Heterologous Expression of the Barley (Hordeum vulgare L.) Xantha-f, -g and -h Genes that Encode Magnesium Chelatase Subunits

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Abstract

Biosynthesis of chlorophyll involves several enzymatic reactions of which many are shared with the heme biosynthesis pathway. Magnesium chelatase is the first specific enzyme in the chlorophyll pathway. It catalyzes the formation of Mg-protoporphyrin IX from the insertion of Mg$^{2+}$ into protoporphyrin IX. The enzyme consists of three subunits encoded by three genes. The three genes are named Xantha-h, Xantha-g and Xantha-f in barley (Hordeum vulgare L.). The products of the genes have a molecular weight of 38, 78 and 148 kDa, respectively, as mature proteins in the chloroplast. Most studies on magnesium chelatase enzymes have been performed using recombinant proteins of Rhodobacter capsulatus, Synechocystis sp. PCC6803 and Thermosynechococcus elongatus, which are photosynthetic bacteria. In the present study we established a recombinant expression system for barley magnesium chelatase with the long-term goal to obtain structural information of this enigmatic enzyme complex from a higher plant. The genes Xantha-h, -g and -f were cloned in plasmid pET15b, which allowed the production of the three subunits as His-tagged proteins in Escherichia coli BL21(DE3)pLysS. The purified subunits stimulated magnesium chelatase activity of barley plastid extracts and produced activity in assays with only recombinant proteins. In preparation for future structural analyses of the barley magnesium chelatase, stability tests were performed on the subunits and activity assays were screened to find an optimal buffer system and pH.

Keywords Magnesium chelatase · Barley · Xantha · Chlorophyll biosynthesis · Mg-chelatase · Protoporphyrin

1 Introduction

One of the characteristic steps of chlorophyll biosynthesis is performed by the enzyme magnesium chelatase, which inserts Mg$^{2+}$ into the tetrapyrrole protoporphyrin IX to produce Mg-protoporphyrin IX [1–4]. Early studies on barley (Hordeum vulgare L.) mutants suggested that the enzyme requires three gene products encoded by Xantha-h, Xantha-g and Xantha-f [5]. Later, characterization of a "photosynthetic gene cluster" of 45 kbp in the photosynthetic purple bacteria Rhodobacter capsulatus and Rhodobacter sphaeroides identified three genes named bchI, bchD and bchH, which are the structural genes of the three components of magnesium chelatase [6–9]. The magnesium chelatase genes in the cyanobacterium Synechocystis sp. PCC 6803 were also studied and named chlI, chlD and chlH [10, 11]. In order to avoid confusion, the genes are often referred to by their original names and the proteins as I, D and H corresponding to the gene products of Xantha-h/bchI/chlI, Xantha-g/bchD/chlD and Xantha-f/chlH, respectively. The size of the I, D and H subunits are typically in the range of 38–45 kDa, 60–85 kDa and 129–155 kDa, respectively. The specific size of the barley proteins are 38, 78 and 148 kDa as mature proteins after import into the chloroplast.

It was found early that the magnesium chelatase reaction is powered by hydrolysis of ATP [12, 13]. This is now understood in the view that the I and D subunits belong to the large group of AAA + proteins (ATPases Associated with various cellular Activities), which form a two-ringed
structure with three dimers of I in one ring and three dimers of D in the other ring [14]. The H subunit has been suggested to be the catalytic subunit and the target of the ID complex [15]. During catalysis, ATP is specifically hydrolyzed by the I subunits. No ATP hydrolysis has been associated with the D subunit although the N-terminal half of the D subunit is very similar to the I subunit [11, 16].

Most biochemical and structural analyses of magnesium chelatase have been performed with proteins from R. capsulatus, Thermosynechococcus elongatus and Synechocystis sp. PCC 6803 (Table 1). In general, the three subunits are conserved when compared between different species. It can therefore be expected that the overall catalytic mechanism of magnesium chelatase is identical in all bacteriochlorophyll and chlorophyll containing organisms. Still, the I, D and H subunits of plants are distantly related to especially the corresponding proteins of R. capsulatus. To gain a better knowledge of plant magnesium chelatase we have in the present study established an expression system for the three barley magnesium chelatase genes in Escherichia coli and purified the recombinant subunits. The protein subunits showed magnesium chelatase activity and will be used in future studies of the magnesium chelatase structure.

