Improving the Secretion of a Methyl Parathion Hydrolase in *Pichia pastoris* by Modifying Its N-Terminal Sequence

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Abstract

*Pichia pastoris* is commonly used to express and secrete target proteins, although not all recombinant proteins can be successfully produced. In this study, we used methyl parathion hydrolase (MPH) from *Ochrobactrum* sp. M231 as a model to study the importance of the N-terminus of the protein for its secretion. While MPH can be efficiently expressed intracellularly in *P. pastoris*, it is not secreted into the extracellular environment. Three MPH mutants (N66-MPH, D10-MPH, and N9-MPH) were constructed through modification of its N-terminus, and the secretion of each by *P. pastoris* was improved when compared to wild-type MPH. The level of secreted D10-MPH was increased to 0.21 U/mL, while that of N9-MPH was enhanced to 0.16 U/mL. Although N66-MPH was not enzymatically active, it was secreted efficiently, and these results demonstrate that the secretion of heterologous proteins in *P. pastoris* may be improved by modifying their N-terminal structures.

Citation: Wang P, Huang L, Jiang H, Tian J, Chu X, et al. (2014) Improving The Secretion of a Methyl Parathion Hydrolase in *Pichia pastoris* by Modifying its N-Terminal Sequence. PLoS ONE 9(5): e96974. doi:10.1371/journal.pone.0096974

Editor: Israel Silman, Weizmann Institute of Science, Israel

Received September 17, 2013; Accepted April 14, 2014; Published May 7, 2014

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Funding: This work was supported by the National High Technology Research and Development Program of China (863 Program, 2013AA102804) and the National Natural Science Foundation of China (NSFC, Grant no. 30900839). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

*Pichia pastoris* is a methylotrophic yeast that has been genetically engineered to express heterologous proteins for use in research, as well as industrial and pharmaceutical applications [1]. The use of *P. pastoris* as a host offers many advantages compared with other expression systems. For example, the yeast can be used to express large amounts of extracellular proteins at relatively little cost [2]. The preferential secretion of recombinant proteins allows for the direct isolation of target proteins from culture media, eliminating the need for high-cost, low-yield cell disruption. Furthermore, this feature limits the toxicity issues resulting from intracellular accumulation of target proteins [2]. In the *P. pastoris* expression system, signal peptides are commonly included in commercially available vectors, and so can be attached to the recombinant protein, causing it to be exported from the cell. Because *P. pastoris* secretes few intrinsic proteins, the recombinant protein forms the major polypeptide species found in the extracellular growth medium, which facilitates purification of the heterologous protein [3].

However, the *P. pastoris* system also has several limitations. Many recombinant proteins that should be secreted are retained inside the yeast cell [4–6]. Furthermore, these proteins are often mis-folded and degraded intracellularly, which can be a major problem [7]. Because the system is used widely, a number of strategies for enhancing protein production have been developed. However, most strategies do not target the protein itself, but instead modify other aspects of the process, such as the host [8], promoter [8], signal peptide [9,10], chaperone [11], fusion proteins [12], protease [13], fermentation conditions [14–18], codon optimization [19], or gene copy number [20]. Although these methods can effectively enhance the expression of some proteins, it is challenging to enhance the secretion of certain proteins that are retained in the cell. Previous studies have shown that the rational design of the internal regions of a protein can enhance its secretion by *P. pastoris* [21–24]. Therefore, some secretion signals that affect protein secretion must exist in the internal regions of proteins. It may therefore be necessary to modify the sequence or structure of a protein to enhance its secretion.

In this study, we used methyl parathion hydrolase (MPH) from *Ochrobactrum* sp. M231 to investigate the effect of modification of its N-terminus on secretion. Organophosphorus hydrolases (OPHs) play important roles in the decontamination and bioremediation of environments polluted by organophosphate pesticides [25–33]. MPH, isolated by our lab from *Ochrobactrum* sp. M231 in 2008 [29], can efficiently and specifically degrade methyl parathion, but it cannot be secreted from *P. pastoris* [29,33–35]. However, another OPH, OPHC2, isolated from *Pseudomonas pseudoalcaligenes*, also by our laboratory [32], has a similar three-dimensional structure to MPH. It has been over-expressed and efficiently secreted at concentrations of up to 5.5 g/L in *P. pastoris* [31,32]. These two proteins have the same function and share a sequence identity of 47.7%. However, they have different secretion patterns when expressed in the same *P. pastoris* expression system with an identical promoter and signal peptide. Although it has been reported that integrating 12 copies of the MPH expression cassette into the *P.
plasmids (N66-MPH and N9-MPH) were then generated through homologous recombination, as described previously [35], using the vector. N9-MPH was generated from homologous recombination of three fragments: fragment 1 (using primers N9-F and pET30-R) and fragment 2 (using primers MPH-F-66 and pET30-R) and fragment 3 (using primers OPHC2-F-1 and pET30-R). The plasmids containing wild-type and mutant vectors were transformed into Top10 cells and their fidelity was confirmed by sequencing.

