Development of novel interferon alpha2b muteins and study the pharmacokinetic and biodistribution profiles in animal model

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ABSTRACT

Novel human interferon alpha2b (hIFNα2b) muteins were developed by substituting cysteine residue (C) at positions 2 and 99 with aspartic acid residues (D). The mutein forms were then studied for pharmacokinetic profile. In addition, the influence of charge on the protein structure was tested in vivo for the biodistribution pattern. Codon substitutions were performed by Polymerase Chain Reaction (PCR)-based site-directed mutagenesis on a previously constructed synthetic hIFNα2b open reading frame (ORF) cloned in pET32b expression plasmid. The result of nucleotide sequencing analysis confirmed that all codons were replaced successfully without any additional mutation. Three mutant forms of hIFNα2b ORF were overexpressed in Escherichia coli BL21 (DE3) resulted in three muteins: hIFNα2b C2D, hIFNα2b C99D, hIFNα2b C2D C99D. To follow the kinetic and localization of the mutein interferon after intravenous administration, Tc99m was used to label the proteins. In particular of elimination half-life, it was shown that hIFNα2b C2D C99D > hIFNα2bC2D > hIFNα2bC99D > wild type. hIFNα2b C2D C99D mutein showed highest blood accumulation after 30 minutes administration. Taken together, the charge of hIFNα2b seems to be responsible for the fate of hIFNα2b in vivo.

Keywords: Mutein; Human Interferon Alpha2b; Amino Acid Substitution; PCR Based Site Directed Mutagenesis; Tc99m labeling; Pharmacokinetic; Biodistribution; Protein Charge

1. INTRODUCTION

Interferon (IFN) is a cytokine produced and secreted by almost all eucaryotic cells as a response to viral, bacterial, antigen, or mitogen stimuli. Based on their receptor types on the cell membrane surface, IFN is classified into type I which consists of IFNα, IFNβ, IFNτ, IFNω and type II which consists of IFNγ. hIFNα2b, as a subclass of IFNα, is a glycoprotein consisting of 165 amino acids with size of 19271 Dalton. The molecule’s O-glycosylation at threonine position 106 is not important for its biological activity. hIFNα2b has two disulfide bridges formed by cysteins (between positions 1 and 98, and between 29 and 138). hIFNα2b is used as standard therapeutic protein for hepatitis B and hepatitis C treatments. In addition, the potential use to treat several types of cancer e.g. multiple myeloma, chronic myeloid leukemia, non-Hodgkin’s lymphoma, renal cell carcinoma, epidermoid cervical cancer, head and neck tumours, melanoma and medullary thyroid carcinoma was also demonstrated [1-3]. However, therapeutical outcome of hIFNα2b is not satisfied due to rapid elimination through renal clearance. Several strategies to improve hIFNα2b half-life have been reported, including pegylation, albumin fusion and glycosylation [3-5].

Previously, a synthetic ORF based on optimized codon for high expression in E. coli encoding for hIFNα2b was successfully constructed using thermodynamically balanced inside out method. The ORF was cloned and overexpressed in E. coli BL21 (DE3). The recombinant hIFNα2b was produced as a fusion protein of 37 kDa containing f-methionine as first amino acid, thioredoxin and polihistidine tags at its N terminus. The protein was confirmed as hIFNα2b using Nano LC MS/MS resulting 80% amino acids coverage [6]. In this report, novel muteins of this protein were developed to improve the negative charge on the protein. The main aim was to prevent renal clearance thus improving the plasma half-life.

There are 12 amino acids (Leu30, Lys31, Arg33, His34, Phe36, Arg120, Lys121, Gln124, Tyr122, Tyr129, Lys131,
and Glu132) in hIFNα2b are involved in the biological activity [7]. Among 4 cysteine residues, cysteine at position 1 and 98 which form a disulfide bridge was not required in biological activity of interferon alpha2b [8]. Even, the disruption of this disulfide bridge by serine substitutions resulted in higher antiviral activity [9]. In this report, we substituted those cysteines with aspartic acid to improve the net negative charge of the protein. Aspartic acid is the amino acid having lowest pI i.e. 2.77. By substituting of other amino acids with aspartic acid, the net charge of hIFN-α2b will be more negative. The expectation was to introduce a repulsion force on interferon when approaching basal renal membrane which is covered by negative protein matrices. To do so, PCR-based site-directed mutagenesis (SDM) was applied. SDM-based on PCR is widely used to test the role of particular residues in the structure, activity and ligand-binding capacity of a protein by introducing mutation into target DNA [10]. Comparing to mutant selective by mutagenic agents, SDM method is much more effective yielding desired mutation in the successful range of 50% - 100% [11]. Three mutant forms of hIFNα2b we developed, namely C2D, C99D, and C2D C99D, will be compared for the enhancement in the pharmacokinetic profile and the accumulation pattern in the body after intravenous injection.