### 2 Material and Methods

#### 2.1 General DNA Techniques

Clones of barley Xantha-h, Xantha-g and Xantha-f inserted into plasmid pET15b were ordered from GenScript (GenScript). Competent E. coli cells were prepared as described by Mandel and Higa [17]. Plasmids were maintained in E. coli XL1-Blue (Stratagene). Expression of genes cloned in pET15b were performed in E. coli BL21(DE3)pLysS.

Plasmids were isolated with the Jet Quick® Plasmid DNA Purification Kit (Life Technologies Corporation).

#### 2.2 Expression of Xantha-h, Xantha-g, Xantha-f

E. coli BL21(DE3)pLysS was transform with plasmids and plated on Tryptose Blood Agar Base (Difco) containing chloramphenicol (25 μg/ml) and ampicillin (50 μg/ml), followed by overnight incubation at 37 °C. Transformants were collected and used to inoculate one liter of LB medium containing 25 mg/ml chloramphenicol and 100 mg/ml ampicillin to an optical density at 600 nm (OD$_{600}$) of 0.05–0.1. The cultures were incubated at 30 °C on a rotary shaker (200 rpm). IPTG (isopropyl β-D-1-thiogalactopyranoside) was added to a concentration of 1 mM when the cultures had reached OD$_{600}$ 0.5–0.7. Incubation continued overnight at 30 °C, except the culture with E. coli BL21(DE3)pLysS/pET15bXanF which was moved to 16 °C and incubated for 40 h. All handling of XanF during expression, purification and activity measurements were performed without direct input of light since the H-subunit of R. capsulatus is known to be light sensitive [18].

#### 2.3 Purification of Recombinant Proteins

Cells of E. coli producing recombinant XanH, XanG and XanF were harvested by centrifugation at 11,325×g for 20 min at 4 °C. Pellets were washed with binding buffer composed of 20 mM imidazole, 0.5 M NaCl, 20 mM Tris–HCl pH 8.0. Aliquots of cell pellets corresponding to 250 ml of cultures were stored at −80 °C. Upon usage, a cell pellet was thawed and resuspended in binding buffer supplemented with 1 mM PMSF (phenylmethanesulfonyl fluoride) and 4 mM DTT (dithiothreitol). A French press at 16,000 psi (pounds per square inch) was used to lyse

| Organism                | Subunit       | Method                  | Tertiary structure | Reference |
|-------------------------|---------------|-------------------------|--------------------|-----------|
| R. capsulatus           | BchI          | X-ray crystallography   | Hexamer            | [33]      |
| R. capsulatus           | BchI          | Negative-stain EM       | Hexamer            | [34]      |
| R. capsulatus           | BchH          | Negative-stain EM       | Monomer            | [35]      |
| R. capsulatus           | BchID complex | Cryo-EM                 | Trimer of dimers   | [14, 36] |
| B. melitensis           | CobST complex | Cryo-EM                 | Trimer of dimers   | [37]      |
| T. elongates            | ChlH          | Negative-stain EM       | Monomer            | [38]      |
| T. elongates            | ChlH          | Negative-stain EM, Small-angle X-ray scattering | Monomer | [39] |
| Synechocystis sp. PCC6803 | ChlH       | Negative-stain EM       | Heptamer           | [40]      |
| Synechocystis sp. PCC6803 | ChlI        | X-ray crystallography   | Hexamer            | [41]      |
| Synechocystis sp. PCC6803 | ChlH        | X-ray crystallography   | Dimer              | [42]      |

EM electron microscopy
cells followed by centrifugation at 48,384×g, 20 min, 4 °C. XanH and XanF proteins were further purified from the soluble fractions. Similar to the *R. capsulatus* D-subunit, XanG formed inclusion bodies and was obtained in the pellet fraction. The XanG protein was solubilized from the pellet fraction according to the method to solubilize the *R. capsulatus* D-subunit [19]. The pellet containing XanG was resuspended in binding buffer containing 6 M urea and incubated for 1 h on ice before centrifugation at 5000×g for 20 min at 4 °C. XanG was further purified from the urea containing supernatant.