Strains, Plasmids, and Media

The GenBank accession numbers of Ochrobactrum sp. M231 MPH and OPHC2 are ACC63894 and CAE53631, respectively. Escherichia coli strain Top10 and the plasmid pGEM-T were purchased from Promega Corp. (Madison, WI, USA). E. coli strain BL21(DE3) and the expression plasmid pET-30a(+) were purchased from Novagen (Darmstadt, Germany). P. pastoris strain GS115 and the vector pPIC9 were purchased from Invitrogen (Carlsbad, CA).

E. coli cells were cultured aerobically at 37°C in Luria-Bertani medium. Minimal dextrose (MD) medium, buffered complex glycerol medium (BMGY), yeast extract peptone dextrose medium (YPD), and buffered complex menthol medium (BMMY) were prepared according to the manufacturer’s instructions (Invitrogen).

The Design of MPH Mutants

The SCHEMA software [38] was used to compare recombination sites in the N-termini of MPH and OPHC2, using standard parameters, resulting in N-terminal blocks comprising 66 amino acids of MPH, and 68 amino acids of OPHC2. We designed a chimera (N66-MPH), where a 68-amino-acid block of OPHC2 was used to replace the 66 N-terminal amino acids of MPH. Wild-type MPH has a coil structure formed by 10 amino acids at its N-terminus, which might prevent the secretory pathway from recognizing its α-factor signal peptide. We therefore designed a deletion mutant (D10-MPH, in which the 10 N-terminal amino acids of MPH had been removed), and a chimera (N6-MPH, in which the 9 N-terminal amino acids of MPH were replaced by those from OPHC2).

Construction of Expression Vectors in E. coli

Mutant vectors were synthesized using the plasmids pET-30a(+)—mph [33] and pET-30a(+)—ophc2 [31], recombinant plasmids containing mph and ophc2, respectively, as the templates. Two plasmids (N66-MPH and N9-MPH) were then generated through homologous recombination, as described previously [35], using the primers listed in Table S1 in File S1. N66-MPH was generated from homologous recombination of three fragments: fragment 1 (using primers MPH-F-66 and pET30-R) and fragment 2 (using primers MPH-R-0 and pET30-F) were amplified from the pET-30a(+)—mph plasmid, while fragment 3 (using primers OPHC2-F-I and OPHC2-R-68) was amplified from the pET-30a(+)—ophc2 vector. N9-MPH was generated from homologous recombination of two fragments: fragment 1 (using primers N9-R and pET30-F) and fragment 2 (using primers N9-F and pET30-R) were amplified from the pET-30a(+)—mph plasmid. The D10-MPH gene was amplified from pET-30a(+)—mph with the primers D10-F and D10-R. The PCR products were digested with EcoRI and NdeI, and cloned into pET-30a(+). To verify the inserted genes, DNA sequencing was performed at the State Key Laboratory of Crop Genetic Improvement, Chinese Academy of Agricultural Sciences (Beijing, China). The verified plasmids were transformed into competent E. coli BL21(DE3) cells for expression.

Expression, Purification, and Quantification of Wild-type MPH and Mutants

A single colony of the transformed E. coli was cultured in Luria-Bertani liquid medium containing 50 μg/mL kanamycin (LB-kan) at 37°C overnight, and then inoculated to fresh LB-kan (1:100 dilution) and incubated again at 37°C. When the OD600 of the culture reached 0.5, isopropyl β-D-1-thiogalactopyranoside (final concentration, 0.4 mM) was added. Cultures were incubated for an additional 18–20 h at 16°C. The cells were then collected by centrifugation and disrupted by sonication. The recombinant proteins were purified with Ni-NTA Superflow (QIAGEN, USA) according to the manufacturer’s instructions. The final concentration of the purified protein was determined using the Bio-Rad Protein Assay kit (Bio-Rad, Hercules, CA, USA).

