Soluble Expression of Humanized Anti-CD20 Single Chain Antibody in *Escherichia coli* by Cytoplasmic Chaperones Co-expression

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**Abstract**

**Background:** CD20 is an important cell surface receptor that is used for target therapy of B cell lymphoma and some related blood diseases due to vital function of CD20. In previous studies, a Rituximab based humanized single chain variable fragment (scFv) antibody showed good reactivity against B cell related cancer cells. But this recombinant protein produced Inclusion Bodies (IBs) in *Escherichia coli* (E. coli) cytoplasm. The aim of this study was to investigate the effect of coexpression with cytoplasmic chaperones on expression and solubility of humanized anti-CD20 scFv in *E. coli*.

**Methods:** For this purpose, the fragment coding for anti-CD20 huscFv subcloned into the pET22b (+) and transformed into the E. coli BL21 (DE3) was evaluated. In order to inhibit the production of IBs, the effects of co-expression with cytoplasmic chaperones GroEL, DnaK, GroES, Tig, DnaJ and GrpE were investigated.

**Result:** Coexpression with cytoplasmic chaperones led to increased soluble expression of anti-CD20 recombinant protein. Among investigated chaperones, pKJE7 chaperone plasmid containing DnaJ, GrpE, DnaK chaperone genes had significant effects with an expression yield of 325 μg/ml soluble anti-CD20 scFv.

**Conclusion:** The result of this study demonstrated remarkable effect of pKJE7 chaperone on enhancement of soluble expression of anti-CD20 huscFv antibody in *E. coli*.

**Keywords:** Molecular chaperones, Non Hodgkin lymphoma, Single chain antibody

**Introduction**

CD20 is a cell surface marker that is expressed on B cell and presented during the stage of pre B cells. CD20 directly converts the phospholipase C-gamma (PLCγ) to active Ca2+ transport pump across the plasma membrane 1. Non Hodgkin Lymphoma (NHL) is one of the malignant diseases that is associated with overexpression of CD20 on B cell surface. Hence, anti-CD20 antibodies are approved as an effective treatment method for targeted therapy of B cell malignancies. Rituximab is an anti-CD20 monoclonal antibody (mAb) 2 that is utilized for treating NHL often culminating in long-lasting response 3. In recent years, various formats of mAb have been developed, such as Fragment antigen-binding (Fab), single chain variable fragments (scFv) and disulfide-stabilized Fv antibody fragment (dsFv). Among them, the scFv format has been extensively studied for various applications 4. Recombinant scFv is generated by joining VH and VL domains through a short polypeptide linker or disulphide bond 5 and *Escherichia coli* (*E. coli*) are generally used as host cells 6-8. IBs are composed of proteins with inappropriate structures. To overcome this problem, various strategies have been developed including the use of molecular and chemical chaperones 9,10.

Molecular chaperones are now well established as cytoplasmic and periplasmic chaperones 11,12. GroEL, DnaK, GroES, Tig, DnaJ and GrpE are major cytoplasmic chaperones in *E. coli*. It has been shown that cytoplasmic chaperones inhibit the rate of Inclusion Bodies (IB) formation leading to increase in solubility of recombinant proteins 13,14. The cytoplasmic chaperones are generally expressed in derivatives of pACYC.
Soluble Expression of Anti-CD20 scFv in E. coli

Materials and Methods

Bacterial strains, media, and plasmids

E. coli DH5α, BL21 (DE3) and expression vector pET22b (+) were purchased from Novagen. IPTG, Taq DNA polymerase was obtained from Promega Company. Restriction enzyme and T4 DNA ligase were purchased from Takara Shuzo (Kyoto, Japan). The pUC57-huscFv construct was obtained from V. Ahmadzadeh. The chaperon plasmid set was purchased from TAKARA Bio Inc. (Table 1). Also, the Raji cell line, Ni-NTA resin and protein size marker were obtained from Pasteur Institute of Iran, Qiagen and Fermentas, respectively.

Cloning and expression of humanized single chain antibody

The pUC57 containing target gene (pUC57-hscFv construct) was double digested by restriction enzymes MslI (MscI) and XhoI (at 5’ and 3’ ends, respectively). The insert (huscFv) was isolated and sub-cloned into the pET22b expression vector containing pelB leader sequence to facilitate expression of recombinant protein in the periplasmic space. For confirmation of this recombinant construct (pET22b-huscFv), PCR was used and restriction analysis and sequencing were carried out.

