Construction of New Ligation-Independent Cloning Vectors for the Expression and Purification of Recombinant Proteins in Silkworms Using BmNPV Bacmid System

Doucet Daniel, Kato Tatsuya, Thompson James R., Park Enoch Y.

journal or publication title: PLoS ONE
volume: 8
number: 5
page range: e64007
year: 2013-05-10

(C) 2013 Kato et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

URL: http://hdl.handle.net/10297/7326
doi: 10.1371/journal.pone.0064007
Construction of New Ligation-Independent Cloning Vectors for the Expression and Purification of Recombinant Proteins in Silkworms Using BmNPV Bacmid System

Tatsuya Kato1, James R. Thompson2, Enoch Y. Park1,3,4*

1 Laboratory of Biotechnology, Department of Applied Biological Chemistry, Faculty of Agriculture, Shizuoka University, Suruga-ku, Shizuoka, Japan, 2 Department of Physiology and Biomedical Engineering, Mayo Clinic College of Medicine, Rochester, Minnesota, United States of America, 3 Laboratory of Biotechnology, Integrated Bioscience Section, Graduate School of Science and Technology, Shizuoka University, Suruga-ku, Shizuoka, Japan, 4 Laboratory of Biotechnology, Research Institute of Green Science and Technology, Shizuoka University, Suruga-ku, Shizuoka, Japan

Abstract

A ligation independent cloning (LIC) system has been developed to facilitate the rapid and high-efficiency cloning of genes in a Bombyx mori expression system. This system was confirmed by the expression of human microsomal triglyceride transfer protein (hMTP) fused with EGFP in silkworm larvae and pupae. Moreover, hMTP and human protein disulfide isomerase (hPDI) genes were inserted into two LIC vectors harboring gcLINK sequences and were combined by using the LIC through gcLINK sequences. The constructed vector was incorporated into the Bombyx mori nucleopolyhedrovirus (BmNPV) bacmid, and injected into silkworm larvae. The expressed hMTP-hPDI complex was purified from the fat bodies of silkworm larvae. This LIC vector system was applied to express the E1, E2, and E3 subunits of human γ-ketoglutarate dehydrogenase (KGDH) in silkworm larvae. The expressed proteins were purified easily from fat bodies using three different affinity chromatography steps. The LIC vectors constructed as described in this report allow for the rapid expression and purification of recombinant proteins or their complexes by using the BmNPV bacmid system.

Citation: Kato T, Thompson JR, Park EY (2013) Construction of New Ligation-Independent Cloning Vectors for the Expression and Purification of Recombinant Proteins in Silkworms Using BmNPV Bacmid System. PLoS ONE 8(5): e64007. doi:10.1371/journal.pone.0064007

Editor: Daniel Doucet, Natural Resources Canada, Canada

Received November 21, 2012; Accepted April 8, 2013; Published May 10, 2013

Copyright: © 2013 Kato et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported partly by Strategic Young Researcher Overseas Visits Program for Accelerating Brain Circulation from Japan Society for the Promotion of Science (JSPS), Japan, and funded partly by Promotion of Nanobio-technology Research to support aging and welfare society from Ministry of Education, Culture, Sports, Science and Technology (MEXT), Japan. 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.

* E-mail: acypark@ipc.shizuoka.ac.jp

Introduction

Generally, to produce recombinant proteins, target genes are amplified by PCR with primers containing restriction enzyme sites. PCR products and expression vectors are digested by restriction enzymes and ligated using T4 DNA ligase. Using this protocol, the available restriction enzymes depend on the nucleotide sequences of target genes, and different enzymes should be used when many target genes are cloned in parallel.

Ligation-independent cloning (LIC) has been developed as a new cloning method, which eliminates the use of restriction endonuclease digestion and ligation of PCR products [1,2]. LIC circumvents the limitations of traditional gene cloning methods because any PCR products can be inserted into LIC-compatible cloning vectors without restriction enzymes and T4 DNA ligase. In LIC, PCR primers are designed to have LIC-compatible 5’ extension sequences. PCR products and vectors are treated with T4 DNA polymerase in the presence of a single deoxyribonucleotide triphosphate, which generates specific 12–20 nucleotide single stranded overhangs. The PCR products are then annealed with vectors which have complementary overhangs, and transfor-
purification of human α-ketoglutarate dehydrogenase (KGDH) subunits (E1, E2, E3) in silkworm larvae showed the feasibility of using LIC vectors. Moreover, an improved LIC vector, allowing the fusion of two expression plasmids into one was also constructed for expression of a protein complex based on the ‘LINK’ sequence [11,12]. A newly created ‘LINK’ sequence was incorporated into the improved LIC vector and used for successful co-expression of human microsomal triglyceride transfer protein (hMTP) and human protein disulfide isomerase (hPDI) protein complex.

