ISOLATION AND AUTOXIDATION PROFILE OF FISH MYOGLOBIN FROM HOKI (MACRURONUS MAGELLANICUS)

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Abstract : Myoglobin is known to be present exclusively in cardiac and red skeletal muscles, but not in white skeletal muscles. Thus, to date, only a few studies on myoglobin from fish species with white flesh have been reported. For comparative examination, we directly isolated myoglobin from cardiac muscle of hoki (Macruronus magellanicus), one of the most important commercial fish species with white muscle. The ferrous myoglobin was separated from its ferric met-form by anion exchange column chromatography. The absorption spectra of hoki myoglobin were similar to those of bigeye tuna skeletal myoglobin, in both oxy- and met-forms. However, hoki oxymyoglobin was found to be susceptible to autoxidation in 0.1 M buffer (pH 7.2) at 25°C, with its rate being more than 3 times higher than that of bigeye tuna oxymyoglobin.

Key words : fish myoglobin, hoki, oxymyoglobin, autoxidation

INTRODUCTION

In vertebrates, myoglobin (Mb), a mobile carrier of oxygen, is known to be present in cardiac and red skeletal muscles, and plays an important role in maintaining aerobic metabolism both as an oxygen store and as a facilitator of oxygen diffusion1). Studies of teleost fish Mbs have provided important information on molecular evolution of vertebrate Mb, because of their remarkable species diversity compared to any other class of vertebrate2). Structural and stability properties of fish Mb has been investigated mainly in fish with red muscles, such as tuna and mackerel, in which Mb is found in high concentration2-5). Only a few studies, however, have reported on Mb from fish species with white skeletal muscle6).

Hoki (Macruronus magellanicus), which is closely related to cod, is one of the most important commercial fish species with white muscle. It is found in the Southeast Pacific and Southwest Atlantic Oceans, off the coast of southern Chile and Argentina, at depths ranging from 60 to 600 m7).

In this paper, we described procedures for isolating oxygen binding ferrous Mb (oxyMb) directly from hoki cardiac muscle, which contains large amounts of Mb, almost comparable to those in mammalian cardiac muscles. Using this native preparation, we examined for the first time the autoxidation rate of hoki oxyMb to ferric Mb (metMb).

MATERIALS AND METHODS

Oxymyoglobin preparation

According to the purification procedure for tuna Mb8), hoki Mb from cardiac muscles was prepared at low temperatures (0-4°C). In order to avoid rapid autoxidation of oxyMb, CO-saturated buffers were used during the procedure. The frozen hearts (ca. 50 g) were thawed quickly and minced twice in a 2-fold volume (w/v) of 10 mM Tris–HCl buffer (pH 8.7) containing 1 mM EDTA. After extraction for 30 min, insoluble materials were removed by centrifugation. The extract was then fractionated between 70% and 90% saturation with ammonium sul-
fate at pH 8. The Mb precipitate was centrifuged, dissolved in a minimum volume of 2 mM Tris-HCl buffer (pH 8.7) containing 1 mM EDTA, and dialyzed against the same buffer. The crude Mb solution was then passed through a Sephadex G-50 (GE Healthcare Japan) column (4.5×90 cm) with a 2 mM Tris-HCl buffer (pH 8.7) containing 1 mM EDTA to separate Mb from hemoglobin. The hoki Mb was applied to a DEAE-cellulose column, which had been equilibrated with a 2 mM Tris-HCl buffer at pH 8.7. After contaminant proteins were washed out completely, metMb was eluted with a 5 mM Tris-HCl buffer (pH 8.7) for the first. Then, the ferrous CO-form was eluted out with a 10 mM Tris-HCl buffer (pH 8.7), and immediately stored in liquid nitrogen until use. For autoxidation experiments, hoki oxyMb was produced from the CO-form in an O₂-saturated buffer (pH 8.7), by flash-photolysis for 3 min using 300-watts photoreflector lamp (Toshiba) in an ice bath (0–4°C). The concentration of the Mb was determined, after conversion into cyano-met-form, using an absorption coefficient of 9.0 mM⁻¹cm⁻¹ at 540 nm. This coefficient value was obtained on the basis of the pyridine hemochromogen method.

Autoxidation rate measurement

The rate of autoxidation of oxyMb was measured in a 0.1 M buffer (pH 7.2) at 25°C, according to our standard procedure. For example, a 900 µl solution of the appropriate buffer was placed in a test tube and incubated in a water bath maintained at 25 ±0.1°C. The reaction was started by adding 100 µl of fresh oxyMb solution (100–150 µM), and the changes in the absorption spectrum over 450–700 nm were recorded at measured intervals of time. For the final state of the runs, oxyMb was completely converted into its ferric met-form by the addition of potassium ferricyanide. Absorption spectra were recorded on a Hitachi U-3300 spectrophotometer equipped with a thermostatically controlled cell holder (within ±0.1°C).

RESULTS

Oxymyoglobin preparation

In the procedures for isolating native ferrous Mb, an essential step was the chromatographic separation from ferric metMb by a DEAE-cellulose column. The metMb was eluted out with a 5 mM buffer, and then ferrous CO-Mb was obtained with a 10 mM buffer (Fig. 1). Obtained oxy- and met-Mbs showed a single band in SDS-polyacrylamide gel electrophoresis, respectively, with the molecular mass of about 15 kDa. The oxygenated form was produced from the CO-form by flash photolysis in an O₂-saturated buffer (pH 8.7). The spectrum of the met-form was obtained at pH 7.0. The expanded scale was used for the visible spectra.

