The Structural Characterization of Recombinant Human Interferon Gamma

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Abstract: The recombinant human interferon gamma (rhIFN-γ) that is produced in E. coli by a method based on recombinant DNA technology, increases macrophage cytotoxicity. In this study to evaluate the structural characterization of rhIFN-γ, considered two conditions: fresh sample (4°C) and old sample (6 months in 25°C) of rhIFN-γ. Fresh sample (4°C) and old sample (6 months in 25°C) analyzed by analytical methods for determination of biological activity and structural analysis (deamidated and oxidized, heterodimers, monomer and aggregates, covalent dimer and SDS-PAGE). The structural characterization of them was compared.

Key words: rhIFN-γ, structural characterization, recombinant, E. coli

INTRODUCTION

The human interferons (hIFNs) are a family of proteins. The initial discovery of which was based on their ability to inhibit viral growth in target cells (Friedland, 1996). Their biological effects in vivo include antiviral activity, cell growth inhibitor and immunomodulator activity (Pitha, 2007; Dheda et al., 2005; Perez et al., 1990; Chen et al., 1992). IFN-γ is more potent than the other two IFNs (α,β) in its immunomodulatory activities. The antiproliferative effects make it a potentially useful against cancer (kidney cell carcinoma, colon cancer and rheumatoid arthritis). The IFN-γ is secreted by human antigen stimulated T-lymphocytes in response to viral infection and various other inducers. The native human interferon gamma is composed of 143 amino acid residues with a total molecular mass of 20-25 kDa. It is glycosylated and does not contain cysteine residues (Khalilzadeh et al., 2004; Mohammad-Mosaabadi et al., 2007). In 1986, hIFN-γ cDNA was cloned and expressed in Escherichia coli. This rh-IFN-γ produced in E. coli is not glycosylated and has methionine as its N-terminal residue. The total molecular mass of rh-IFN-γ is 17 kDa that it is physiologically active (Pitha, 2007). The protein consists of non covalent dimers of two identical monomers.

Recombinant proteins expressed in genetically transformed cells must accurately represent the natural molecules that they are intended to replace or complement. They should also fulfill the minimum requirements for biological efficacy, safety and quality criteria as do other preparations intended for pharmaceutical use (Zhang and Tong, 1992).

The deferent testing is done for evaluation of this recombinant product so as biological activity and structural analysis.

This study describes the characterization of final product of rhIFN-γ as structural analysis: covalent dimer and oligomer, deamidated, oxidized, monomer and aggregate form of hIFN-γ in fresh and old sample that described and comparison of the electrophoretic pattern and biological activity was done.

MATERIALS AND METHODS

Analysis of covalent dimer: Size Exclusion Chromatography (SEC) with HPLC method was used to check purity of rhIFN-γ and the presence of undesired form including covalent dimer. The column (7.5×300 mm, 10,000-300,000 Dalton), ultrapak TSK G3000 SW LKB was equilibrating with sodium phosphate buffer (0.2M, pH 6.8) at a flow rate of 1 mL min⁻¹ and mixture of molecular mass standards, BSA (66 kDa), trypsin inhibitor (20.1 kDa), lysozyme (14.4 kDa) as resolution solution, fresh sample and old sample of rhIFN-γ as tests solution were used. The effluent was monitored at 210 nm.

Analysis of deamidated and oxidized forms and heterodimers: Ion Exchange Chromatography (IEC) with HPLC method was used to check these forms of rhIFN-γ. The column (IEC sp-825 Shodex 8 mm ID × 75 mm L⁻¹) was equilibrated with ammonium acetate 0.05 M, pH 6.5 (buffer A) and developed with a linear gradient (100-0%).

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1087
(0-100%) ammonium acetate 1.2 M, pH 6.5 (buffer B) in a 
57 min period at flow rate 1.2 mL min⁻¹ in 35°C and tests 
solution (fresh and old) were injected and detected. The 
effluents were monitored at 280 nm.