Materials and Methods

Construction of LIC vectors for the expression of recombinant proteins fused with affinity tag C-terminally

Plasmid construction is illustrated in Figures 1 and 2. To construct various LIC vectors, LIC-specific forward (Cassette F) and reverse primers (Cassette R) were designed as shown in Figure 1A. Cassette F contains the LIC sequence, the thrombin cleavage site, and the 5′-sequence of the EGFP gene. Cassette R contains the 6× histidine tag, Strep-tag II sequences, and 3′-sequence of the EGFP gene. PCR was performed using these primers, and the EGFP gene was used as a template. The PCR product treated with XhoI and KpnI was inserted into the XhoI–KpnI site of pFastBac1 (Invitrogen, San Diego, CA, USA). The constructed plasmid was named pFastEGFP-SH-LIC (Figure 1B). EGFP was removed from pFastEGFP-SH-LIC by NheI and self-ligation, and pFast-S-H-LIC was produced. Digestion of pFast-SH-LIC by MluI and PvuII, respectively, and self-ligation, produced pFast-S-LIC and pFast-H-LIC.

Two modified LINK sequences allowing the joining of plasmids in tandem were also designed [12]. gcLINK1 and gcLINK2 include the GC rich recognition sites for the restriction enzyme AscI (gcLINK1) or AscI and NcoI (gcLINK2) (Figure 2B). The construction of tandem plasmids allows the co-expression of two subunits of a protein complex [11,12]. Two expression cassettes can be in tandem connected in one vector through gcLINK1 and 2 sequences. gcLINK-specific primers (gcLINK1-F, gcLINK2-R) (Figure 1A) were used to insert gcLINK sequences into pFast-S-LIC and pFast-H-LIC. gcLINK1-F and gcLINK2-R primers have gcLINK1 and gcLINK2 sequences, respectively. The region containing the polyhedrin promoter, LIC sequence, and the SV40 polyadenylation signal region was amplified using these primers and pFast-S-LIC or pFast-H-LIC as a template. Each amplified fragment treated by BstGI and XhoI was inserted into the BstGI–XhoI site of pFastBac1. Thus, pFastgcLINK-S-LIC and pFastgcLINK-H-LIC were constructed (Figure 1B).

LIC reaction

Plasmid construction by LIC reaction is also described in Figure 2. pFast-S-LIC and pFast-H-LIC were treated with SmaI and incubated at 65°C for 20 min to inactivate enzyme activity. T4 DNA polymerase (2.5 U) (Fermentas INC, Glen Burnie, MD, USA), which has both activities of 5′-3′ polymerase and 3′-5′ exonuclease, and 2.5 mM dTTP, were added into reaction mixtures and incubated at 25°C for 30 min, followed by heating at 65°C for 20 min to inactivate T4 DNA polymerase. Because of the 3′-5′ activity of polymerase the nucleotides (“A”, “G” and “C”) are removed from 3′-ends until the first “T” residue is reached. When T4 DNA polymerase comes to this first “T”, its polymerase reaction dominates and its 3′-5′ exonuclease reaction stops. This reaction leads two specific 5′-overhangs in this LIC vector of 10 and 12 bases, respectively.

hMTP and hPDI genes were amplified by PCR using Human MTP F (5′-CGAAGAGGAGGCTCACTAT- GATTCTTCTTGGTGTG 3′), Human MTP R (5′ GGCAAGCAGACCGCTAACCAGGCTTGAAGT 3′), Human PDI F (5′- CGAAGAGGAGGCTCACTATGCTGGCGCCGGGTCTGTG 3′), and Human PDI R (5′ GCACCGAGACCGCTAACCAGGCTTGAATC 3′) primers, respectively. The ER retention sequence (last 4 amino acids of hPDI) was removed from the intact hPDI sequence. Amplified genes were purified and treated with T4 DNA polymerase according to the protocol described above. Then, T4 DNA polymerase-treated hMTP and hPDI were mixed with T4 DNA polymerase-treated pFastgcLINK-S-LIC and pFastgcLINK-H-LIC, respectively, and heated at 65°C for 5 min. The mixtures were then cooled to room temperature for 1 h to anneal each other. Two microliters of 25 mM EDTA was added to each mixture before E. coli DH10B (Invitrogen) transformation. The constructed plasmids were named pFastgcLINK-hMTP-S and pFastgcLINK-hPDI-H. Applying the same protocol described above, pFastBac-hMTP-EGFP-SH was constructed using pFastEGFP-SH-LIC and LIC genes amplified by PCR.

