. Genomic DNA was digested using EcoRI or XbaI, and hybridized with a digoxigenin-labeled PeLUMPS cDNA. Lane PC, pUT2 plasmid digested with HindIII and XbaI as a positive control; lane 1, M4; lanes 2–5, representative transgenic M4 with pUT1 (M4-1D, M4-1C, M4-1A, and M4-1B); lanes 6–9, representative transgenic M4 with pUT2 (M4-2A, M4-2B, M4-2D, and M4-2E). The numbers on the left and right sides of the gels indicate molecular size expressed in kilobase pairs. Arrowheads indicate restriction fragments containing the chromosomal LUMPS sequence.

Additional file 3: Construction of pUT1 in two-step PCR. Prub_F and Tact_R carry HindIII and KpnI site, respectively, that were used for the cloning into pBluescript II sk(+).

Abbreviations
5-FOA: 5-fluoroorotic acid; CDR: coding region; ODCase: orotidine-5'-phosphate decarboxylase; OPRase: orotate phosphoribosyltransferase; PeC7N1: actin gene; PeRBCS: 1,5-bisphosphosphate carboxylase/oxygenase small subunit gene; PeTUBULIN1: beta-tubulin gene; PeLUMPS: uridine monophosphate synthase gene; TALEN: transcription activator-like effector nuclease; UMPs: uridine monophosphate synthetase; Ura+: uracil synthesis; UV: ultraviolet light.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
YK designed the experiments, carried out molecular genetic studies, drafted this manuscript and revised it critically for important intellectual content; KO carried out UV mutagenesis and molecular genetic studies and helped to draft the manuscript. JJ designed and carried out molecular genetic studies including genetic transformation and helped to draft the manuscript. YY designed and carried out TALEN mutagenesis and molecular genetic studies, analyzed the data, and helped to draft the manuscript. SH conceived the study, participated in its design and coordination, and helped to draft the manuscript. All authors read and approved the final version of the manuscript.

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