Study on the Interaction of Retinoic Acids to Human Serum Albumin by Fluorescence and Circular Dichroism Spectroscopy

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Abstract

Retinoic acids (RAs) are considered to be endocrine disruptor chemicals and toxic environmental priority pollutants. In this paper, the interactions between RAs and human serum albumin (HSA) were examined by steady state fluorescence, time-resolved fluorescence and circular dichroism spectroscopy (CD). The RAs quenched the fluorescence of the protein remarkably and the mechanism of quenching was found to be static in nature. Synchronous fluorescence studies suggested that the polarity around the tryptophan (Trp) residues and tyrosine (Tyr) residues was not altered in the presence of RAs. The thermodynamic parameters of the binding reactions (ΔGθ, ΔHθ, ΔSθ) were measured, and they indicated the presence of hydrophobic forces and hydrogen interactions in the RAs–HSA interactions. The alterations of HSA secondary structure in the presence of RAs were confirmed by CD and time resolved fluorescence spectroscopy.

Keywords: Retinoic acids; Human serum albumin; Fluorescence; Circular dichroism

Introduction

Retinoic acids (RAs) are essential for physiological processes such as reproduction, cell proliferation and differentiation, vision, and embryonic development [1]. However, extensive research has proved that RAs are among the most potent known animal teratogens [2]. Exposure of embryos to exogenous RAs causes a spectrum of malformations, including defects of the neural tube and central nervous system; skeleton, palate, ear, and other craniofacial malformations; defects in the heart, thymus, and urogenital system; as well as missing or duplicate limbs and digits in the offspring of humans, rodents, chickens, the African clawed frog (Xenopus laevis), and fish [3–10]. When frogs were exposed to RAs during metamorphosis, a suite of abnormalities, such as reductions and deletions of the hind limb, bony triangles, and eye deformities, could occur [11–15]. These abnormalities observed in the laboratory are similar to those found in eutrophic habitats [16,17].

Human serum albumin (HSA) is the most abundant protein in plasma, which functions in the maintenance of colloid osmotic blood pressure and in the binding and transportation of various ligands such as fatty acids, hormones, and drugs, then transports them between tissues and organs [18]. It has been shown that the distribution, free concentration, and metabolism of various ligands can be significantly altered as a result of their binding to HSA [19]. Ligand interactions at the protein binding level will in most cases significantly affect the apparent distribution volume of the ligands and also affect the elimination rate of ligands. Therefore, investigating the interactions of RAs and HSA are significant for knowing their transports and distributions in the body and clarifying their action mechanisms. As of yet, however, no work has been reported for the mechanism of these interactions and the detailed physicochemical characterizations of RAs binding to HSA.

In this paper, we present a spectroscopic analysis of the interaction of HSA with four RAs (Figure 1) such as Retinoids (RED), Acitren (ACE), all-trans-Retinal (REA) and 4-keto all-trans-retinoic acid (REO) in aqueous solution at physiological conditions, using constant protein concentration and various RAs compositions. Structural information regarding RAs binding mode and the effects of RAs–HSA complexation on the protein stability and secondary structure are reported here.

Keywords: Retinoic acids; Human serum albumin; Fluorescence; Circular dichroism

Materials

HSA (fatty acid content <0.05%), Retinoids, Acitren, all-trans-Retinal, 4-keto all-trans-retinoic acid were purchased from the Sigma Chemical Company and used without further purification. All other reagents were of analytical reagent grade. Ultrapure water was used throughout the experiments. The solution of HSA was prepared in phosphate buffers (1 μM, pH 7.40).

Apparatus and methods

All the steady-state fluorescence spectra were recorded on a LS-50 B spectrofluorimeter (Perkin Elmer, USA) equipped with 1.0 cm quartz
cells. The excitation wavelength was 280 nm and the slit widths of
emission and excitation were set 5.0-5.0 nm.

Synchronous fluorescence spectra of were measured under the
same conditions as the steady-state fluorescence. For synchronous
fluorescence, Δλ (λem-λex) were set at 15 nm and 60 nm, respectively.

The fluorescence lifetime measurements were done in a Horiba
Jobin Yvon Fluoro Max-4 time correlation single photon counting
(TCSPC) system, using a 265 nm nanosecond diode laser. The time
ranges are 0.11 ns/channel, in 4096 effective channels. Data were globally
fitted to the appropriate exponential model after deconvolution of the
instrument response function by an iterative deconvolution technique,
using the IBH DAS6 fluorescence decay analysis software.

