Functional expression of CCL8 and its interaction with chemokine receptor CCR3

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Abstract

Background: Chemokines and their cognate receptors play important role in the control of leukocyte chemotaxis, HIV entry and other inflammatory diseases. Developing an efficient method to investigate the functional expression of chemokines and its interactions with specific receptors will be helpful to assess the structural and functional characteristics as well as the design of new approach to therapeutic intervention.

Results: By making systematic optimization study of expression conditions, soluble and functional production of chemokine C-C motif ligand 8 (CCL8) in Escherichia coli (E. coli) has been achieved with approx. 1.5 mg protein/l culture. Quartz crystal microbalance (QCM) analysis exhibited that the purified CCL8 could bind with C-C chemokine receptor type 3 (CCR3) with dissociation equilibrium constant ($K_D$) as $1.2 \times 10^{-7}$ M in vitro. Obvious internalization of CCR3 in vivo could be detected in 1 h when exposed to 100 nM of CCL8. Compared with chemokine C-C motif ligand 11 (CCL11) and chemokine C-C motif ligand 24 (CCL24), a weaker chemotactic effect of CCR3 expressing cells was observed when induced by CCL8 with same concentration.

Conclusion: This study delivers a simple and applicable way to produce functional chemokines in E. coli. The results clearly confirms that CCL8 can interact with chemokine receptor CCR3, therefore, it is promising area to develop drugs for the treatment of related diseases.

Keywords: Agonist, CCL8, Chemokine, Chemokine receptor CCR3, Expression, Interaction

Background

Chemokines are a class of structurally related chemotactic cytokines, which along with their cognate receptors can regulate a variety of cellular functions, including immunodeficiency virus type I infection, cancer metastasis, arthritis, asthma and neurodegenerative diseases [1, 2]. Chemokines and their derived antagonists have been demonstrated as an effective way for treatment of allergic diseases [3, 4]. Therefore, chemokines are attracting more and more interests in therapeutic potential and extensively demonstrated in the pharmaceutical development [5, 6].

Despite the simplicity of these small cytokines, structural and functional studies of chemokines are far more complicated than initially expected [2]. Firstly, it is challenging to obtain functional chemokines in considerable amount (milligram) from natural sources [7]. Over-expression of chemokines in E. coli can be achieved, but tends to form inclusion bodies [8], where refolding of chemokines are time consuming and costly effective [9]. Secondly, due to the promiscuity of the chemokine system, the chemokine receptor can bind with various chemokines, such as C-C chemokine receptor type 1 (CCR1) can bind with chemokine C-C motif ligand 3 (CCL3), chemokine C-C motif ligand 5 (CCL5), chemokine C-C motif ligand 7 (CCL7) and chemokine C-C motif ligand 23 (CCL23). Conversely, one chemokine can also interact with different receptors, for example CCL5 (Rantes) can interact with CCR1, CCR3 and C-C chemokine receptor type (CCR5) [10, 11], inducing different responses [12]. The crosstalk of chemokines and chemokine receptors are still not clear so far [1, 13]. Therefore, developing an efficient method for functional expression of chemokines and their clear interactions with receptors will be helpful for designing new medicines with high efficacy and low side effects [3].

CCL8 (monocyte chemotactic protein-2, MCP-2) belongs to the CC chemokine sub-family [1], which has been reported as an agonist of C-C chemokine receptor type 2 (CCR2) and CCR5 [14], and plays a pivotal role in...
the control of leukocyte chemotaxis, HIV entry and other inflammatory diseases [15–17]. Despite its important medical purpose, CCL8 as well as other chemokines tends to form inclusion bodies when overexpressed in *E. coli* [8, 18]. There is increasing demand for developing protocols to obtain milligrams quantity of soluble and functional CCL8 for biological studies and drug screening [19–21]. The cross interaction of CCL8 with another important allergic related chemokine receptor CCR3 [9, 22] are still highly controversial. Several reports in the literature indicate that CCL8 is one of the potential agonist of CCR3 [1, 6, 23], while others are not involved [2, 12, 14, 24]. To our knowledge, still no experimental reports exist on interactions of CCL8 with CCR3, and their detailed binding kinetics, thermodynamics and functional responses are still not well characterized. Here, a simple and efficient protocol for soluble and functional production of CCL8 in *E. coli* was established.

Binding assay of CCL8 with CCR3 in vitro was carried out using QCM method. Internalization and chemotaxis of CCR3 expressing cells induced by CCL8 were also characterized. Our work provides an efficient way to produce functional chemokines in *E. coli*, and provides the detailed interaction informations of CCL8 with CCR3 for potential drug development and treatment of related diseases.