PET-22b-hscFv construct was transformed to BL21(D3) and was grown in LB medium complemented with 100 μg/ml ampicillin. The culture then was incubated with shaking (150 rpm) at 25°C until culture was induced at A600=0.7 using 1 mM IPTG for 4 hr. Finally, protein expression was analyzed by SDS-PAGE followed by visualization with Coomassie Brilliant Blue staining and quantitative analysis by using image processing program called ImageJ (National Institute of Health, Bethesda, MD).

Soluble expression of huscFv by co-expression of pET22b-huscFv with chaperones plasmids

For soluble expression of huscFv, E. coli BL21 (DE3) containing pET22b-huscFv bacterial cells were transformed with various chaperone plasmids. Five chaperones including plasmids pG-KJE8, pGro7, pKJE7, pG-T12 and pPT16 were used. The results from these transformations give the BL21(DE3)/pET22b-huscFv/pG-KJE8, BL21(DE3)/pET22b-huscFv/pGro7, BL21(DE3)/pET22b-huscFv/pKJE7, BL21(DE3)/pET22b-huscFv/pPT16, BL21(DE3)/pET22b-huscFv/pT16. The transformed cells were cultured in 100 ml LB medium containing 20 μg/ml chloramphenicol and 50 μg/ml ampicillin. Induction of cells containing chaperones pGro7, pKJE7 and pG-T12 was carried out by L-arabinose, pG-KJE8 by L-arabinose and tetracycline, and pPT16 by tetracycline. For expression of chaperones related inducers, L-arabinose 0.6 mM for pGro7, pKJE7 and pT16, tetracycline 10 ng/ml for pG-T12 and both L-arabinose and tetracycline for pG-T12 were added into the 100 ml culture medium in the initiation of culture. When the culture OD reached to 0.7, 0.4 mM IPTG was added for induction of huscFv and incubation was continued for 4 hr with shaking (150 rpm) at 30°C. Then, the cells were harvested by centrifugation at 10000 g for 3 min at room temperature. The cells were resuspended in 10 ml of lysis buffer (100 mM NaCl, 50 mM NaH2PO4 at pH=8.0) and were disrupted by sonication (five 30 s pulses interrupted with cooling on ice).

Purification of huscFv

For purification of recombinant protein, the harvested cells were sonicated as described above and soluble part of cells was subjected to affinity chromatography. Briefly, supernatant was applied onto a column containing 2 ml Ni-NTA resin. The column then washed with buffer A (5 mM and 20 mM of imidazole) to remove non-specific proteins. Finally the sample was eluted by elution buffer containing 250 mM imidazole. The purified recombinant protein was analyzed by 12% SDS-PAGE followed by staining with Coomassie Blue G250. The concentration of purified huscFv was analyzed by nanodrop analyser.

Indirect ELISA for antigen-binding activity of anti-CD20 huscFv

To assay the affinity and determine the antigen binding activity of humanized single chain antibody, the Raji cell lysates (10⁶ cells per ml) were coated in ELISA plates (96-well plate) and incubated overnight at 4°C. After coating, the plates were washed three times with PBS and blocked by 300 μl blocking buffer.
(PBS buffer containing 3% BSA and 0.05% Tween 20) (PBS-T) for 1 hr at room temperature. In the next step, serial dilutions of soluble huscFv were added to the wells. After washing with PBS-T, 100 µ of HRP conjugated protein L (1:3000) were added into the wells, incubated for 1.5 hr at RT, and washed with PBS-T. The reaction was developed with 100 µ of TMB at room temperature. Finally, reactions were determined by reading the Optical Density at 450 nm (OD 450) to evaluate the affinity of antibodies.

**Results**

**Construction of expression plasmid pET22b-anti-CD20 huscFv**

For cloning and expression of anti-CD20 huscFv in *E. coli*, the pUC57 plasmid containing the gene coding of huscFv was digested with restriction digestion enzymes MlsI (MscI) and XhoI and the insert was subcloned into the MlsI (MscI) and XhoI site of pET22b (+) expression vector. This subcloning was further confirmed by subsequent restriction digestion analysis (Figure 1).

**Expression of recombinant protein**

For expression of anti-CD20 huscFv, the BL21 (DE 3) containing ET22b-huscFv was cultured in LB broth containing 100 mg/ml ampicillin. When the cell concentration reached approximately to the OD 600 of 0.7, the culture was induced by addition of 1 Mm IPTG for 4 and 24 hr at 25°C. Analysis by SDS-PAGE indicated that anti-CD20 huscFv was efficiently expressed in *E. coli* with a band of about 28 kDa. Analysis for solubility of recombinant huscFv showed that the majority of huscFv was expressed as IBs (Figure 2).