2. MATERIAL AND METHOD

2.1. Bacterial Strains and Plasmids

pET32b-hIFNα2b recombinant plasmid from previous work was used as a template for SDM [6]. Luria Bertani (LB) broth containing 100 µg/mL of ampicillin was used for cultivation and LB containing 0.5 mM of isopropyl thio-

galactopyranoside (IPTG) was used as an inducer in gene expression step.

2.2. Animals

Male Wistar rats with age of 6 - 8 weeks and 250 - 300 g were used for pharmacokinetic study. While, Swiss mice weighing of 30 - 40 g were used for biodistribution assay. Animals used in the experiments received care in compliance with the “Principles of Laboratory Animal Care” and “Guide for the Care and Use of Laboratory Animals”.

2.3. Site-Directed Mutagenesis

Mutagenic primers were designed using DNA Star (DNA Star Inc, USA) as shown in Table 1. First PCR-based SDM was applied to substitute TGT2 into GAT. Recombinant pET32b carrying hIFNα2b ORF was used as a template. PCR was done in 50 µl reaction volume containing 400 µM dNTP, 1 × Pfu buffer, 2.5 U of Pfu DNA poly-

merase, 150 ng of SDMPFORC2D primer, 150 ng of SDMREVC2D primer, and 50 ng template. PCR was performed as follow: initial denaturation at 95°C for 30 s, 12 cycles of denaturation at 95°C for 30 s, annealing at 55°C for 1 m, and polymerization at 68°C for 6.5 m. PCR product was treated by 10 U DpnI at 37°C for 1 h and transformed into E. coli Top 10. Transformants were characterized based on plasmid size, and subsequently digested using EcoRI or HindIII. Nucleotide sequencing of the insert was then performed to evaluate the mutation. Recombinant pET32b that contained mutated hIFNα2b C2D ORF and pET32b carrying wildtype hIFNα2b ORF were used as templates in second SDM step, which was substituting TGC99 into GAC. The PCR was performed as described above. The mutagenic primers used were SDMPFORC99D and SDMREVC99D. The PCR was performed at 54°C for 1 min for annealing.

2.4. Protein Overproduction, Purification and Characterization

The protein overproduction was performed using optimized condition we obtained previously [12]. E. coli BL21(DE3) containing mutated plasmids overnight culture was added (3% v/v) to 1 L of LB broth containing 100 µg·mL−1 of ampicillin. The culture was incubated for about 1.5 h in a shaking incubator at 37°C 200 rpm and IPTG to 0.5 mM final concentration was added to midlog phase of cell culture. Incubation was continued for an additional 3 h. Cell pellet was harvested by centrifugation at 5000 g for 10 min. The cells were resuspended in lysis buffer (50 mM NaCl and 1 mM EDTA) and lysed by sonication at 2.5 Hertz in the presence of 1 mM of phenyl-

methyl sulphonfluoride. To prevent temperature elevation, the cells were sequentially sonicated and cooled on ice for 10 times, each time for 30 s. Soluble proteins were separated from IB by centrifugation at 10,000 g. The soluble protein was purified using Nickel column according to the manufacturer’s protocol (Protino, Germany). To recover IB the pellet were washed with washing buffer (lysis buffer containing 0.5% Triton X-100 pH 7.2) and resuspended in solubilizing buffer (guanidine HCl buffer consisted of 6 M GdnHCl in 50 mM Tris-HCl pH 8 containing 800 mM of 2-Mercaptoethanol (2-ME) for 30 min at

Table 1. Oligonucleotides of PCR-Based SDM used to mutate hIFNα2b.