Enzymatic Properties of Wild-type and Mutant MPH

The standard enzyme assay and determination of kinetic parameters were performed as described by the reference [35]. The determination of the enzymatic properties of wild-type and mutant MPH was performed according to methods described previously [33,34].

Construction of Expression Vector and Transformation of P. pastoris

The wild-type and mutant plasmids were digested with EcoRV and NdeI, and ligated into pPIC9. All plasmids were transformed into Top10 cells and their fidelity was confirmed by sequencing. The plasmids containing wild-type and mutant vectors were extracted using a plasmid extraction kit (TIANGEN, Beijing, China). BglII was used to linearize 5–10-μg recombinant DNA, which was then transformed into P. pastoris GS115 cells using the Gene Pulser system (Bio-Rad; conditions used: 2.5 kV, 25 μF, and 400 Ω). His+ transformants were selected on MD plates, and the genomic DNA was extracted and analyzed by PCR using the primers 5’-AOX1 and 3’-AOX1 (Table S1 in File S1).

Selection of High-producing Recombinant P. pastoris Strains

After transformation, the His+ transformants from the MD plates were grown in 3-mL BMGY, and induced in 1-mL BMMY for 48 h. Next, 100 clones each from the wild type and three mutant MPH transformants (400 clones in total) were assessed for the secretion of the expressed proteins using the standard enzyme assay. We also selected high-producing recombinant P. pastoris strains for the wild type and each of the three mutants for the shake-flask culture.

Expression of MPH and Mutant Proteins in Shake-flask Culture

The colonies of His+ transformants exhibiting MPH activity were inoculated into 45-mL BMGY at 28°C with constant shaking at 200 rpm until the OD600 reached 5.0. Cells were pelleted by centrifugation and resuspended in 15-mL BMMY, then induced at 28°C with constant shaking at 200 rpm for 120 h. Methanol was added to a final concentration of 0.5% (v/v) every 24 h. The culture supernatant and cells were harvested by centrifugation (12,000 g, 3 min, 4°C) to analyze MPH activity according to methods described previously [36]. MPH activity in the supernatant and cells was determined using the standard enzyme assay.
Isolation of Yeast RNA and Quantitative Real-time PCR

Yeast expression cultures were induced for 1 day with 0.5% methanol, and adjusted to an OD_{600} of 8.0. Cell pellets from 1-mL samples of the cultures were collected by centrifugation and lysed by grinding in liquid nitrogen. Total RNA was isolated using TRIzol (TIANGEN, Beijing, China) following the manufacturer’s instructions, and cDNA was synthesized using the TIANGEN RT Kit (TIANGEN, Beijing, China). Duplicate PCR reactions were performed on an ABI PRISM 7700 sequence detection system (Applied Biosystems, Weiterstadt, Germany) with standard conditions (50°C for 2 min; 95°C for 10 min; and 45 cycles of 95°C for 15 sec, and 60°C for 1 min) using TransStart Probe qPCR SuperMix (Transgen, Beijing, China) according to the manufacturer’s recommendations. The primers used for mph were mph-mF and mph-mR (Table S1 in File S1). Yeast gapdh was amplified as a control for normalization using the primers GAPDH-F and GAPDH-R (Table S1 in File S1). All primers were used at a final concentration of 0.2 μM.

Gene Copy Number Determination by Real-time PCR

Genomic DNA was prepared using the TIANamp Yeast DNA Kit (TIANGEN, Beijing, China). Gene copy numbers of mph and mutants were determined according to the absolute quantification method described in the previous section by using the SYBR Green Real-time PCR Master Mix-Plus (Toyobo, Osaka, Japan). The primers and quantitative real-time PCR protocol were as described in the previous section.

Protein Structure Analysis

The tertiary structures of MPH, OPHC2, and N9-MPH were constructed using the Discovery Studio software v.2.5.5 (Accelrys Software Inc., USA). Molecular dynamics simulations (MDS) of the protein structures were carried out for 10 ns at 300 K using the Gromacs v.4.5.5 software, as described previously [33]. The interaction energy was calculated using the VMD v.1.8.6 software, following standard protocols.

