Different solid sample preparation methods affecting the spectral similarity of salmon calcitonin

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Abstract. Salmon calcitonin (sCT) was selected as a model protein drug for investigating its structural similarity in the solid state by four sample preparation methods, such as tape, smeared, CaF2 and film methods. The conformational changes of sCT in the solid state were estimated by using a second-derivative Fourier transform infrared (FT-IR) microspectroscopy. The tape method was acted as a standard reference.

The value of correlation coefficient (r) for smeared method was higher than that of other method, indicating that a novel technique by smearing sCT powder on the surface of KBr pellet was the best optimal sample preparation method.

Keywords: Salmon calcitonin, FT-IR, sample preparation, second derivative, structural similarity

1. Introduction

The therapeutic activity of protein drug is well known to be highly dependent on their conformational structure, how to keep the structural integrity and active conformation of protein drug in the manufacturing processes of production, or shipping and long-term storage of products are the highlighted and critical issues [4,7]. Because the secondary structure is one of the most important conformational information for a protein, thus the secondary structure prediction and determination of proteins are the important events [10,20,25]. Many analytical techniques have been applied to determine the secondary structure of protein [18,22,23], Fourier transform infrared (FT-IR) spectroscopy is one of the most common spectroscopic techniques to quickly examine the conformational changes of protein structure in the solid and liquid states [1,8,16].

Since protein drug formulations are more stable in the solid state than in the liquid state, the solid-state protein products have often been manufactured [8,15,24]. Before FT-IR determination of protein secondary conformation in the solid state, the solid-state protein sample was commonly mixed and ground with KBr powder, and then compressed in a mechanical die press to form a translucent pellet. These two processes of grinding and compression might cause the additional protein structural alterations [6,14], implying that the sample preparation method for protein in the solid state plays an important role during FT-IR determination. In order to avoid both processing effects, a unique solid sample preparation method is needed.

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Calcitonin (CT), a 32-amino acid linear polypeptide hormone, is always used for the therapy of different bone diseases [2,17]. Among different types of CT available for clinic use, salmon calcitonin (sCT) is one of the most potent forms [2,9]. Very little has been reported regarding the structural stability of sCT formulations in the solid state. In this study, we used sCT as a model protein drug to preliminarily examine the influence of different sample preparation methods on the conformational structure of sCT in the solid state.

2. Materials and methods

2.1. Materials

Salmon calcitonin (sCT) was purchased from Polypeptide laboratory A/S, Denmark. The KBr crystals were obtained from Jasco Parts Center (Jasco Co., Tokyo, Japan). Aluminum foil was purchased from Reynolds Metals (VA, USA).

2.2. Sample preparation methods

(1) Tape method: A tiny sample of sCT powder was partly adhered with adhesive tape and fixed on the edge of glass plate.
(2) Smeared method: A trace amount of sCT powder was carefully smeared on the surface of KBr pellet without any compression.
(3) CaF$_2$ method: One drop of 1% (w/v) sCT aqueous solution was dropped on the surface of CaF$_2$ plate and stored at 25°C, 50% relative humidity (RH) condition. After storage for 1 day, the cast film on the foil was formed.
(4) Film method: One drop of 1% (w/v) sCT aqueous solution was dropped on the aluminum foil and stored at 25°C, 50% relative humidity (RH) condition. The cast film on the foil was formed after storage for 1 day.

All the solid samples were stored at 25 ± 2°C, 60 ± 5% RH condition before spectral analysis.

2.3. FT-IR microspectroscopic studies of different sCT samples

Each IR spectrum of sCT sample prepared by tape, smeared or CaF$_2$ method was determined by FT-IR microspectroscopy (Micro FTIR-200, Jasco Co., Japan) equipped with a mercury cadmium telluride (MCT) detector using a transmission technique, according to our previous studies [11,12]. All the spectra were obtained at a 4 cm$^{-1}$ resolution and at 100 scans. However, the IR spectrum of sCT film prepared by a film method was also determined by using a reflectance technique [13]. The reflectance IR spectra were collected at an angle of incidence centered at 30°. All the determinations were undertaken at 25 ± 2°C and 60 ± 5% RH condition.

2.4. Data acquisition and handling

2.4.1. Spectral analysis

A software of spectral manager for window (Jasco Co., Tokyo, Japan) and GRAMS spectroscopy software suite (Version 7, Thermo Electron Co., MA, USA) were used for data acquisition and handling. Second-derivative spectral analysis was applied to locate the position of the overlapping components in the amide bands and assigned to different secondary structures [21].
2.4.2. Structural similarity

In order to quantify the structural similarity of sCT samples prepared by different methods, the spectral correlation coefficient analysis between two second-derivative IR spectra was applied. A mathematical procedure proposed by Prestrelski et al. was used to calculate the spectral correlation coefficient ($r$) between two second-derivative IR spectra as follow [19]:

$$ r = \frac{\sum_{i}^{n} x_i y_i}{\sqrt{\sum_{i}^{n} x_i^2 \sum_{i}^{n} y_i^2}} $$

where $x_i$ and $y_i$ are the spectral absorbance values of the reference and comparison spectra respectively, at the $i$th frequency position in the amide I region.