<table>
<thead>
<tr>
<th>Primers</th>
<th>Nucleotide sequence (5’→3’)</th>
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<tbody>
<tr>
<td>SDMPFORC2D</td>
<td>CGA ATT CCC ATA TGG ATG ACC TGC CGC</td>
</tr>
<tr>
<td>SDMREVC2D</td>
<td>GCG GCA GGT CAT CCA TAT GGG AAT TCG</td>
</tr>
<tr>
<td>SDMPFORC99D</td>
<td>C CGT GAG GCC GAC GTT ATC CAA GGT G</td>
</tr>
<tr>
<td>SDMREVC99D</td>
<td>C ACC TTG GAT AAC GTC CGC CTC CAG G</td>
</tr>
</tbody>
</table>
room temperature. Subsequently, solubilized pellet was centrifuged at 15,000 g for 15 min and 4°C to separate hIFNα2b and unsolubilized material. hIFNα2b from IB was refolded by 9 days dialysis in refolding buffer (0.2 mM EDTA, 0.25 mM Dithiothreitol, 50 mM Tris and 0.4 M urea; pH 8.0). The All crude proteins and purified muteins (as soluble and IB) were analyzed using 15% Sodium Dodecyl Sulphate Polyacrylamide Gel Electrophoresis (SDS-PAGE) under denaturing condition. To evaluate the protein conformation, purified muteins were also characterized using 10% native PAGE. Imidazol was removed from purified protein using nanosep centrifugal concentration with 10 kDa cut off (Pall life science). Protein concentration was determined using Bradford method based on coomassie blue staining.

2.5. IFNα2b Radiolabeling

Interferons were labeled using 99mTc radioactive. Three different concentrations of SnCl2·2H2O (10, 30, and 50 µg/µL) were tested to reduce 99mTcO4-. SnCl2·2H2O was carefully weighed and diluted in 1 mL of HCl 0.1 N. The solution was added to 50 µg of IFNα2b and 2 mCi of 99mTcO4- . The mixture was homogenized and incubated at 37°C for 15 min. Purification was done using Sephacryl S-300 HR column. The purity of IFNα2b-99mTc was checked using paper chromatography with 10 cm of whatman 3 mm paper as stationary phase and dry acetone as a mobile phase. Percent of unreduced 99m-TcO4- was calculated as follow:

\[
\% \text{ unreduced } 99m\text{TcO}_4^- = \frac{\text{Activity of } 99m\text{TcO}_4^- \text{(cps)}}{\text{Total activity (cps)}} \times 100%
\]

Labeling reaction was carried out using optimal condition consisted of 50 µg of SnCl2·2H2O, 100 µg of IFNα2b and 10 mCi of 99mTcO4-. A similar reaction, but without protein was performed as a control. The mixtures were homogenized, incubated for 15 min, and loaded into Sephacryl S-300 HR column. 0.05 mM of phosphate buffer pH 7.4 was used as elution buffer. 1 mL of each fraction was collected and measured using dose calibrator (RI Deluxe IsotopCalib II, Victoreen, 139,000 N).

2.6. Pharmacokinetics Study

Four groups of animals (n = 3) were used: group received 99mTc-IFN-α2b wild type, 99mTc-IFN-α2b mutein C99D, 99mTc-IFN-α2b mutein C2D; and 99mTc-IFN-α2b mutein C2D C99D. Each group was divided into 2 classes: 10 and 30 min observations. Each radiolabeled interferon was intravenously injected with dose of 100 µCi. The biodistribution of radiolabeled interferons was observed in 2 different periods: 10 and 30 min. After each time period of administration of the radiolabeled protein, the mice were sacrificed. The blood and several organs (liver, kidney, heart, lung, intestine and lymph) were collected and weighed. The radioactivity was measured using single channel analyzer as described previously for pharmacokinetic study.