To connect pFastgcLINK-hMTP-S with pFastgcLINK-hPDI-H, pFastgcLINK-hMTP-S and pFastgcLINK-hPDI-H were digested by NcoI and AscI, respectively, and each digested plasmid was treated with T4 DNA polymerase in the presence of 2.5 mM dATP or dTTP. NcoI-Digested pFastgcLINK-hMTP-S and an ∼ 2.8 kbp AscI-digested fragment including the hPDI gene were purified. Overhang sequences of LINK1 and 2 regions after T4 DNA polymerase treatment are shown in Figure 2B. Each purified fragment was mixed, heated, and then cooled at room temperature to anneal each other at the complementary sequences. Two microliters of 25 mM EDTA was added to each mixture before E. coli DH5α transformation. The constructed plasmids were designated as pFastgcLINK-hMTP-S/hPDI-H (Figure 2B).

Construction of LIC vectors for the expression of recombinant proteins fused with an affinity tag in N-terminal

A LIC cassette for the expression of recombinant proteins fused with an affinity tag in the N-terminal was amplified by PCR using primers F2 and R2, and pFast-SH-LIC, which was constructed in this study as a template (Figure 3). An amplified fragment containing the Start codon, Strep Tag II sequence, 6× histidine tag sequence, TEV protease cleavage site, and LIC site (SpI) was ligated into the XhoI–KpnI site in pFastBac1. gcLINK1 and gcLINK2 sequences were inserted into the resulting plasmid (pFast-N-SH-LIC) by the same manner as described in “Construction of LIC vectors for the expression of recombinant proteins fused with affinity tag C-terminally” in Materials and Methods. The resulting plasmid, pFastgcLINK-N-SH-LIC, had 5 SpI sites.
Rapid Expression of Proteins Using LIC Vector

(A) pFastgclINK-S-LIC

LIC reaction

hMTP

NarI

pFastgclINK-hMTP-S

(B) pFastgclINK-hMTP-S/hPDI-H

LIC reaction

hMTP

Ascl

pFastgclINK-hPDI-H

(C) pFastgclINK-hMTP-S

LIC annealing sequence

Start codon

LIC annealing sequence

5’ CGGAGGGGAGGCCCTACCATGATTTCTTCTTCTGCCTGGTTAGGCTGGCTTGCA...

hMTP gene

Thrombin cleavage site
Four of the Sfl sites in this vector, except for that in the LIC sequence, were mutated by using the Quickchange Lightning Multi site-Directed Mutagenesis Kit (Agilent Technologies, Inc., Santa Clara, CA, USA) using the following primers (pF-sp441: 5’ GGAATTTTACAAAGATTAACGTTTATAAT 3’, pF-sp5572: 5’ GATAAATGCTTACAAAGATGAAAGGAGAAGTAA 3’, pF-sp3986: 5’ CTGTATTCCGGAAAGATTAATAGATCGATCCGATTGATGCTG 3’, pF-sp3996: 5’ TTTTTTTTTTTTTTTTTTGTATTCCGGAAAGATTAATAGATCGATCCGATTGATGCTG 3’, E2-R: 5’ TACTTCCAATCCAATGCTTTGCTTGTTGAGGCCC 3’, E2-F: 5’ TCTTACCCATGCACTGACATTGGAGCATAATG 3’, E3-F: 5’ TGTACCTTCAATGCACTGACATTGGAGCATAATG 3’, E3-R: 5’ TGTACCTTCAATGCACTGACATTGGAGCATAATG 3’). Each amplified gene was inserted into pFMgcLINK-F-LIC vector. Then, pFMgcLINK-S-E1, pFMgcLINK-H-E2, and pFMgcLINK-N-S-LIC, pFMgcLINK-N-H-LIC, and pFMgcLINK-N-S-H-LIC, pFMgcLINK-N-H-LIC, were constructed. Recombinant BmNPV bacmid DNA (10 μg) containing helper plasmid was mixed with a one-tenth volume of DMRIE-C (Invitrogen) reagent and this mixture was injected into silkworm larvae or pupae.

Expression and purification of expressed proteins from fat bodies of silkworm larvae or pupae injected with recombinant BmNPV bacmid DNA

hMTP-hPDI complex. The fat bodies collected from 10 silkworm pupae were suspended in 25 mL of phosphate-buffered saline (PBS, pH 7.4) and sonicated 3 times for 30 sec each time with 1 min intervals. After centrifugation of the resultant homogenate at 8000 x g for 10 min, the collected supernatant was mixed with 0.5 mL of PBS-equilibrated TALON Metal Affinity Resin (Clontech, Mountain View, CA, USA) and stirred at 4°C for 1 h. This mixture was centrifuged at 3000 x g for 2 min, and the precipitated resin was washed with 120 mL of PBS. Proteins bound to resin were eluted with 2 mL of PBS containing 200 mM imidazole. This eluent was mixed and gently stirred with 0.25 mL of PBS-equilibrated Streptactin Superflow (QIAGEN K.K., Tokyo, Japan) at 4°C for 1 h. This mixture was centrifuged at 3000 x g for 2 min and the supernatant was removed. After washing the affinity gel with 5 mL of PBS, hMTP-hPDI complex was eluted with 2 mL of PBS containing 5 mM desthiobiotin.