Fig. 1. DEAE-cellulose column chromatography of hoki myoglobin. After gel filtration, Mb solution was applied to a DEAE-cellulose column (2.0×10 cm) equilibrated with a 2 mM Tris-HCl buffer (pH 8.7) under O₂-saturated condition. The metMb was eluted out with a 5 mM buffer, and then Mb-CO was eluted with a 10 mM buffer.

Fig. 2. Absorption spectra of the hoki oxyMb (bold line) and metMb (broken line). The concentration was 10 µM for each form. The oxy-form was produced from the CO-form by flash photolysis in an O₂-saturated buffer (pH 8.7). The spectrum of the met-form was obtained at pH 7.0. The expanded scale was used for the visible spectra.
CHARACTERIZATION OF HOKI MYOGLOBIN

The \( \gamma \)-peak of hoki metMb was higher than that of the oxyMb with an absorbance ratio of \( \gamma_{\text{met}}/\gamma_{\text{oxy}} = 1.25 \).

Stability property of hoki oxymyoglobin

Under air-saturated conditions, oxyMb (MbO\(_2\)) is oxidized easily to metMb with generation of the superoxide anion as

\[
\text{MbO}_2 \xrightarrow{k_{\text{obs}}} \text{metMb} + \text{O}_2^-. \tag{1}
\]

where \( k_{\text{obs}} \) represents the first-order rate constant observed at a given pH\(^{10} \). Therefore, the rate of autoxidation is given by

\[
-\frac{d[\text{MbO}_2]}{dt} = k_{\text{obs}} \cdot [\text{MbO}_2]. \tag{2}
\]

Fig. 3A shows such example of spectral changes with time for the autoxidation reaction of hoki oxyMb in a 0.1 M phosphate buffer (pH 7.2) at 25°C. This process of autoxidation was followed by a plot of experimental data as a \(-\ln([\text{MbO}_2]/[\text{MbO}_2]_0)\) versus time \( t \), where the ratio of oxyMb concentration after time \( t \) to that at time \( t=0 \) can be monitored by the absorbance changes at 577 nm (\( \alpha \)-peak of hoki oxyMb). Its first-order plot, Fig. 3B, is also shown to obtain the rate constant of \( k_{\text{obs}}=9.1\times10^{-2} \) h\(^{-1} \) from the slope of the line. Regarding the half-life for conversion of oxyMb to metMb, this corresponded to 8 h for hoki fish.

DISCUSSION

The reversible and stable binding of oxygen to the heme iron(II) is the basis of Mb function. However, the oxygenated form of Mb is known to be oxidized easily to ferric metMb, which cannot bind molecular oxygen and is therefore physiologically inactive, with generation of the superoxide anion\(^{10,11} \). Using a variety of native oxyMbs isolated directly from various sources, the reaction mechanism for autoxidation of oxyMbs has been investigated\(^{12,13} \).

In the present study, we directly isolated Mb from the cardiac muscle of hoki fish and examined its autoxidation reaction. Because of its rapid autoxidation, it was essential to convert the ferrous form of hoki Mb into CO-form by the use of CO-saturated buffers prior to the purification. The native oxygenated form was produced from the obtained CO-form by flash photolysis\(^8 \) in O\(_2\)-saturated buffer.

The absorption spectra of hoki Mb were similar to those of bigeye tuna Mb\(^{31} \). The \( \gamma \)-peak of hoki metMb was higher than that of the oxyMb (Fig. 2). Shikama and Matsuoka\(^14 \) have examined a dozen Mbs from various species for their spectrophotometric properties, and found that the proteins can be divided into two groups on the basis of the absorbance ratio of the \( \gamma \)-peak of the acidic met-form to that of the oxy-form, that is to say, the \( \gamma_{\text{met}}/\gamma_{\text{oxy}} \) ratio. Values higher than 1.0 were obtained for vertebrate Mbs containing the distal histidine, whereas those of less than 1.0 were seen for the ratio of the Mbs lacking this residue. It should be noted that the distal histidine, which is highly conserved in almost
vertebrate Mbs, plays an important role in stabilization of the oxygen molecules bound to the heme iron(II) by hydrogen-bond formation. Judging from this spectral criterion, a value of $\gamma_{\text{met}}/\gamma_{\text{oxy}}=1.25$ for hoki Mb indicates that this heme protein has the distal histidine residue.

On the other hand, it was revealed that the autoxidation rate of hoki oxyMb was higher than those of bigeye tuna oxyMb ($2.9 \times 10^{-2} \text{ h}^{-1}$) in a 0.1 M buffer at 25°C. Madden et al. compared the Mbs autoxidation rates of several fish species and found that the rates were closely related to body temperature; that is, Mbs from fish species with higher body temperatures were more stable to autoxidation at pH 7.5, 37°C. As the body temperature of hoki, which inhabits deep sea, may be lower than that of tunas (approximately 20–30°C), our result is consistent with their report.

When compared with usual Mbs containing the distal histidine residue which had been examined so far, hoki oxyMb is found to be susceptible to autoxidation in a 0.1 M buffer (pH 7.2) at 25°C. The autoxidation rate differences are considered to be caused by the hydrophobicity of heme pocket. In addition, it has been suggested that the autoxidation property of fish Mbs is associated with the instability of its overall molecular structure. In order to understand the stability properties of hoki oxyMb in detail, analysis of the pH dependency of autoxidation rate as well as the determination of the primary sequence of hoki Mb is required.

REFERENCES