### Results

#### Optimization of expression conditions

To achieve soluble production of CCL8, the culture conditions of *E. coli* harboring pET28 a-His$_{12}$-CCL8 plasmid were systematically optimized, such as culture temperature, induction phase and induction concentrations. The induction phase plays an important role for soluble production of CCL8. As shown in Fig. 1a, the best induction phase is found as induction at OD$_{600nm}$

![Fig. 1](image)

**Fig. 1** Optimization of induction phase, inducer concentration and induction temperature. Production of CCL8 was characterized using dot-blot. The intensities of dot-blot were averaged and error bars were calculated based on three times experiments. **a** Effect of induction phase on the expression of pET28a-CCL8. **b** Effect of inducer concentration on the expression of pET28a-CCL8. **c** Effect of expression temperature on the expression of pET28a-CCL8.
of 0.4-0.6 when induced with 0.5 mM Isopropyl β-D-Thiogalactoside (IPTG). Later or earlier induction would significantly result in lower yield of soluble aimed proteins. As biologically active cytokine, the accumulation of CCL8 would be harmful for *E. coli* cells. Therefore, total induction time can also obviously influence yield of soluble CCL8 (Fig 1a). It is observed that within 10 h, the cell density increased with increasing culture time. However, after that the cell density decreased as time prolonged, resulting in lower yield of soluble proteins. Furthermore, culture temperature and concentration of inducer also influence the growth of *E. coli* cells and show obvious effects on soluble production of CCL8 (Fig. 1b and c). After optimization, maximum production of soluble CCL8 was obtained when induced by 0.5 mM IPTG at OD600nm as 0.4-0.6, with induction temperature as 18°C for totally 10 h induction. The total expression level of His12-tagged CCL8 fusion protein was determined as 12-16 mg protein/l culture using dot blot method where the purified chemokine CCL11 was used as a standard.

**Purification and characterization of CCL8**

Six liters of *E. coli* cells containing pET28a-His12-CCL8 plasmid were cultured at optimized conditions and then harvested by centrifugation. After cell lysis, the soluble His12-tagged CCL8 fusion protein was purified via Ni2+ affinity chromatography. The eluted protein shows a distinct band at 14 kDa, which is consistent with the theoretical molecular weight of His12-tagged CCL8 fusion protein. The result was further confirmed by western blot analysis using anti-6×His tag monoclonal antibodies. The yield of His12-tagged CCL8 fusion protein was determined as 5-8 mg/l culture with purity over 90%, and the averaged recovery rate is about 50% in fusion protein purification. The fusion protein was then digested by His6-TEV (Tobacco Etch Virus) enzyme to remove the His12-tag fusion part. After that, the digested mixture was passed through another Ni2+ affinity chromatography column, on which the His12-tagged fusion part and His6-TEV were captured, and CCL8 without a tag was recovered as flow-through. The apparent molecular weight of purified CCL8 on the SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) was 8.9 kDa (Fig. 2a), which is agree with its theoretical molecular weight. The result was further confirmed with mass spectrometry analysis (Fig. 2b). The final yield of CCL8 after digestion and purification was about 1.5 mg protein/l culture, corresponding to a recovery yield of 19-30% from fusion proteins. The secondary structure of CCL8 was analyzed using circular dichroism (Fig. 2c), which exhibit a mixture of α helix and β-sheets, similar as other typical chemokines [1, 22]. The secondary structure content of the purified CCL8 was estimated on Dicroweb as 22% α helix and 18% β-sheets, which is well consistent with the 25% α helix and 20% β-sheets of crystal structure analysis [14]. It suggested that the purified CCL8 has been well folded into a reasonable structure.

**Binding assay of recombinant CCL8 with CCR3**

In order to determine whether CCL8 bind with CCR3 in vitro, the binding assay of CCL8 with chemokine receptor CCR3 was carried out using QCM method. From Fig. 3, typical QCM binding and dissociation curves are clearly observed compared with the control, suggesting that CCL8 can bind with CCR3 in vitro. By fitting these curves, the $K_D$ value between CCR3 and CCL8 was obtained as 1.2 x 10^{-7} M, which is comparable to native agonists of CCR3, i.e. CCL11 and CCL24. They bind with CCR3 with $K_D$ as 3.7 x 10^{-7} M and 3.0 x 10^{-7} M, respectively [22]. However, the association rate $k_a$ and dissociation rate $k_d$ values for CCL8 binding with CCR3 are 8.6 x 10^{4} M^{-1} s^{-1} and 0.00106 s^{-1} respectively (Table 1), which is much slower than those of CCL11 and CCL24 (Table 1). These results suggest that CCL8 is a potential agonist of CCR3 with similar binding affinity as CCL11 and CCL24, but slower kinetic binding rate [22]. It should be noted that the data obtained for CCR3 binding with CCL11 and CCL24 were determined using surface plasma resonance (SPR) methods, while the data for CCR3 binding with CCL8 were characterized using QCM, the difference of $K_D$ value was generated from the two different methods.