S-E1. The homogenate supernatant was obtained from the collected fat bodies of 5 silkworm larvae by the same manner as the purification of hMTP-hPDI complex. Supernatant was mixed with 0.5 mL of Strept-Tactin Superflow equilibrated with Tris-buffered saline (TBS, pH 7.5) and gently stirred at 4°C for 1 h. This mixture was centrifuged at 3000 x g for 2 min and the supernatant was removed. After washing the affinity gel with 120 mL of TBS, S-E1 was eluted with 2 mL of TBS containing 5 mM desthiobiotin.

H-E2. H-E2: Homogenate supernatant was obtained from fat bodies collected from 5 silkworm larvae by the same manner as the purification of hMTP-hPDI complex. Supernatant was mixed with 0.5 mL of TALON Metal Affinity Resin equilibrated with TBS and gently shaken at 4°C for 1 h. This mixture was centrifuged at 3000 x g for 2 min and the supernatant was removed. After washing the affinity gel with 120 mL of TBS, H-E2 was eluted with 2 mL of TBS containing 200 mM imidazole.

F-E3. Homogenate supernatant was obtained from fat bodies collected from 5 silkworm larvae by the same manner as the purification of hMTP-hPDI complex. Supernatant was mixed with 0.5 mL of anti-FLAG M2 antibody agarose (Sigma Aldrich Japan, Tokyo). This mixture was centrifuged at 3000 x g for 2 min and the supernatant was removed. After washing the affinity gel with 120 mL of TBS, F-E3 was eluted with 2 mL of TBS containing 100 μg/mL of FLAG peptide (Sigma Aldrich Japan).

SDS-PAGE and protein analysis

Recombinant protein samples were subjected to SDS-PAGE on 10 or 12% polyacrylamide gels using the Mini-protean II system (Bio-Rad Co. Ltd, Hercules, CA, USA). Total proteins on SDS-PAGE gel were detected with Coomassie Brilliant Blue (CBB) R-250 or silver staining. For the specific detection of fluorescent hMTP-EGFP fusion protein on SDS-PAGE gels, samples were mixed only with sample buffer and not boiled [14]. Fluorescent bands were detected using Molecular Imager FX (Bio-Rad).
(A) Cassette F2: 5' GCCTCTAGACACATGCTGTCCTGCACCCCG 3'
KpnI SspI MluI
Cassette R2: 5' GGGGTACGCTCCCTGACTCCTTTGTACGGTGCTGGAGGAAGCTG 3'
LIC site
TEV protease cleavage site
6 x His tag

(B) Cassette F2 
6 x His tag 
LIC site 
Start codon 
PvuII 
SspI 
MluI 

Cassette R2: 

XbaI, KpnI 
Ligation 
PvuII 
PvuII 
SspI 

XbaI, KpnI 
Ligation 
PvuII 
PvuII 

PCR 

XbaI, KpnI 
Ligation 
PvuII 
PvuII 

Mutation at four SspI sites in pFastBac vector 

(pFastN-SH-LIC) 

(gcLINK1) 

(gcLINK2) 

MluI 
PvuII 
Ligation 

MluI 
PvuII 
Ligation 

MluI 
PvuII 
Ligation 

MluI 
PvuII 
Ligation 

Insertion of each gene by LIC 

(FMgcLINK-S-E1) 

(FMgcLINK-H-E2) 

(FMgcLINK-F-E3) 

(C) FMgcLINK-S-E1

LIC annealing sequence
5' TACTCCCAATCCATGACACCTGCTGAAGCC 3'

Stop codon

LIC annealing sequence
5' GTCTCTGAACTCTCTGAGGGATAAGTTGAAGCC 3'

TEV protease cleavage site

E1 gene (41-1023 a.a.)
Fluorescent fusion proteins that were not boiled showed a slightly different molecular mass on SDS-PAGE gels as compared to boiled samples.

Protein concentration was determined using Protein Assay Kit II (Bio-Rad) based on the Bradford method.

Dialysis was performed in PBS overnight at 4°C using a dialysis membrane (WAKO Pure Chem. Ind., Ltd. Osaka, Japan).

Dihydrolipoamide dehydrogenase (DLD) activity was assayed according to Huo et al [15]. In brief, 1 mL of assay mixture [0.5 mM lipoamide, 0.1 mM NADH, 1 mM EDTA, 50 mM potassium phosphate (pH 6.7)] was incubated at room temperature, and 10 µL of enzyme was added to initiate the enzyme reaction. The decrease of absorbance at 340 nm was measured, and enzyme activity was calculated using the extinction coefficient of 6.22 mM cm⁻¹ for NADH. One unit of enzyme activity was defined as the amount of enzyme capable of catalyzing the conversion of 1 µmol of NADH to NAD⁺ per minute.