**Analysis of monomer and aggregates:** The column
(7.5×300 mm 10,000-500,000 Dalton, ultropak TSK G3000 
SW LKB was used to check monomer and aggregates 
form of rhIFN-γ. After equilibrating of column with 
potassium chloride (1.2 M) in solution A (0.59 g L⁻¹ 
 Succinic acid and 40 g L⁻¹ mannitol pH 5) at flow rate 
0.8 mL min⁻¹, 20 µL of each, resolution solution (BSA + 
Irukio), tests solution (fresh and old) were injected and 
detected at 214 nm.

**Analysis of purity of rhIFN-γ by SDS-PAGE:** This 
analysis was carried out by SDS-PAGE. The test was 
performed under reducing conditions using resolving gels 
of 15% acrylamide and silver staining as the detection 
purities fresh and old samples of rhIFN-γ.

**Assay of biological activity:** The method relies on viral 
cytopathic effect (CPE) inhibition using semi-micro 
dyebinding technique in IFN-treated cells and comparing 
activity of rhIFN-γ with standard samples of IFN-γ.

The assay was performed by incubating a fixed count 
of vero (African green monkey kidney) cell line with serial 
dilutions of rhIFN-γ followed by challenging the cells with 
defined plaque-forming units of Vesicular Stomatitis Virus 
(VSV), all according to standardized procedure (Tsanev 
and Ivanov, 2001; Mohammadian-Mosaabadi et al., 2005).

The interferon activity was calculated as the reciprocal of 
the dilution in the well of the titre plate where 50% of the 
vero cell monolayer is protected from the CPE of 
challenging virus. The standard interferon gamma was 
used in this test is WHO standard as: Interferon gamma, 
human 88/606 (National Institute of Biological Standard 
and Control Blanche Lane, England).

**RESULTS AND DISCUSSION**

The rhIFN-γ is a single chain protein has a molecular 
mass of 17 kD. The active form of it is considered to a 
 homo dimer in which two monomers are none covalently 
bond in anti parallel Orientation (Tsanev and Ivanov, 
2001; Walsh Gary, 2003). So, in characterization of rhIFN-γ 
is necessary to determine possible covalent dimer forms 
as undesired product that form in the assessment 
of covalent dimers was done by gel filtration HPLC.

Figure 1 shows the fresh sample in 4°C of rhIFN lacks 
the covalent dimer forms but HPLC analysis of a rhIFN-γ 
stored for 6 months in 25°C, shows the peaks with 
approximately 35 kD and higher in comparison to 
molecular weight standards (BSA, Trypsin inhibitor, 
Lysozyme) that seems belongs to covalent dimer forms. 
Since, the rh IFN-γ has no cysteine residue; the forms are 
not disulfide linked dimer.

Since, one source of protein variability comes from 
oxidative protein modification to analysis of rhIFN-γ The 
examination of oxidized and deamidated forms of Protein 
was considered. Generally, these modified forms are 
produced during manufacturing process or storage of

![Fig. 1: Analysis of covalent dimer, comparison of graphs from (—), standard proteins which contains Bovine Serum Albumin (66 kD) peak No 1, Soybean Trypsin Inhibitor (20.1 kD) peak No 2 and lysozyme (14.4 kD) peak No 3 and (-----), fresh gamma interferon with any peak and shoulder before main peak, (—), old sample (6 mounts in 25°C) gamma interferon](image-url)
proteins. In order to determination of deamidated and oxidized forms, the fresh sample in 4°C of rhIFN and the old sample stored 6 months in 25°C was examined by strong cation exchange chromatography. The data are shown in Fig. 2, in old sample principal peak (rhIFN-gamma) is about 27.4 min and heterodimers 25.4 min, deamidated and oxidized forms in 26.2 min were eluted at shorter retention time, relative to principal peak. Fresh sample contains any unmodified (deamidated, oxidized, heterodimers) gamma interferon.