The recombinant XanH, XanG and XanF were purified by immobilized metal ion affinity chromatography (IMAC) using 1 ml HisTrap™ FF Crude columns (GE Healthcare). After loading the soluble fractions through the column, washing was performed with 10 column volumes of binding buffer followed by 10 column volumes of wash buffer composed of 40 mM imidazole, 0.5 M NaCl, 20 mM Tris–HCl pH 8.0. Protein was eluted with elution buffer composed of 250 mM imidazole, 0.5 M NaCl, 20 mM Tris–HCl pH 8.0. Six M urea was included in all buffers for purification of XanG. The eluted proteins were desalted with NAP-10 columns (GE Healthcare). XanH was desalted into 20 mM Tricine-NaOH pH 8.0. XanG was desalted into 6 M urea, 20 mM Tris–HCl pH 8.0. XanF was desalted into 50 mM Tricine-NaOH pH 8.0, 250 mM NaCl, 50 mM MgCl₂. Protein concentrations were determined by Bradford Reagent (BioRad).

### 2.4 Preparation of Barley Chloroplast Proteins

Seeds of barley cultivar Bonus were planted into two 50 × 60 cm trays in moist vermiculate and grown in darkness at room temperature. After eight days, etiolated yellow seedlings were illuminated for 4–5 h before harvest. The top 7 cm of the primary leaves were collected and homogenized in a blender with exchangeable razor blades in grinding medium composed of 0.4 M D-mannitol, 20 mM Tricine-NaOH pH 9.0 and 1 mM DTT. The mixture was then filtered through a double layer of nylon tissue and centrifuged at 2000×g, 4 °C, 5 min. The pellet was resuspended in 50 ml grinding medium and aliquots of 6 ml were layered on 8 ml Percoll cushions (40% Percoll [Sigma-Aldrich] and 1 mM DTT in grinding buffer) in 15 ml tubes and centrifuged at 3220×g, 4 °C, 40 min with a swing-out rotor. The pellet was resuspended in grinding medium, transferred to a fresh tube and centrifuged at 2000×g, 4 °C for 5 min. The plastids in the resulting pellet were resuspended and lysed in one ml of lysis buffer composed of 20 mM Tricine-NaOH pH 9.0, 1 mM DTT and 1 mM PMSF. Next, the lysate was ultracentrifuged at 100,000×g, 4 °C for 20 min using a fixed angle rotor. Supernatants were collected and glycerol was added to a final concentration of 10%. Aliquots were stored at −80 °C and used in enzyme assays.

### 2.5 Magnesium Chelatase Assays

Magnesium chelatase assays were performed in total volumes of 50 μl and contained 20 mM Tricine-NaOH pH 9.0, 4 mM ATP, 10 mM creatine phosphate, 15 mM MgCl₂, 2 μM deuteroporphyrin IX, 2.5 units creatine phosphokinase and recombinant barley magnesium chelatase subunits. Assays were stopped after 30 min by addition of 950 μl acetone:water:32% ammonia (80:20:1, vol:vol:vol) followed by centrifugation for 2 min at 17,000×g to remove precipitated protein. The assays were kept in dark during mixing and incubation to prevent photo-oxidation. The supernatants were analyzed on a Shimadzu RF-5301PC spectrophotometer with excitation wavelength set at 408 nm and emission detected between 550 and 650 nm. Excitation and emission slit widths were 5 nm and 10 nm, respectively. The amount of formed Mg-deuteroporphyrin IX was determined from the peak at 585 nm. For testing the effect of different buffer systems 20 mM MOPS-NaOH (pH 6.5, 7.0 and 7.5), 20 mM Tris–HCl (pH 7.5, 8.0, 8.5 and 9.0), 20 mM Tricine-NaOH (pH 7.5, 8.0, 8.5 and 9.0), 20 mM HEPES–NaOH (pH 7.0, 7.5 and 8.0) and 20 mM CHES–NaOH (pH 8.5, 9.0, 9.5 and 10.0) were used.