Results and Discussion

Expression of hMTP-EGFP fusion protein using pFast-SH-LIC vector

The hMTP-hPDI complex was expressed in silkworm larvae using newly constructed LIC vectors. hMTP is located in the lumen of endoplasmic reticulum (ER) as a heterodimer with human protein disulfide isomerase (hPDI) [16,17]. This heterodimer catalyzes the transfer of triacylglycerol cholesteryl ester to phospholipid, and is involved in the assembly and secretion of very-low-density lipoprotein and chylomicrons. Bovine MTP was co-purified with bovine PDI from bovine liver homogenate [18]. Moreover, the hMTP was precipitated when expressed solely in insect cells, but it was solubilized and purified by only co-expression with hPDI [19].

pFastBac-EGFP-SH-LIC contains the EGFP gene, Strep-tag II, and 6 histidine tag sequences downstream of the LIC region (SmaI site). EGFP gene, Strep-tag II, and 6 histidine tag sequences can be removed by thrombin due to the presence of a thrombin cleavage site between the LIC site and these tag sequences. The hMTP gene could be inserted with high efficiency at the LIC site by using the LIC reaction (identified 5 colonies that all had the hMTP gene-inserted vector). The LIC reaction is highly specific and the efficiency of its reaction is ~100% [1,2,9]. Moreover, the hMTP-EGFP-SH gene sequence was confirmed to be correct in this plasmid. To confirm that hMTP-EGFP-SH fusion protein could be expressed in-frame using this vector, a recombinant BmNPV-CP-hMTP-EGFP-SH bacmid was prepared and injected into silkworms and pupae. Green fluorescence was detected in silkworm larvae and pupae (Figure 4A). Fat bodies collected from larvae showing green fluorescence were suspended with PBS and sonicated to extract the expressed hMTP-EGFP fusion protein. A specific GFP fluorescent band on SDS-PAGE was observed in the homogenate of the bacmid-injected larva’s fat body compared to that of a mock injected fat body (Figure 4B). A non-specific fluorescent band was also observed in both homogenates, indicating that this band was from proteins inherent in silkworm larvae. These results indicate that the pFastBac-EGFP-SH vector works well and allows the expression of EGFP fusion protein.

Figure 3. Sequences of primers used for amplification of LIC cassette containing start codon, Strep-tag II sequence, 6× histidine tag sequence, TEV protease cleavage site sequence, and LIC site for N-terminal tagged protein expression, and scheme for LIC construction vectors. (A) LIC cassette for N-terminal tagged protein expression was amplified using primer F2 and primer R2. Amplified LIC cassette has start codon, Strep-tag II sequence, 6× histidine tag sequence, TEV protease cleavage site sequence, and SspI site in the LIC site. (B) pFMgcLINK-S-E1, pFMgcLINK-H-E2, and pFMgcLINK-F-E3 were constructed for each KGDH subunit expression in silkworms. (C) The sequence of E1 gene in pFMgcLINK-S-E1 vector is shown. E1 gene was successfully inserted into the LIC site in-frame in this vector. The sequences of E2 and E3 genes in constructed vectors are not shown, but these genes were also successfully inserted in-frame into the vector.

doi:10.1371/journal.pone.0064007.g003

Figure 4. Expression of hMTP-EGFP fusion protein in silkworm larvae and EGFP fluorescence analysis of hMTP-EGFP fusion protein. (A) Expression of hMTP-EGFP fusion protein in silkworm larvae and pupae. (B) EGFP fluorescence analysis of hMTP-EGFP fusion protein in homogenate of silkworm larvae fat body on a SDS-PAGE gel. Lane 1: fat body of mock, lane 2: fat body of larvae into which bacmid DNA harboring hMTP-EGFP fusion gene was injected.

doi:10.1371/journal.pone.0064007.g004
Construction of the co-expression vector of hMTP and hPDI

Co-expression vectors have been improved in the baculovirus expression system [20,21], also in silkworms [22], but a digestion process using restriction enzymes is absolutely needed for cloning the genes of interest. In this study, a co-expression vector that does not require an enzymatic ligation reaction for gene cloning was constructed.