**Soluble expression of huscFv by co-expression of pET22b-huscFv constructs with chaperones plasmids**

In order to investigate the contribution of different molecular chaperones in soluble expression of anti-CD20 huscFv in *E. coli* BL21 (DE3), this anti-CD20 huscFv was co-expressed with different chaperones plasmids. The result indicates that chaperones enhanced soluble expression of huscFv. Co-expression of the huscFv-anti-CD20 with chaperone plasmid sets including pGro7, pG-KJE8, pTf16, pKjE7 and pG-Tf2 indicated enhancement in solubility in comparison to anti-CD20 huscFv that was expressed without the chaperones and the result was analyzed by NanoDrop which is shown in table 2. Importantly, the co-expression of pKJE7 containing GrpE/DnaK/DnaJ has the highest outcome (up to 50%) on soluble expression of recombinant huscFv compared to pGro7, pG-KJE8, pTf16 and pG-Tf2 chaperons (Figure 3).

**Purification of huscFv**

After cell harvest and lysis by sonication of the cells, soluble fractions were collected by centrifugation at 10000 g for 10 min and recombinant protein was purified by affinity chromatography method from supernatant of *E. coli* cell lysate. The result showed that purity of the recombinant protein was in high level and...
Figure 3. SDS-PAGE analysis of co-expressed anti-CD20 huscFv with different cytoplasmic molecular chaperones after induction with 0.4 mM IPTG for 4 hr at 30°C, washed and resuspended in lysis buffer 10 ml of lysis buffer (100 mM NaCl, 50 mM NaH2PO4 at pH=8.0). Lane 1: the standard protein weight marker, Lane 2: whole cell lysate supernatant from uninduced cells without chaperone plasmid set, Lane 3: whole insoluble pellet from uninduced cells without chaperone plasmid set. Lanes 4-6: samples of different extracts from E. coli BL21 (DE3)/pET22b-huscFv/pG-KJE7 (Lane 4: supernatant from uninduced cells, Lane 5: supernatant from 4 hr, Lane 6: pellet from 4 hr). Lanes 7-9: samples of different extracts from E. coli BL21 (DE3)/pET22b-huscFv/pGro7 (Lane 7: supernatant from uninduced cells, Lane 8: supernatant from 4 hr, Lane 9: pellet from 4 hr). Lanes 10-12: samples of different extracts from E. coli BL21 (DE3)/pET22b-huscFv/pKJE7 (Lane 10: supernatant from uninduced cells, Lane 11: supernatant from 4 hr, Lane 12: pellet from 4 hr). Lanes 13-15: samples of different extracts from E. coli BL21 (DE3)/pET22b-huscFv/pG-T2 (Lane 13: supernatant from uninduced cells, Lane 14: supernatant from 4 hr, Lane 15: pellet from 4 hr). Lanes 16-18: samples of different extracts from E. coli BL21 (DE3)/pET22b-huscFv/pTf16 (Lane 16: supernatant from uninduced cells, Lane 17: supernatant from 4 hr, Lane 18: pellet from 4 hr).

determined by SDS-PAGE and seemed as a single band (Figure 4). Afterwards, the concentration of refolded huscFv protein was determined by NanoDrop which is shown in table 2.

ELISA result

ELISA experiments were accomplished with anti-CD20-huscFv. The results demonstrated that huscFv is refolded by pKJE7, pG-KJE8 and pG-T2 chaperones had higher binding activity compared to huscFv without chaperone co-expression. On the other hand, the relative ELISA results of huscFv refolded by pGro7 and pTf16 chaperones are similar to huscFv without chaperone co-expression. All of the chaperones, huscFv refolded by pKJE7, revealed the highest binding activity.

Discussion

CD20 antigen expressed on cell surface of immature B cells is a validated target for treatment by anti CD20 mAb, Rituximab. This mAb is a chimeric monoclonal antibody and is approved for treatment of B-cell Non-Hodgkin’s Lymphoma (NHL) disease 16.