### 2.6 Phylogenetic Analysis

Barley XanH (GenBank: ABF72535.3), XanG (GenBank: AAZ32779.1) and XanF (GenBank: KAE8772866.1) polypeptide sequences were queried against UniProtKB (www.uniprot.org) to retrieve homologs from other species. Transit peptides were identified by ChloroP [20] and removed from the plant polypeptides before alignments were made. Protein sequences showing ≥ 40% identity to the query sequence were used for phylogenetic analysis. To perform phylogenetic analysis, protein sequences were initially aligned using the MUSCLE algorithm implemented in MEGAX with default setting [21]. A maximum likelihood phylogenetic tree was constructed using the maximum likelihood heuristic method named Nearest Neighbor Interchange implemented in MEGAX with default setting [21]. A maximum likelihood phylogenetic tree was constructed using the maximum likelihood heuristic method named Nearest Neighbor Interchange implemented in MEGAX. The bootstrap consensus tree inferred from 1000 replicates is taken to represent the evolutionary history of the analyzed sequences. Branches which correspond to partitions replicated in bootstrap replicates of less than 50% collapse.
3 Results

3.1 Phylogenetic Analysis of Magnesium Chelatase Subunits

Magnesium chelatase occurs in a broad variety of photosynthetic species that use bacteriochlorophyll or chlorophyll as the light absorbing pigment. We selected 44 different proteins belonging to the three magnesium chelatase subunits and performed a phylogenetic analysis (Fig. 1). Within each clade corresponding to each subunit, the proteins of the photosynthetic purple bacterium R. capsulatus are the most divergent. The proteins of the cyanobacteria T. elongatus and Synechocystis are relatively closely related to each other. The magnesium chelatase proteins of higher plants are distinctly different to those of R. capsulatus, T. elongatus and Synechocystis. This encouraged us to set up a pipeline for production of barley magnesium chelatase in order to get insight of this protein complex in higher plants.

3.2 Construction of Expression Plasmids

In order to obtain recombinant subunits of barley magnesium chelatase, Xantha-h, -g and -f were expressed from plasmid pET15b in E. coli, which resulted in N-terminally His-tagged proteins that could be purified with immobilized metal ion affinity chromatography (IMAC). We chose to use N-terminal His-tags, which have been successful in most magnesium chelatase expression systems. C-terminal His-tags have sometimes been used for the I subunit since an early report found that an N-terminal His-tag on the I subunit was inhibitory [19]. In barley and other plants, the magnesium chelatase subunits are imported into the chloroplast with help of a transit peptide. ChloroP [20] was used to predict the transit peptide sequence of the three subunits. The genes, corresponding to the mature proteins, were ordered from GenScript and cloned between the NdeI–XhoI restriction sites of pET15b. The synthetic genes were codon optimized for expression in E. coli. The expression plasmids were named pET15bXanH, pET15bXanG and pET15bXanF and the resulting polypeptides are shown in Fig. 2.

3.3 Production and Purification of Barley Magnesium Chelatase Subunits

The barley magnesium chelatase genes were expressed in E. coli BL21(DE3)pLysS from plasmids pET15bXanH, pET15bXanG and pET15bXanF. E. coli BL21(DE3) was less efficient (data not shown). XanG formed inclusion bodies, which had to be solubilized with urea. This is also the case for the R. capsulatus ortholog [19]. Thus, all buffers used for purification of XanG contained 6 M urea. Also, XanH and XanF tended to form inclusion bodies but enough protein remained in supernatant fractions after centrifugation for further purification. The subunits were purified by IMAC.
followed by desalting. Typically, 1–8 mg of protein were obtained from 250 ml of bacterial culture (Fig. 3).

### 3.4 Active Recombinant Barley Magnesium Chelatase Subunits

Magnetese chelatase activity can be measured with protein extract isolated from barley chloroplasts [22]. In the current study we prepared such plastid extract, which showed magnesium chelatase activity when used in enzymatic assays demonstrating that all magnesium chelatase subunits are present in the plastid extract. We hypothesized that addition of any of the three barley magnesium chelatase subunits produced in *E. coli* as recombinant proteins would further increase the enzymatic activity. Thus, combining the barley plastid extract with fractions from the heterologous expression system in *E. coli* allowed us to work with one expression system at the time until an active recombinant magnesium chelatase subunit was obtained. Therefore, various purification fractions of recombinant XanF, XanG or XanH were added to magnesium chelatase assay mixtures containing barley plastid extract and a stimulation of activity indicated that an active recombinant subunit had been prepared (data not shown). When all three subunits showed activity in these assays, they were successfully combined with each other without plastid extracts. In different assays, the amount of two subunits were kept constant and the third unit was varied (Fig. 4). In our current model of the enzymatic mechanism based on the *R. capsulatus* magnesium chelatase, the D subunit forms a hexameric platform for the stepwise assembly of the I subunits into a two-tiered hexameric ring [3].