**Internalization assay**

Internalization of GPCR’s has been considered as a possible way of receptor desensitization after agonist stimulation [25]. The induced internalization of CCR3 with CCL8 on the cell surface was assessed using the TRx-HEK293 cells (Thermo Fisher) stably transfected with CCR3. The cells were treated with 100 nM CCL8, and the internalization process of CCR3 was imaged on a Nikon A1 confocal microscopy. After 30 min stimulation, CCR3-EGFP protein (enhanced green fluorescent) tended to form large aggregates and transferred into cytosol, which means that obvious internalization of CCR3 could be induced when treated with CCL8 (Fig 4).

**Chemotaxis test**

Chemotaxis assays are useful tools for the evaluation of chemotactic ability of agonists. To explore the chemotactic ability of CCL8 inducing CCR3 expressing cells, chemotaxis test was carried out on Transwell chambers using TRx-HEK293 cells stably transfected CCR3-EGFP. From Fig. 5, it can be seen that CCL8 induce
obvious chemotactic migration of TRx-HEK293 cells stably transfected CCR3, enabled that CCL8 can interact with CCR3 in vivo, confirms that CCL8 is one of agonists of CCR3. Compared with other agonists of CCR3, the chemotactic index of CCL8 is similar to CCL5, but lower than those of CCL11 and CCL24 (Fig. 5). The results obtained from the binding assay in vitro shows that CCL8 has similar binding affinity but slower binding and dissociation rate with CCR3 than those of CCL11 and CCL24, which may be an agreement for its weaker chemotactic index than CCL11 and CCL24.

**Discussion**

Chemokines are clinically significant proteins, their structural and functional studies have attracted more and more interests nowadays [26, 27]. To obtain a high concentration of biological active chemokines in milligram with high purity is challenging tasks because of its naturally low yield and the formation of inclusion bodies when overexpressed in *E. coli* [8]. There are normally two ways to produce large amount of chemokines: one is overexpression of chemokines as inclusion bodies and then refolded them from denatured conditions [8, 28]. Another approach is expressing chemokines as fusion proteins.
protein with different fusion tags, such as maltose binding protein (MBP), glutathione-S-transferase (GST) [19], and NusA [29]. Although this fusion chemokines have been proved to retain their binding activity with their cognate receptors, but to exclude any possible nonspecific interactions, the fusion part always need to be removed by exogenous enzyme digestion and further chromatography methods. The two methods mentioned above are usually low yield, time consuming and costly. By systematical efforts on optimization of expression conditions, the formation of inclusion bodies can also be significantly decreased, and soluble and functional expression of cytokines can be achieved [22, 30]. Our previous study has demonstrated soluble production of human chemokines CCL11 and CCL24, and here we present the protocol for soluble production of biologically active CCL8. We found that, although chemokines have similar tertiary structure and functions, but different optimal expression conditions are needed to achieve high level soluble production. For example, CCL5 (RANTES) cannot achieve milligram production of soluble and functional protein even with similar methods, suggested that it is necessary to develop a different protocol for production of various chemokines.

Table 1 The thermodynamic and kinetic constants for CCR3 binding with different ligands

| Agonists | $k_a$ ($M^{-1} s^{-1}$) | $k_d (s^{-1})$ | $K_D (M)$ |
|----------|------------------------|---------------|---------|
| CCL8     | $8.6 \times 10^4$      | $0.00106$     | $1.2 \times 10^{-7}$ |
| CCL11$^a$| $2.0 \times 10^4$      | $0.0740$      | $3.7 \times 10^{-7}$ |
| CCL24$^a$| $6.3 \times 10^3$      | $0.1900$      | $3.0 \times 10^{-7}$ |

$^a$Data were taken from (Wang et al. [22])