Due to the size and cost related drawbacks of full length monoclonal antibodies, development of smaller versions of these antibodies using low cost expression system of E. coli is the subject of great interest 17. In spite of low cost and high yield of protein production in E. coli, this expression system suffers from several disadvantages. Expression of eukaryotic proteins in E. coli generally results in production of the recombinant proteins as inclusion body that lack correct 3D structure 18. IBs are composed of proteins with inappropriate structures that are the result of protein expression in non-physiological conditions in the cytoplasm of E. coli. Periplasmic expression is one of strategies to overcome this impediment. Hence, pelB leader peptides were used to allow translocation of unfolded recombinant protein to the oxidizing space of periplasm. The disulphide bonds have an important role in correct folding of recombinant proteins that lead to high stability. So, productions of recombinant proteins in soluble form are the subject of extensive researches. For increasing soluble expression of anti-CD20-huscFv, various approaches are used including co-expression with molecular chaperone, use of chemical chaperones and optimization of the culture conditions. The present study aimed to evaluate the effects of co-expression with different chaperons on soluble expression of recombinant anti-CD20 huscFv 9,19,20. For this purpose, the sequence encoding for Rituximab based hscFv was transformed into the BL21 (DE3) for expression.

In our study, partial soluble expression of the huscFv was observed without using any molecular chaperons. Hu X et al indicated that the use of low temperature and low rate of inducer concentration led to enhancement in correct folding and finally increased concentration of soluble and functional form of recombinant proteins 21.

On the other hand, for enhancement of the huscFv soluble expression, various plasmids containing chaperone combinations have been used that revealed remarkable enhancement (up to 50%) of soluble expression in the cytoplasm. The related studies are in agreement with our results 22,23. Among different chaperone plasmids checked by co-expression, the pKJE7 plasmid

Figure 4. SDS-PAGE analysis of the purified recombinant huscFv in E. coli

Lane 1: the standard protein weight marker, Lane 2: purified huscFv/pTf16, Lane 3: purified huscFv/pGro7, Lane 4: purified huscFv/pG-KJE8, Lane 5: purified huscFv/pKJE7, Lane 6: purified huscFv/pG-T2.
set containing DnaK, DnaJ, GrpE chaperone genes demonstrated higher soluble yield in comparison to pGro-7, pG-KJE8, pTf16 and pG-Tf2 plasmids. High level of expression of huscFv with pKJE7 plasmid set depended on close activation between DnaK, DnaJ, GrpE chaperone proteins. The results were in agreement with studies done by Bo et al. They reported that anti-BNP scFv was expressed in soluble form by using pKJE7 plasmid set that has the highest amount of scFv, while increasing expressed soluble form of anti-BNP scFv led to reduction of anti-BNP scFv expression 23. In another study, Heo et al reported the enhancement of soluble expression of anti-c-Met scFv with co-expression of cytoplasmic chaperones 24. On the contrary, Nishihara et al showed that chaperone proteins containing GroEL and GroES have synergic effect with GrpE, DnaK, DnaJ in helping folding an Allergen of Japanese Cedar Pollen, Cryj2, in E. coli 19.

However, in our study, there were not any relative synergistic effects between GroEL and GroES proteins with GrpE, DnaK, DnaJ chaperone proteins. Also, in this study, binding activity of soluble huscFv produced by co-expression of combination of various cytoplasmic chaperones was analyzed by relative ELISA. The result showed that all of the soluble fractions had positive effects in binding activity, among them co-expression with plasmid pKJE7 led to 50% increase in huscFv binding activity compared with and without chaperone coexpression. Increasing the binding activity of huscFv co-expressed with plasmid pKJE7 depended on the absence of both GroESL and DnaKJE. These results indicated that there is not any constructive collaboration between GroELS and DnaKJE. Similar results were previously reported for recombinant protein production in the cytoplasm. Sonoda H et al proved that co-expression of DnaKJE with GroELS has negated the effects of GroELS 25. Furthermore, Sonoda H et al proved that in the cytoplasmic production system, co-expression of GroES/GroEL displays 4.6 fold evaluation in antigen- binding activity 26.

Conclusion
The result of this study indicates that co-expression of pKJE7 cytoplasmic chaperone with huscFv leads to 50% increase in soluble expression. Furthermore, the result of binding activity revealed that all chaperon plasmids had positive effect in huscFv binding activity. Among them, the maximum huscFv binding activity was obtained when huscFv was co-expressed with pKJE7 plasmid chaperone set.

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Conflict of Interest
The authors declare that they have no competing interests.

Ethical Approval
This article does not contain any studies with human participants or animals performed by any of the authors.

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