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**Fig. 2** The polypeptide sequences of the barley magnesium chelatase subunits as produced from pET15bXanH, pET15bXanG and pET15bXanF. The sequence marked with grey is derived from pET15b and contains the His-tag
Heterologous Expression of the Barley (*Hordeum vulgare* L.) Xantha-\(f\), -\(g\) and -\(h\) Genes that…

excess of the D subunit over the I subunit lowers the activity [16]. In contrast, increasing amounts of I and H subunits lead to saturation of these subunits and hyperbolic curves. Our initial experiments with the barley magnesium chelatase subunits suggest that they follow the same enzymatic mechanism (Fig. 4).

### 3.5 Finding Optimal Conditions for Barley Magnesium Chelatase

Since structural analysis is the long-term goal of our work with the barley magnesium chelatase, it was important to obtain basic information about optimal buffer systems and storage conditions. MOPS, Tris, Tricine, HEPES and CHES where used to cover different pH ranges and used in magnesium chelatase assays with the recombinant subunits. The barley magnesium chelatase appeared to be relatively tolerant regarding the range of pH. Good activity was obtained in all buffer systems between pH 8.0–9.5, with the exception of CHES at pH 8.5 (Fig. 5).

The purified subunits were tested for their thermal stability by exposing them to different storage conditions over 24 h. The temperatures varied from −80 to 37 °C and also included repeated freeze thawing of a samples stored at −20 °C. After 24 h the samples were analyzed by SDS-PAGE. In general, the subunits withstood the treatments. The only negative effect was seen in the case of XanF stored at 30 and 37 °C (Fig. 6).

### 4 Discussion

Chlorophyll and heme are the most abundant tetapyrroles in biological systems. They share the first initial catalytic steps of the tetapyrrole biosynthetic pathway [4, 23]. A major difference between them is the bound metal ion, which is magnesium in chlorophyll and iron in heme. Protoporphyrin IX is their last common intermediate and used by magnesium...
chelatase or ferrochelatase for the insertion of Mg$^{2+}$ or Fe$^{2+}$, respectively. Although the porphyrin substrate is the same, the two enzymes show no obvious similarities. While ferrochelatase is a monomer or homodimer, magnesium chelatase is a large protein complex composed of three gene products and requires ATP for catalysis [24]. Interestingly, there are two different cobalt chelatases involved in synthesis of cobalamin (vitamin B$_{12}$). The cobalt chelatase participating in the anaerobic pathway is structurally similar to ferrochelatase, while the cobalt chelatase involved in the aerobic pathway belongs to the AAA+ protein family like magnesium chelatase and requires input of energy in the form of ATP [25]. Today, the structural information available for the AAA+ class of chelatases is obtained from a limited group of organisms; *R. capsulatus*, *Synechocystis* sp. PCC 6803, *T. elongatus* and *Brucella melitensis* (Table 1). We would like to obtain structural insight into the magnesium chelatase of plants and therefore established an expression system of the barley magnesium chelatase genes since barley has a long history in the field of magnesium chelatase research. The oldest mutant in barley, *xantha-f.10*, was induced 1953 by X-ray in the cultivar Bonus [26]. A total of 20 barley mutants are available and the mutations have been described at DNA level [27–31]. The barley magnesium chelatase proteins are most different from those of *R. capsulatus*, which is obvious from a phylogenetic analysis of the three subunits (Fig. 1). *R. capsulatus* produces bacteriochlorophyll under anaerobic conditions and the differences in the polypeptide sequences are probably adaptations to an anaerobic environment [32]. In alignments between the *R. capsulatus* D polypeptide to the D subunits of plants, several gaps are found [3]. Several gaps are also found in alignments of H subunits but the *R. capsulatus* H sequence has also an addition of an iron-sulfur cluster sequence that has been suggested to be involved in regulation of magnesium chelatase activity in this bacterium [32]. We hope that the present heterologous expression system of barley magnesium chelatase will allow us to perform studies on a plant magnesium chelatase and reveal plant specific information of this key enzyme of chlorophyll biosynthesis.

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**Author Contributions** MH conceived the idea of the work; RM, DS, MH and HMY performed experiments, analyzed data and wrote the manuscript.

**Compliance with Ethical Standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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