Chemokines participate in leukocyte trafficking and function, implicated in health and disease. Heterophilic interactions between chemokines and chemokine receptors may modulate their activities. However, systematic evaluation of interactions between chemokines and their specific receptors has not been established [31]. Human CCR3 (hCCR3) and CCL8 both belong to the CC chemokine-chemokine receptor sub-family [1], and plays pivotal roles in the control of leukocyte chemotaxis, HIV entry and other inflammatory diseases [15–17]. Unfortunately, the interaction of CCR3 and CCL8 is still not well characterized, therefore highly debated. Here using QCM binding assay as well as internalization and chemotaxis tests, we confirmed that CCL8 is one of the agonist of chemokine receptor CCR3, and their detailed kinetic and thermodynamic binding constants in vitro and in vivo were also characterized. Compared with other ligands of CCR3, CCL8 shows similar interaction with CCR3 as the universal ligands CCL5, but different from the native ligands of CCR3, such as CCL11 and CCL24. Our results provide important information for CCL8-CCR3 interaction system. These facts wukk deliver a great advantage in potential drug design and treatment of related diseases.

Conclusions

A simple and reliable method for soluble expression of chemokine CCL8 has been established with a yield of 1.5 mg protein /l culture. The purified CCL8 can bind with chemokine receptor CCR3 in vitro with $K_D$ value as $1.2 \times 10^{-7}$ M, which is similar as CCL11 and CCL24. Obvious internalization and chemotactic migration of CCR3 expressing cells can be observed when induced with CCL8, which confirmed that CCL8 is one of the agonist of chemokine receptor CCR3. The antagonist development based on CCL8 will be a useful tool for development of efficient drugs and treatment of related diseases.

Methods

Recombinant plasmid construction

Gene of human CCL8 (NCBI locus: 1ESR_A) was commercially synthesized by GenScript Bio-company (China) with optimized codon usage for over- expression in E. coli. The gene was amplified by PCR using a forward primer as 5’-AGTGGATCCATCATCACCACCATGATGAGCATC -3′ and reverse primer as 5’-ACTAAGCTTTTAATACGCTTGTTTCTGAGAAAAA TCTGATC C AG-3’. The PCR product were digested with BamHI and Hind III, and then ligated into pET28a vector (Novagen, USA) at the downstream of a T7 promoter. To facilitate downstream purification, another 6 x His tag was added onto the N terminus of CCL8 gene, resulting in pET28a-His<sub>12</sub>-CCL8 plasmid. A
TEV enzyme digestion site was also inserted between the 6 × His tag and CCL8 gene for removal of any fusion part after purification. After confirmed by DNA sequencing, the recombinant plasmid was then transformed into *E. coli* strain BL21 (DE3) (Novagen) for optimization and protein expression.

**Optimization of expression conditions**

In order to achieve maximal soluble production of CCL8, the expression conditions of cells harboring pET28a-His₁₂-CCL8 were optimized systematically using a similar method as previously reported [22]. Briefly, colonies from the transformation plates were cultured in 5 ml Luria-Bertani broth (LB) medium overnight at 37°C with shaking. 1 ml of overnight culture was then inoculated into 100 ml of fresh Terrific Broth (TB) medium supplemented with 50 μg/ml kanamycin, and IPTG were applied to induce overexpression of target proteins. The induction phase, concentration of inducer and induction temperature were optimized by inducing at OD₆₀₀nm as 0.4-0.6, 0.6-0.8, 0.8-1.0, 1.0-1.2 and 1.4-1.6, and IPTG concentrations of 0.1 mM, 0.2 mM, 0.3 mM, 0.5 mM and 1 mM and induction temperature as 18°C, 25°C and 37°C, respectively. After induction with IPTG, samples were taken at regular times, and then lysed and centrifuged at 12000 g for 10 min at 4 °C. Supernatant were then taken and analyzed using dot-blot with mouse anti-His monoclonal antibody as primary antibody and HRP (horse radish peroxidase)-labeled goat anti-mouse antibody as secondary antibody. The blot was finally stained using the Amersham ECL plus, and then detected on a FLA-5100 imaging system (Fuji, Japan). The intensity of each dot was analyzed with MultiGauge Ver.3.X software (Fuji, Japan) according to manufacturer’s instructions (Additional file 1: Figure S1).