The tape method was acted as a standard reference, since this method maintained a native form of sCT without any treatment. The $r$ value provides a measure to compare each IR spectrum of a given sCT sample prepared by one of different methods to that of sCT sample prepared by a tape method. All the spectral comparisons were performed in the amide I region (1700–1600 cm$^{-1}$). Each spectrum was baseline-offset corrected and area-normalized. Comparison of identical spectra gives a value of 1.0. The larger the changes in conformation detected, the greater the difference between two spectra obtained, leading to a smaller $r$ value.

3. Results and discussion

Infrared spectra of sCT sample prepared by four different methods (tape, smeared, CaF$_2$ and film) within the range of 3700–2800 and 1800–1000 cm$^{-1}$, are shown in Fig. 1. It is evident that all the IR spectra from four sample preparation methods seemed to be similar. The peaks at 3296–3230 cm$^{-1}$ were originated from the vibrations of NH stretching mode of sCT. The peaks near 2957–2959 cm$^{-1}$ were due to the asymmetric CH$_3$ and CH$_2$ stretching bands of sCT, while the peaks at 2871 cm$^{-1}$ was associated

Fig. 1. Infrared spectra of sCT prepared by four different methods within the range of 3700–2800 and 1800–1000 cm$^{-1}$. Key: (a) tape method; (b) smeared method; (c) CaF$_2$ method; (d) film method.
with the symmetric CH$_3$ stretching mode of the side chains of sCT. The amide I, II and III bands were located at 1657–1658, 1544–1547 and 1254–1256 cm$^{-1}$, respectively. The peaks at 1407–1409 cm$^{-1}$ were due to the symmetric COO$^-$ stretching band and/or the deformation of CH$_2$ and CH$_3$. The shoulders at 1081–1083 cm$^{-1}$ were mainly corresponded to the contributions of C–O and C–N stretching modes [3,5]. Because all samples exhibited the similar IR spectra, suggesting that four sample preparation methods did not markedly alter the conformational structure of sCT in the solid state.

The amide I band in the IR spectrum is particularly more sensitive to protein secondary structure than other amide bands, since the amide I band arising from the C=O and N–H groups is susceptible to hydrogen bonding and coupling between transition dipole of adjacent peptide bonds in the structure of proteins. Thus the amide I band is the most suitable probe used to differentiate different secondary structures of protein. The second-derivative spectra have been applied to resolve the overlapping bands within the amide I band and to express their secondary structure [21]. Since the height of a second-derivative IR peak is proportional to the square of the original peak height with an opposite sign, the position and height of a second-derivative peak may quantitatively reflect the secondary structure of sCT. To further verify the structural similarity of sCT prepared by different sample preparation methods, a spectral correlation coefficient analysis between two second-derivative amide I spectra was carried out. The IR spectrum of sCT prepared by tape method was compared with that of the IR spectrum of sCT prepared by smeared, CaF$_2$ or film method using the second-derivative spectra in amide I region (Fig. 2A).

![Fig. 2. Second-derivative amide I spectra of sCT prepared by different methods (A) and correlation of peak intensity of sCT prepared by tape method with that of the peak intensity of sCT prepared by smeared, CaF$_2$ or film method using the second-derivative spectra in amide I region (B–D). Key: (A) baseline corrected and area-normalized spectra; (B) tape method vs. smeared method; (C) tape method vs. CaF$_2$ method; (D) tape method vs. film method.](image-url)
This was accomplished by calculating the correlation coefficient ($r$) for the spectrum of sCT by using tape preparation method as a reference. It is apparent that the calculated $r$ value was 0.983 for smeared method vs. tape method, 0.957 for CaF$_2$ method vs. tape method, or 0.956 for film method vs. tape method, respectively (Fig. 2B–D). The correlation coefficients were very close to unity, suggesting no appreciable structural change. In other word, four sample preparation methods used here did not cause the additional sCT structural alterations, suggesting that these four sample preparation methods may be suitable for studying the protein secondary structure whether in powder or film form. Because the sCT powder was a native form (tape method) and the $r$ value of smeared method was higher than that of the CaF$_2$ method or film method, indicating that by smearing sCT powder on the surface KBr pellet (smeared method) was the best optimal sample preparation method.

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References


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