In the purification process of rhIFNg during unfolding and refolding reactions, Some portion of monomers doesn’t assemble in to active dimer forms and remain as monomers forms. Protein aggregation is a common issue encountered during manufacture of bio pharmaceutical drug. Although the aggregated forms are removed during purification process by SEC chromatography and so on but it seems, some these forms because of stresses can lead to protein aggregation, are observed in final product. So, the rhIFNg was analyzed by analytical SEC based on difference in size of monomer and aggregates that the peaks of monomer, native dimer and aggregated are isolated. Figure 3 shows that fresh sample is followed

Fig. 2: Analysis of deamidated and oxidized forms and heterodimers, comparison of graphs from old sample (6 months in 25°C) gamma interferon (—) which contains principal peak (rhIFN-gamma) is about 27.4 min and Heterodimers (25.4 min). Deamidated and oxidized forms (26.2 min) were eluted at shorter retention time, relative principal peak, with (—) fresh gamma interferon without unmodified form

Fig. 3: Analysis of monomer and aggregates, comparison of graphs from standard proteins (—), which contains Bovine Serum Albumin (66 kD) with retention time equal to 8.85 min, commercial gamma interferon with retention time 10.13 min, with (----) rhIFN gamma and (—) old sample (6 months in 25°C) rhIFN gamma
the pattern of commercial rhIFNγ in standard sample that contains native dimer form without any monomer and aggregate form. The old sample has a little aggregate form and no monomer.

In Fig. 4, the comparison of electrophoretic pattern is shown that the old sample has a band between 35 and 25 kD bands of standard sample that it may be is related to covalent dimer and a band between 14.4 and 18.8 kD region is belonged to monomer form of IFN in while the fresh sample has a one band in monomer region.

The most convenient method for interferon assay is quantitating its antiviral activity or inhibition of cytopathic effect (CPE). In this research, we looked for a fast and sensitive method to quantitative antiviral activity of this cytokin based on cytopathic effect/dye uptake. After standardization of method, activity of fresh and old IFN-γ was compared to standard sample (WHO standard) (Fig. 5). In regard to biological activity of standard sample, 20×10⁶ IU mg⁻¹, the biological activity of fresh sample is near the WHO standard, but biological activity of old sample approximately two times decreased. Antiviral activity of rhIFN gamma in vero cell line shown in Fig. 6.

According to obtained results, we can observe that rhIFN-γ is a sensitive protein. Modifications of rhIFN-γ such as deamidated and oxidized, monomer, covalent dimer and aggregate happened in during life time (0 and 6 months) and temperature change (4 and 25°C). These

Fig. 4: SDS-PAGE analysis of rhIFN-gamma. Lane 1 is MW standards from MBI Fermentes Company which contain β-galactosidase 116.0 kD, BSA 66.2 kD, ovalbumin 45.0 kD, LDH 35 kD, Bsp981 restriction endonuclease 25 kD, β-lactoglobulin 18.4 kD and lysozyme 14.4 kD. IFN-gamma band should stand between the last two bands. Lane 2 is old sample (6 months in 25°C) rhIFN-gamma and Lane 3 fresh rhIFN-gamma

Fig. 5: Determination of anti-viral activity of IFN-gamma
Fig. 6: Antiviral activity of rhIFN gamma in vero cell line. Normal Vero cell line without any treatment (a), Vero cell line that treated with 500 ng mL⁻¹ of IFN-γ(WHO) and challenged with 1/50,000 dilution of VSV stock after 48 h (b), Vero cell line that treated with 500 ng mL⁻¹ fresh sample of IFN-γ (4°C) and challenged with 1/50,000 dilution of our VSV stock after 48 h (c), Vero cell line that treated with 500 ng mL⁻¹ old sample of IFN-γ (6 months in 25°C) and challenged with 1/50,000 dilution of this VSV stock after 48 h (d).

modifications change the structure of rhIFN-γ and decrease approximately two times biological activity. It may be these assessments are so necessary for this protein using as a recombinant pharmaceutical drug.

REFERENCES