**Expression and purification of CCL8 in *E. coli***

Six liters of *E. coli* cells were harvested and resuspended in phosphate buffered saline (PBS) buffer. After broken on an ultra-high pressure homogenizer, the crude extract was obtained by centrifugation at 12,000 g for 20 min at 4 °C, and then applied to a HiTrap™ Chelating HP column (GE Healthcare). The impurities were washed off with PBS buffer containing 250 mM imidazole (pH 7.4) and the target CCL8 fusion protein was eluted with PBS buffer containing 500 mM imidazole (pH 7.4). The eluent was then buffer exchanged into 1 × PBS, and 6 ×
His-tagged TEV enzymes (plasmids expressing TEV enzymes was obtained as courtesy from Alan Fersht’s lab at LMB, UK) were added for digestion of His$_{12}$-CCL8 fusion proteins. The digestion mixture was passed through another HiTrap™ Chelating HP column, and the cleaved fusion part with His$_{12}$-tag and the His$_{6}$-TEV enzyme will be captured on the column, and CCL8 is recovered in the flow-through fraction. The purified CCL8 was collected and characterized using SDS-PAGE, mass spectrum and circular dichroism spectroscopy respectively.

Quartz crystal microbalance measurement

The binding assay of CCL8 with CCR3 in vitro was carried out on a QCM-Z500 (Biolin Scientific, Sweden) instrument using Q-sense His tag capturing sensor at 25 °C. The sensor was first activated with 0.5 mM NiCl$_2$. The 6 × His tagged CCR3 purified from stably transfected HEK293 cells was prepared as our previously reported [9] and immobilized on the sensor. Different concentrations of CCL8 were flowed through the sensor, respectively, and the binding of CCL8 to the immobilized CCR3 was monitored in real time with a flow rate of 50 μl/min. 5 μM BSA (bovine serum albumin) was used as control to exclude any possible non-specific interaction. The sensorgrams were fitted globally with a 1:1 binding model [22].

Internalization assay

The TRx-HEK293 cells stably transfected CCR3-EGFP previously constructed in our lab [9] was used for the receptor internalization assay. The cultured cells were cultured in Dulbecco's Modified Eagle Medium (DMEM) medium containing 0.5% bovine serum, and before internalization assay, cells was washed with HEPES buffer, and then stimulated with 100 nM CCL8. Images of CCR3-EGFP on cell surface before and after stimulation were captured by a confocal microscopy (Nikon A1, Japan) at different incubation times.

Chemotaxis test

TRx-HEK293 cells stably transfected with CCR3-EGFP [9] were resuspended ($1 \times 10^4$) in DMEM containing 0.5% (v/v) bovine serum albumin. 100 μl cell suspensions were placed in the upper wells of Transwell chambers (Corning, USA) containing bare filter with a pore size of 8 μm. The CCL8 (100 nM) in 500 μl same medium was placed in the lower chambers. After incubation at 37 °C for 4 h, cells migrated through the filter were stained with crystal violet and counted by image J software. Migration index is obtained as values of migrated cell numbers induced with chemokines divided by migrated cell numbers without chemokines.

Additional file

Additional file 1: Figure S1. Optimization of induction phase. Production of CCL8 was characterized using dot-blot. The intensities of dot-blot were averaged and error bars were calculated based on three times experiments. (A) Dot-blot images for optimization of induction phase. (B) Histogram for effect of induction phase on the expression of pET28a-CCL8. (TIFF 1708 kb)

Abbreviations

BSA: bovine serum albumin; CCL11: Chemokine C-C motif ligand 11; CCL23: Chemokine C-C motif ligand 23; CCL24: Chemokine C-C motif ligand 24; CCL3: Chemokine C-C motif ligand 3; CCL5: Chemokine C-C motif ligand 5; CCL7: Chemokine C-C motif ligand 7; CCL8: Chemokine C-C motif ligand 8; CR1: C-C Chemokine receptor type 1; CCR2: C-C Chemokine receptor type 2; CCR3: C-C Chemokine receptor type 3; CCR5: C-C Chemokine receptor type 5; DMEM: Dulbecco's Modified Eagle Medium; E. coli: Escherichia coli; EGFP: enhanced green fluorescent protein; GST: glutathione-S-transferase; HRP: horse radish peroxidase; IPTG: isopropyl β-D-Thiogalactoside; Kd: dissociation equilibrium constant; LSB: Luria-Bertani broth; MBP: maltose binding protein; PBS: phosphate buffered saline; QCM: Quartz crystal microbalance; SDS-PAGE: sodium dodecyl sulfate polyacrylamide gel electrophoresis; SPR: surface plasma resonance; TB: Terrific Broth; TEV: Tobacco Etch Virus

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

BG and TS designed the research plan. TS, JL, YS and ZW conducted the experiments. BG, JL, and TS analyzed the data and interpreted the results. BG and TS drafted the manuscript, and all authors revised the manuscript. The authors declare that they have no competing interests.

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