Results

Three-dimensional Models of MPH, OPHC2, and N9-MPH

The 3D structures of MPH, OPHC2, and N9-MPH were modeled based on the crystal structure of Pseudomonas sp. MPH (PDB reference: 1P9E) [40]. The resulting model (Figure 1) is a dimer that can be described as a ββ/βα sandwich, which is typical of metallo-hydrolase/oxidoreductase folding. As expected, the structures of MPH and OPHC2 are similar, since they have high sequence identity. Based on the models of the 3D protein structures, the recombination sites at the N-terminus were identified as residues 66 of MPH and 68 of OPHC2 using the Schima software [38].

We next assessed the importance of the N-terminus for protein secretion. As shown in Figure S1 in File S1, the nine residues at the N-terminus of MPH and OPHC2 are different, and only three residues are consistent when the sequences are aligned. However, the total sequence identity between these regions of MPH and OPHC2 is 47.7%. In addition, wild-type MPH has a coil structure in the first 10 amino acids of its N-terminus, which might prevent recognition of its α-factor signal peptide by the secretory pathway. Therefore, two MPH mutants (D10-MPH and N9-MPH) were constructed to assess the effect of these residues on the secretion of MPH. In D10-MPH, the 10 amino acids from the N-terminus were deleted, while in N9-MPH, the 9 N-terminal amino acids of MPH were replaced by those from OPHC2.

Expression of MPH and Mutant Proteins in P. pastoris

To investigate the effects of the mutations on the expression and secretion of MPH in P. pastoris, pPIC9-based yeast expression constructs (pPIC9-N66-mph, pPIC9-D10-mph, pPIC9-N9-mph) were generated. One hundred clones each from wild-type and three mutant MPH transformants were grown, and the secretion of the expressed proteins was assessed using the standard enzyme assay. As shown in Figure 2, the supernatants from most D10-MPH and N66-MPH transformants showed MPH activity higher than 0.1 U. In contrast, supernatant from wild-type MPH transformants exhibited activity at 0.05 U. The N9-MPH clone, which had 34 mutated amino acids compared with wild-type MPH, had low enzymatic activity, even though SDS-PAGE confirmed its expression and secretion by P. pastoris (data not shown). We also selected the following high-producing recombinant P. pastoris strains of each gene for the next shake-flask culture: MPH-24# (0.03 U/mL), N66-MPH-5# (0.02 U/mL), D10-MPH-56# (0.15 U/mL), and N9-MPH-70# (0.15 U/mL).

Production and Activity of Wild-type and Mutant MPH Proteins

The four transformants (MPH-24#, N66-MPH-5#, D10-MPH-56#, and N9-MPH-70#) were cultured and the MPH activities in methanol-induced cultures at 28°C were calculated. As shown in Figure 3, the maximum activity was observed after 108 h for D10-MPH-56# and N9-MPH-70#, while N66-MPH-5# showed the lowest extracellular MPH activity. The proteins in each supernatant were collected after 120 h of induction and analyzed by SDS-PAGE. As shown in Figure 4, N66-MPH-5#, D10-MPH-56#, and N9-MPH-70# all showed distinctive bands with a molecular weight of ~35 kDa. The bands were subsequently identified as the appropriate MPHs by mass spectroscopy (data not shown). However, no band corresponding to MPH-24# was observed. These data suggest that N66-MPH, D10-MPH, and N9-MPH, but not MPH-24#, can be efficiently secreted from yeast cells. We also detected the intracellular MPH activity. As shown in Figure S2 in File S1, the strain containing wild-type MPH had higher intracellular MPH activity than the three mutant strains (N9-MPH, D10-MPH, and N66-MPH).
Specifically, the intracellular MPH activity of *P. pastoris* strain MPH-24# reached 1.1 U/mL after 24 h, and eventually increased to 3 U/mL after 120 h (Figure S2b in File S1). These results indicate that the mutations facilitated secretion of the expressed proteins from the cell. Meanwhile, intracellular proteins are released upon cell lysis, which may explain why we did not detect MPH activity in supernatants from the initial cultures of wild-type MPH, but later identified weak MPH activity (Figure 3, Figure S2 in File S1). These experiments were repeated three times with similar results. During the 5-day induction period, the four selected transformants had comparable growth rates (Figure S3 in File S1). This suggests that the increased MPH levels produced by the three
We employed the molecular mass of 38 kDa on SDS-PAGE (Figure S4 in File S1). The expressed protein migrated as a single band with an approximate P. mutan mutants as wild-type MPH was not secreted from E. coli expressed in the pastoris are shown in Table 2. All of the mutants, except N66-MPH, had activities measured. The kinetic parameters of the enzymes MPH were cloned and expressed in N9-MPH) were single-copy clones (Table 1). Therefore, the normalization. Data are expressed as the means of duplicate samples, and error bars indicate standard deviation (SD).