hMTP and hPDI genes were inserted using the LIC reaction into constructed pFastgcLINK-S-LIC and pFastgcLINK-H-LIC, respectively, (Figure 1) and then pFastgcLINK-hMTP-S and pFastgcLINK-hPDI-H were constructed as shown in Figure 2A. The efficiency of inserting hMTP and hPDI genes into pFastBac-S-LIC and pFastBac-H-LIC, respectively, was ~100%. Sequences of hMTP connected with Strep-tag II (hMTP-S) and hPDI connected with 6× histidine tag (hPDI-H) were confirmed without any insertion and deletion of nucleotides (Figure 2C, only hMTP-S sequence is shown). pFastgcLINK-hMTP-S and pFastgcLINK-hPDI-H were connected by the LIC reaction as shown in Figure 2A. Normal and small sized transformants appeared after E. coli transformation of the LIC reaction. pFastgcLINK-hMTP-S/hPDI-H, which harbors hMTP-S and hPDI-H genes, was obtained from small sized transformants. pFastgcLINK-hMTP-S/hPDI-H was obtained from all 4 small transformants, while pFastgcLINK-hMTP-S was obtained from all 4 normal transformants. The ratio of small transformants in all transformants was ~50%; however, pFastgcLINK-hMTP-S/hPDI-H was obtained with 100% efficiency. A recombinant BmNPV bacmid for co-expression of hMTP-S and hPDI-H was constructed using pFastgcLINK-hMTP-S/hPDI-H.

Co-expression of hMTP-hPDI complex in silkworm larvae and pupae using constructed co-expression vector

A recombinant BmNPV-CP--hMTP-S/hPDI-H bacmid harboring hMTP-S and hPDI-H genes was injected to 10 pupae and silkworm larvae and reared for 6 d. The collected bacmid-injected pupae were suspended with PBS and homogenized by sonication. hMTP-hPDI complex was purified by TALON affinity resin and Strep-Tactin agarose because hMTP and hPDI were connected with Strep-tag II and 6× histidine tag, respectively. Two protein bands appeared after Strep-Tactin agarose chromatography (Figure 5A). The molecular weight of each band (hMTP-S and hPDI-H) was estimated to be ~100 and 58 kDa, respectively.

Table 1. Purification of dihydrolipoamide dehydrogenase expressed in fat body of silkworm larvae.

|Volume (ml) | DLD activity (U) | Protein concentration (mg) | Specific activity (U/mg) | Purification (-fold)* | Recovery (%) |
|------------|------------------|---------------------------|-------------------------|----------------------|-------------|
|Fat body homogenate 20 | 0.32 | 121.00 | 2.64×10^3 | 1 | 100 |
|Anti-FLAG agarose 2 | 0.42 | 0.13 | 3.23 | 1224 | 130 |

*Defined by dividing specific activity by that of fat body homogenate.
doi:10.1371/journal.pone.0064007.t001
from each amino acid sequence. These values corresponded to 2 bands observed on a SDS-PAGE gel stained with silver staining. Moreover, no bands were observed in the Strep-Tactin agarose eluent except for hMTP-S and hPDI-H (Figure 3A). This indicates that hMTP-hPDI complex could be highly purified by TALON affinity resin and Strep-Tactin agarose. hMTP forms inclusion bodies when solely expressed in the baculovirus expression system [19], but hMTP formed the soluble complex with hPDI in pupae by co-expression with hPDI. This phenomenon is similar to a previous report on improvement of IgG solubility by co-expression of BiP in Trichoplusia ni cells using a baculovirus expression system [23]. Thirteen micrograms of purified hMTP-hPDI complex was obtained from 10 silkworm pupae. This value is less than that of human prorenin-human prorenin receptor complex (70 µg/15 silkworm larvae) reported previously [24]. However, the hMTP-hPDI complex from hemolymph of BmNPV-CP--hMTP-S/hPDI-H bacmid DNA-injected silkworm larvae couldn’t be purified. This suggests that hMTP-hPDI complex resides only in the endoplasmic reticulum (ER), and therefore, the yield of purified hMTP-hPDI complex was low compared to the yield of human prorenin-human prorenin receptor complex. To improve the recovery of purified hMTP-hPDI complex, purification by Strep-Tactin agarose was performed after dialysis of the hMTP-hPDI complex eluent of TALON affinity chromatography against PBS. Recovery yield (10 µg/10 silkworm pupae) was constant irrespective of dialysis (Figure 5B). When high-throughput purification of many recombinant proteins is performed in parallel using 2 purification steps, dialysis after the first chromatography is the rate-determining step. This affinity chromatography combination (6× histidine and Strep-tag II tags) is very useful for high-throughput protein purification because the dialysis step between TALON affinity and Strep-Tactin agarose chromatography can be omitted.

Expression and purification of each human KGDH subunit in silkworm larvae using LIC vectors constructed for N-terminal tagged fused protein expression

Human KGDH is in the mitochondrial matrix and is a multi-enzyme complex in the TCA cycle. KGDH is composed of multi-copies of E1 (α-ketoglutarate dehydrogenase), E2 (dihydrolipoamide succinyltransferase), and E3 (dihydrolipoamide dehydrogenase). KGDH plays an important role in controlling the reductive potential (NADH/NAD+) in mitochondria [25]. Moreover, KGDH is a rate-limiting enzyme in the TCA cycle and may be involved in neurodegenerative disease [26]. Thus, expression of each KGDH subunit was investigated in silkworm larvae using LIC vectors without its mitochondrial targeting signal.