2.7. Biodistribution Assay

HepG2 cell line was maintained in DMEM containing penicillin, streptomycin, amphotericin B, gentamycin and 10% heat-inactivated fetal bovine serum (FBS) at 37°C in a humidified incubator with 5% CO2. After 70% confluence, the adhering cells were washed with PBS and detached using 0.25% trypsin/EDTA. Cells (6 × 105/well) were seeded to 96 wells plate and incubated overnight. The medium was replaced by 150 µL of fresh medium containing IFN wildtype or muteins (concentration ranged from 0.25 to 5 µg/mL). Cell in medium alone was used as a negative control. Proliferation of cells was determined at 72 h after IFN treatment. The cells were washed using PBS and the viable cells were measured based on MTT conversion into formazan by mitochondrial reduce-tase. 20 µL of MTT solution (5 mg/mL) was added to
each well and incubated for 3 h. The medium was discarded and insoluble formazan crystals were dissolved in 200 µL of dimethyl sulfoxide. The plate was shaken for 1 min and absorbance was measured at 515 nm. The assay was analyzed in three independent experiments.

2.9. Statistical Analyses

All data were analyzed with SPSS program. Statistical differences between the control and IFN-treated groups were calculated by unpaired student’s t-test and considered significant at P < 0.05.

3. RESULT

3.1. Site-Directed Mutagenesis

The sequences of the oligonucleotides used in PCR based SDM was presented in Table 1. By using LB agar containing ampicillin selection, there were 60 transformants identified at first SDM step. Five recombinant plasmids were isolated and characterized. Migration analysis showed that all recombinant plasmids had similar size to wild-type (Figure 1(a)). Single digestion analysis using EcoRI or HindIII gave single band with 6.4 kb in size as expected (Figure 1(b)). Nucleotide analysis was performed to determine whether the recombinant plasmids contained the mutated hIFNα2b ORF. Using Seqman DNA Star alignment software, the sequences data confirmed that four plasmids harbored desired nucleotide change without any additional mutation (Figure 2). These recombinant plasmids were used as templates for second SDM step.

At second SDM step, it produced 70 transformants from each template. Five of them were characterized by mi-

Figure 1. Characterization of recombinant plasmids (a) Migration analysis: lane 1 = pET32b-hIFNα2b wildtype, lanes 2 - 6 = recombinant plasmids isolated from transformants; (b) Digestion analysis: lane 1 = DNA marker, lane2 = pET32b-hIFNα2b wildtype uncut, lane 3 = pET32b-hIFNα2b wildtype cut by EcoRI, lanes 4 - 8 = recombinant plasmids isolated from transformants cut by EcoRI, lanes 9 - 13 = recombinant plasmids isolated from transformants cut by EcoRI.

Figure 2. Nucleotide sequence analysis of transformants recombinant plasmid resulted from first SDM step using T7 promoter and terminator primers.
igration and single digestion analyses. All recombinants plasmid showed similar size to the wildtype with 6.4 kb in size (data not shown). Nucleotide sequence analysis showed that four plasmids from each template have been successfully mutated without any undesired mutation (Figure 3). The two steps PCR-based SDM resulted

Figure 3. Nucleotide sequence analysis of transformants recombinant plasmid resulted from second SDM using T7 terminator primers: (a) pET32b-hIFNα2b as a template; (b) pET32b-hIFNα2b C2D as a template.
in three mutated recombinant plasmids: pET32b-hIFNα2b C2D, pET32b-hIFNα2b C99D, and pET32b-hIFNα2b C2D C99D.

3.2. Protein Overproduction, Purification and Characterization

All mutated recombinant plasmids were transformed into *E. coli* BL21 (DE3) to express the ORFs. The overproduction was done at 25°C for 3 h with 200 rpm vigorous shaking and 0.5 mM IPTG. Total proteins were characterized in 15% SDS PAGE. As shown in Figure 4(a), at induced condition, all muteins showed similar band size (about 37 kDa) to wildtype form. The muteins were purified using nickel affinity column to obtain soluble protein (Figure 4(b)) as well as isolated IB (Figure 4(c)). The muteins amount both as soluble and IB were measured by Bradford assay. The muteins yield was ranging from 8.7 - 8.9 mg/L culture as soluble with 14% - 15% recovery and 28.7 - 28.8 mg/L culture as solubilized IB with 50% - 51% recovery. The renaturation of solubilized IB was performed according to previous work [12]. Based on non reducing SDS PAGE result, it was shown that solubilized IB was refolded partially. Characterization of the muteins conformation using non denaturing PAGE showed that the wildtype form gave two different conformations (Figure 5 lane 1) as also shown by mutein C2D and C99D. In contrast, mutein C2D C99D showed one band indicating that only one conformation was resulted.