**Discussion**

In this study, MPH from Ochrobactrum sp. M231 was selected and the importance of its N-terminus in its secretion by P. pastoris was determined. We swapped the corresponding block of sequence from OPHC2 based the Schema software analysis [38] and removed the N-terminal block of MPH according to the 3D protein structures. Our results revealed that the N-terminal region plays an important role in secretion. In addition, the improved secretion of the MPH mutants was not due to differences in growth rate, mRNA expression, gene copy number, or stability. Although the mutant proteins had reduced catalytic efficiency, the secretion of both D10-MPH and N9-MPH was improved significantly compared to wild-type MPH, as demonstrated by SDS-PAGE (Figure S5 in File S1).

Because the N-terminus of the protein plays an important role in its secretion, we used MDS to predict structural differences between the proteins. As shown in Figure 1, the N-terminus of each chain (A or B) directly interacts with the other chain, and thus plays an important role in dimer formation. The total interaction energy between chains A and B was calculated by MDS modeling (Figure S6 in File S1). MPH and OPHC2 have similar interaction energies between the chains. Specifically, the average interaction energies between chains A and B of MPH, OPHC2, and N9-MPH during the final 5 ns of MDS were –858.3, –847.3, and –627.1 kcal/mol, respectively. However, the average interaction energies of the N-terminus (the first 10 amino acids) and other regions of the protein for the final 5 ns of MDS were –670.0 and –530.9 kcal/mol for MPH and OPHC2, respectively. These results suggest that the N-terminus of OPHC2 was more flexible than that of MPH. As shown in Figure S6 in File S1, the interaction energies of the N-terminus of N9-MPH with the other protein regions was also lower than that of MPH. A more flexible N-terminus is therefore important for optimal protein secretion.

**Figure 5. Analysis of mph mRNA levels.** The expression of the recombinant cDNAs was assessed by quantitative real-time PCR, using GAPDH for normalization. Data are expressed as the means of duplicate samples, and error bars indicate standard deviation (SD).

doi:10.1371/journal.pone.0096974.g005
These results indicate that the N-terminus of the MPH protein contains a key sequence factor that affects its secretion from P. pastoris. In this study, we used a specific method to modify the protein N-terminal sequence. However, highly specific strategies will be necessary to optimize the N-terminal sequences of other proteins, and software should be developed for such work in the future.

Supporting Information

File S1 Supporting figures and tables. This file contains Table S1-Table S2 and Figure S1-Figure S6. Table S1, The enzymatic properties of WT and mutant MPH. Figure S1, The sequence alignment of N-terminal of the three proteins.

Table 1. The copy number calculation of WT and mutant MPH according to absolute (Abs. Q).

| Pichia pastoris strain | Abs. Q (SYBR Green) | Resulting copy number |
|------------------------|----------------------|-----------------------|
| N9-MPH-5#              | 1.1±0.1              | 1                     |
| D10-MPH-56#            | 1.3±0.1              | 1                     |
| N9-MPH-70#             | 0.8±0.1              | 1                     |
| WT MPH-24#             | 1.0±0.1              | 1                     |

doi:10.1371/journal.pone.0096974.t001

Table 2. The kinetic parameters of WT and mutant MPH.

|                  | $k_{cat}$ (min$^{-1}$) | $K_m$ (μM) | $k_{cat}/K_m$ (μM$^{-1}$min$^{-1}$) |
|------------------|------------------------|------------|-------------------------------------|
| WT MPH           | 253.80±11.41           | 73.17±4.70 | 3.47                                |
| D10-MPH*         | 77.53±2.66             | 86.87±4.01 | 0.89                                |
| N9-MPH*          | 208.8±9.14             | 145.5±6.51 | 1.44                                |

*The first 10 amino acids of N-terminus of MPH were deleted.
†The first nine amino acids from the N-terminus of MPH were replaced by those from OPCH2.

doi:10.1371/journal.pone.0096974.t002

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