LIC vectors for the expression of recombinant proteins fused with an affinity tag at the N-terminus were also constructed as shown in Figure 3. The LIC cassette was initially amplified by PCR. The amplified fragment has a start codon, 6× histidine tag sequence, Strep-tag II tag sequence, TEV protease cleavage site sequence, and a LIC site. This LIC site sequence has the 5′- and 3′-UTR regions and a LIC site, which is inserted into LIC vectors constructed in this study, also has a 4 SspI site. Then, 4 SspI sites in the constructed pFastgcLINK-N-SH-LIC vector, except for the 5′-UTR, were mutated so as not to be digested by SspI. LIC vectors for each N-terminally tagged (6× histidine tag, Strep-tag II) protein, pFMgcLINK-N-S-LIC and pFMgcLINK-N-H-LIC, were constructed in the same manner as shown in Figure 1. Moreover, the FLAG tag sequence was replaced with a 6× histidine tag and Strep-tag II module at the MuI and PvuII sites, and pFMgcLINK-N-F-LIC was constructed, indicating that any tag can be inserted into this site instead of the 6× histidine tag and Strep-tag II. This replacement can be available to LIC vectors constructed in Figure 1 for C-terminal tag and protein expression, or the pFastgcLINK-N-S-LIC or pFastgcLINK-H-LIC vector. To confirm the integrity of these vectors, each human KGDH subunit (E1, E2, E3) was inserted into pFMgcLINK-N-S-LIC, pFMgcLINK-N-H-LIC, and pFMgcLINK-N-F-LIC, respectively, using the LIC reaction. The efficiency of inserting the human KGDH subunit (E1, E2, E3) into pFastgcLINK-N-S-LIC or pFastgcLINK-H-LIC vectors was ~100%, and each gene was inserted into each vector in-frame (Figure 3C, only E1 sequence is shown). Each recombinant bacmid was constructed using these vectors, and each subunit was expressed in silkworm larvae and purified from its fat body. Because each subunit lacks its mitochondria targeting signal-coding sequence, each subunit was purified from the supernatant of homogenate by affinity gel chromatography as almost a single band at an estimated molecular weight (E1: 110 kDa, E2: 48 kDa, E3: 55 kDa); except that some amount of purified E3 was degraded and had slightly higher mobility on an SDS-PAGE gel (Figure 6.). E3 is a DLD and DLD activity was assayed using purified E3. E3 was purified from the fat body homogenate by ~1224-fold, and the specific activity of purified E3 was 3.2 U/mg (Table 1). In this study, 55, 30, 130 µg of purified E1, E2, and E3 were obtained from 5 bacmid-injected silkworm larvae, respectively. This verified that LIC vectors pFMgcLINK-N-S-LIC, pFMgcLINK-N-H-LIC, and pFMgcLINK-N-F-LIC also work well for protein expression and purification in silkworms.

Conclusions

LIC and co-expression vectors were constructed using a new gcLINK sequence and BmNPV bacmid. Using these vectors, an hMTP-hPDI co-expression vector was successfully constructed without ligation. Moreover, purification of hMTP-hPDI complex was easily performed by TALON affinity and Strep-Tactin agarose chromatography, because Strep-tag II and 6× histidine tag were fused with hMTP and hPDI, respectively. Moreover, LIC vectors for the expression of recombinant proteins fused with several tags in the N- or C-terminal end were constructed. Each human KGDH subunit was expressed in silkworm larvae and purified from its fat body. This indicates that LIC vectors constructed in this study were available for protein expression and purification in silkworms. Because these vectors facilitate high-efficiency cloning of genes and fusion with purification tags, they are applicable for high-throughput expression of proteins, co-expression of protein complexes, and simultaneous administration of gene expression in the silkworm expression system.

Author Contributions

Conceived and designed the experiments: TK JRT. Performed the experiments: TK JRT EYP. Analyzed the data: TK JRT EYP. Contributed reagents/materials/analysis tools: JRT EYP. Wrote the paper: TK.

References

1. Aalnæs M, Winters-Hauser R, Krogstad P (1999) Ligation-independent cloning of PCR products (LIC-PCR). Nucleic Acids Res 18: 6069–6074.
2. Haun RS, Moss J (1992) Ligation-independent cloning of glutathione S-transferase fusion genes for expression in Escherichia coli. Gene 112: 37–43.
3. Weeks SD, Drinker M, Loll PJ (2007) Ligation independent cloning vectors for expression of SUMO fusions. Protein Expr Purif 53: 40–50.
4. Lee J, Kim SH (2009) High-throughput T7 LIC vector for introducing C-terminal poly-histidine tags with variable lengths without extra sequences. Protein Expr Purif 65: 58–61.

5. Chanda PK, Edris WA, Kennedy JD (2006) A set of ligation-independent expression vectors for co-expression of proteins in *Escherichia coli*. Protein Expr Purif 47: 217–224.