3.3. IFNα2b Radiolabeling

Pure ⁹⁹mTc-IFNα2b with free TcO₂ was shown on paper chromatograph (Figure 6). The most appropriate concentration of SnCl₂·2H₂O to reduce TcO₂ was achieved at 50 µg/mL (Table 2). To remove TcO₂ and TcO₄⁻ after radiolabeling, Sephacryl HR-300 column was used as shown in Figure 7 which represented wildtype form. Similar chromatograms were observed for all muteins (data not shown). Protein characterization of fraction number 10 - 15 using SDS-PAGE showed IFNα2b band corresponding to 37 kDa size.

3.4. Pharmacokinetics Study

Blood protein concentration as a function of time was shown in Figure 8. As depicted in that figure, all mutein forms showed similar profile as the wildtype form but had longer elimination half-life (p < 0.05), 178 min (C2D), 118 min (C99D), 184 min (C2DC99D), and 57 min (wildtype) respectively.

3.5. Biodistribution Assay

After 10 min of intravenous administration, only mutein C2DC99D showed highest accumulation in liver (Figure 9(a)) and showed highest blood accumulation after 30 min of injection (p < 0.05). All mutein forms showed higher renal distribution especially at 10 min (p < 0.05).

3.6. Antiproliferation Assay

As shown in Figure 10, although a dose-dependent activity, no significant different on the antiproliferation effect was observed between wildtype and muteins at all concentration tested.

4. DISCUSSION

Several strategies have been reported to obtain mutation using site-directed mutagenesis. In this study, we applied mutagenic overlapping primers to introduce mutation on our recombinant hIFNα2b. The mutagenic primers were
Figure 5. Muteins characterization by native PAGE. Lane 1 = wildtype, lane 2 = mutein C2D, lane 3 = mutein C99D and lane 4 = C2D C99D.

Figure 6. Paper chromatograph of labeled interferon. The TcO₄⁻ peak was only detected in a standard reaction.

Table 2. The yield of radiolabel interferon using different concentrations of reducing agent.

<table>
<thead>
<tr>
<th>SnCl₂·2H₂O (µg/mL)</th>
<th>TcO₄⁻ (%)</th>
<th>⁹⁹mTc-IFNα₂b + TcO₄⁻ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>3.73</td>
<td>96.20</td>
</tr>
<tr>
<td>30</td>
<td>3.54</td>
<td>96.46</td>
</tr>
<tr>
<td>50</td>
<td>0.36</td>
<td>99.64</td>
</tr>
</tbody>
</table>

Figure 7. Chromatogram of labeled ⁹⁹mTc-IFNα₂b after applying through size exclusion Sephacryl HR-300 column.

Figure 8. Blood protein concentration as a function of time after intravenous injection of radiolabel proteins.

Figure 9. Biodistribution profile of wildtype and mutein forms of ⁹⁹mTc-hIFNα₂b. (a) 10 minutes after administration; (b) 30 minutes after administration.

Figure 10. Antiproliferation analysis of interferon alpha2b in HepG2 cells.
designed using pET32b-hIFNα2b sequence as a template with following consideration: both primers contained desired mutation and annealed to the same sequence of opposite strand, have 25 - 26 bps in length, and have desired mutation in the middle of primer sequences. To increase sequence fidelity, Pfu DNA polymerase enzyme which has 3’ - 5’ exonuclease activity was used. Nucleotide sequence analysis confirmed that most recombinant plasmids (80%) carrying desired mutation with no undesired mutation, 13% showed undesired mutation and only 7% of recombinant plasmids showed no mutation. This means, the efficiency of our method was 80%. This result was in agreement with Hemsley [14] which also claimed 80% efficiency. The presence of undesired mutation (13%) can be due to the lack of Pfu fidelity. It was reported that Pfu error rate was $1.3 \times 10^{-6}$, but Cline [15] observed that the error rate depended on PCR reaction, such as concentration of dNTPs (100 - 300 µM) and MgSO$_4$ (2 - 3 mM) at certain pH (8.5 - 9.1). Out from those conditions, the error rate can be 40-fold higher. The higher dNTPs concentration (400 µM) we used in this study seems to be good condition during PCR reaction. As a result, unfortunately unexpected additional mutations were observed (13%).