6. Qin H, Hu J, Hua Y, Challa SV, Cross TA, et al. (2008) Construction of a series of vectors for high throughput cloning and expression screening of membrane proteins from *Mycobacterium tuberculosis*. BMC Biotechnol 8: 51.

7. Cabrita LD, Dai W, Bottomley SP (2006) A family of *E. coli* expression vectors for laboratory scale and high throughput soluble protein production. BMC Biotechnol 6: 12.

8. Alzari PM, Berglund H, Berrose NS, Blagova E, Busso D, et al. (2006) Implementation of semi-automated cloning and prokaryotic expression screening: the impact of SPINE. Acta Crystallogr D Biol Crystallogr 62: 1103–1113.

9. Bardocz V, Geczi V, Sawasaki T, Endo Y, Meszaros T (2008) A set of ligation-independent in vitro translation vectors for eukaryotic protein production. BMC Biotechnol 8: 32.

10. Li M, Hays FA, Roe-Zurz Z, Vuong L, Kelly L, et al. (2009) Selecting optimum eukaryotic integral membrane proteins for structure determination by rapid expression and solubilization screening. J Mol Biol 385: 620–630.

11. Alexandrov A, Vignali M, LaCount DJ, Quartley E, de Vries C, et al. (2004) A facile method for high-throughput co-expression of protein pairs. Mol Cell Proteomics 3: 934–938.

12. Schirch C, Kummer D, Sounalakakis D, Heinenmann U, Busso K (2007) Vectors for co-expression of an unrestricted number of proteins. Nucleic Acids Res 35: e43.

13. Motohashi T, Shimojima T, Fukagawa T, Maenaka K, Park EY (2005) Efficient large-scale protein production of larvae and pupae of silkworm by *Bombyx mori* nuclear polyhedrosis virus bacmid system. Biochem Biophys Res Commun 326: 564–569.

14. Aoki T, Takashashi Y, Koch KS, Leffert HL, Watabe H (1996) Construction of a fusion protein between protein A and green fluorescent protein and its application to Western blotting. FEBS Lett 384: 193–197.

15. Huo J, Shi H, Yao Q, Chen H, Wang L, et al. (2010) Cloning and purification of recombinant silkworm dihydrolipoamide dehydrogenase expressed in *Escherichia coli*. Protein Expr Purif 72: 95–100.

16. Wetterau JR, Zilversmit DB (1983) Purification and characterization of microsomal triglyceride and cholesterol ester transfer protein from bovine liver microsomes. Chem Phys Lipids 30: 205–222.

17. Wetterau JR, Comb KS, Spinnet SN, Joiner BJ (1990) Protein disulfide isomerase is a component of the microsomal triglyceride transfer protein complex. J Biol Chem 265: 9800–9807.

18. Lamberg A, Jauhiainen M, Mesto J, Ehnholm C, Shoulders C, et al. (1996) The role of protein disulfide isomerase in the microsomal triglyceride transfer protein does not reside in its isomerase activity. Biochem J 315: 533–536.

19. Ritchie PJ, Decourt A, Amey J, Maun CJ, Read J, et al. (1999) Baculovirus expression and biochemical characterization of the human microsomal triglyceride transfer protein. Biochem J 338: 305–310.

20. Aricescu AR, Asenovorg R, Bill RM, Busso D, Chang VT, et al. (2006) Eukaryotic expression: developments for structural proteomics, Acta Crystallogr D Biol Crystallogr 62: 1114–1124.

21. Trowitzsch S, Bieniossek C, Nie Y, Garzoni F, Berger I (2010) New baculovirus expression tools for recombinant protein complex production, J Struct Biol 172: 45–54.

22. Zhou Y, Chen H, Li X, Wang Y, Chen K, et al. (2011) Production of recombinant human DNA polymerase delta in a *Bombyx mori* bioreactor. PLoS ONE 6: e22224.

23. Hsu T, Betenbaugh MJ (1997) Coexpression of molecular chaperone BiP improves immunoglobulin solubility and IgG secretion from *Trichoplusia ni* insect cells, Biotechnol Prog 13: 96–104.

24. Du D, Kato T, Suzuki F, Park EY (2009) Expression of protein complex comprising the human protein and [pro]tein receptor in silkworm larvae using *Bombyx mori* nucleopolyhedrovirus (BmNPV) bacmid for improving biological function, Mol Biotechnol 43: 154–161.

25. McLain AL, Szweda PA, Szweda LI (2011) α-Ketoglutarate dehydrogenase: A mitochondrial redox sensor. Free Radic Res 45: 29–36.

26. Shi Q, Xu H, Yu H, Zhang N, Ye Y, et al. (2011) Inactivation and reactivation of the mitochondrial α-ketoglutarate dehydrogenase complex. J Biol Chem 286: 17640–17648.