When the mutated plasmids obtained were overproduced, the yield and recovery of muteins as soluble and IB were not significantly different as compared to wild-type form [12]. In addition, the refolded muteins showed similar result to the wild type indicating that the substitution did not have any effect on the protein solubility. Characterization using native PAGE showed two different conformations on the wild type form. Disulfide interchange suggested to be responsible for this different which resulted in unexpected conformation. Changing only one of two cysteines residues involved in this interchange conformation did not prevent the interchange conformation. However, when both cysteines were substituted, unexpected conformation was eliminated, indicating the disulfide interchange was disappeared. The denaturing SDS PAGE (data not shown) confirmed that all muteins had the same size (37 kDa) with the wild type indicating that no significant change of molecular weight.

The benefit of amino acid substitution on hIFNα2b we reported here was shown, in particular in the plasma half-life, as well as in the biodistribution profile alterations. Although no change in the pattern of plasma concentration vs time curve, mutein forms of hIFNα2b revealed longer circulation time as compared to wild type form (Figure 8). Both wild type and mutein forms of hIFNα2b showed biphasic curve which indicated that the proteins have a two-compartment model of pharmacokinetic. Improving the circulation time of mutein hIFNα2bs, although minor, seems to be a charge-dependent. Substituting the cysteine residue with aspartic acid resulted in improved net negative charge of the hIFNα2b as shown in our calculated PI value (6.26 for wild type, 5.71 for one point mutation, and 5.31 for 2 points mutation, respectively). The charge selective of renal filtration seems to be one of responsible parameters for the plasma half-life enhancement of the muteins. Size selective renal filtration seems not to be responsible to this phenomenon as we observed no size different between wildtype and muteins as confirmed by size exclusion chromatogram (Figure 7). The glome rular basement membrane (GBM) which functions as a filter is covered by negatively charged matrix proteins: GBM-sulfate proteoglycans, i.e. agrin and perlec an. These matrix layers contribute to an anionic GBM barrier. The endothelial and podocytes may also form a part of a negative filtration barrier by membrane bound sulfate proteoglycan on their surface [16,17]. Although the mutein forms we developed did not result in superior improvement in the plasma half-life of hIFNα2b as compared to other groups [3-5], our approach is the first to report the successful amino acid substitution in relation to the plasma half-life improvement of hIFNα2b. In addition, two points mutation of hIFNα2b (C2D C99D mutein) showed higher liver uptake after 10 min of protein injection among other hIFNα2bs we studied, although minor. Renal accumulation increased significantly in all mutein forms as compared to wildtype, especially at 10 min. Decreasing accumulation both in liver and kidney at 30 min may due to redistribution to the blood as shown in Figure 9. As early interpretation, this seems to be charge-dependent uptake. Although ambiguous, our data on renal accumulation of mutein forms (Figure 9) was in line with Yamasaki et al. [18] result which reported high renal accumulation of aconylated BSA, the more negative charge of BSA. Further careful study must be performed to strengthen our biodistribution data.

Surprisingly, substitution of cysteins group with aspartic acids resulted in maintaining the activity of interferon with IC50 on both wild type and muteins interferon alpha2b was >5 µg/mL as shown in our preliminary in vitro data using HepG2 cells. This report confirmed the finding described by Inamura [19] and Tnani [20].

5. CONCLUSION
The muteins were successfully constructed and produced as 37 kDa fusion protein containing thioredoxin tag at its N terminus. The yields of muteins were ranging 8.8 - 8.9 mg/L from soluble protein and 22.8 - 22.9 from inclusion bodies. All muteins showed longer plasma circulation than the wild type form. In addition to the plasma half-life, the cysteineto aspartic acid substitution of hIFNα2b did not diminish the activity. Although the charge of hIFNα2b seems to be one of responsible parameters for the fate of hIFNα2b in vivo, further careful study is still required to confirm our finding.
REFERENCES