CD measurements were taken with an Applied Photophysics
Chirascan circular dichroism spectrometer using a 1 cm quartz cell. The
wavelength interval is 200-260 nm.

Results and Discussion

Steady-state fluorescence spectra

Fluorescence spectroscopy is widely employed to study proteins
and peptides. The intrinsic fluorescence of HSA is very sensitive to
its microenvironment. Actually, the intrinsic fluorescence of HSA is
almost contributed by tryptophan (Trp) alone, because phenylalanine
has a very low quantum yield and the fluorescence of tyrosine (Tyr)
is almost totally quenched if it is ionized or near an amino group, a
carboxyl group, or a tryptophan [20]. When local surroundings of HSA
were altered slightly its intrinsic fluorescence would weaken obviously,
such factors as protein conformational transition, biomolecule binding,
and denaturation, etc. are responsible for the weakening. Figure 2A-2D
showed the fluorescence emission spectra (λex=280 nm) obtained for
HSA at pH 7.40 with the addition of different RAs compounds. It can be
seen that the fluorescence intensity of HSA decreased in the presence
of RAs. The measurements of intensity and position (due to the change
of polarity around the fluorophore molecule) are useful to learn the
microenvironment of amino acid residues. The fluorescence change
indicates that all four RAs compounds can change the structure of HSA.

Fluorescence quenching studies

Fluorescence quenching can be dynamic, resulting from collisional
encounters between the fluorophore and the quencher, or static,
resulting from the formation of a ground-state complex between the
fluorophore (protein) and the quencher. In both cases, molecular
contact is required between the fluorophore and the quencher for
fluorescence quenching to occur [21]. Dynamic and static quenching
can be distinguished by their difference depending on temperature
[22]. Higher temperature results in faster diffusion and larger amounts
of collisional quenching. This will typically lead to the dissociation of
weakly bound complexes. Therefore, the quenching constant increases
with the increase in temperature for dynamic quenching. It decreases,
however, with increasing temperature for static quenching.

The fluorescence quenching data were analyzed by using the Stern–
Volmer equation [23]:

\[ F/F = 1 + K_q [Q] = 1 + K \tau_s [Q] \]  

where \( F \) and \( F \) are the fluorescence emission peak area of HSA in the
absence and presence of RAs, respectively. \( K_q \) and \( K \) are the Stern–
Volmer quenching constant and the quenching rate constant of HSA,
respectively. \( \tau_s \) is the average lifetime of HSA without RAs (\( \tau_s = 10^{-8} \) s)
[24].

Figure 3 shows the fluorescence quenching curves of HSA by RAs following the Stern–Volmer plots at various temperatures. For
all four RAs, the fluorescence quenching curves have favorable linear
relationships, and the slopes of the quenching curves decreased with the
increasing of temperature. This indicates that the quenching mechanism
between RAs and HSA is the static quenching. The quenching constants
\( K_q \) and \( K \) at different temperatures are listed in Table 1. For all four
RAs, the \( K_q \) values decrease with increasing temperature and the calculated \( K_q \) was far greater than the maximum diffusion-
controlled quenching rate constant of various quenchers with the biopolymer (10^{-12}
Lmol^{-1}s^{-1}). Both results imply that the quenching is not initiated by
dynamic collision but formation of compound.

The \( K_q \) of RAs-HSA systems at 298 K is Acitren>Retinal>Retinoic
acid>Retinoids. We can conclude that the binding force of Acitren in
vivo is more stable than Retinal, retinoic acid, Retinoids.

Binding parameters of interaction

When small molecules bind independently to a set of equivalent sites
on a macromolecule, the binding constant (\( K_b \)) and possible number of
binding sites (\( n \)) can be determined from the following equations [25]:

\[ \log\left(\frac{F_0 - F}{F}\right) = \log K_b + \log[Q] \] (2)

In above equations, [Q] is the RAs concentration. \( K_b \) and \( n \) are the
binding constant and the number of binding sites, respectively. \( K_b \) and \( n \) can be obtained from the slope and intercept of the plots of \( \log(F_0-
F)/F \) versus \( \log[Q] \). The binding constants \( K_b \) and binding sites \( n \) were
listed in Table 2. The values of \( K_b \) indicate that there is a great binding
between RAs and HSA. The number of binding sites \( n \) of RAs and HSA
at is approximately 1.0, which indicates that there are 1.0 binding sites
in HSA for RAs during their interaction.

Generally, the forces acting between a drug and a biomolecule
may include hydrogen bonds, van der Waals forces, electrostatic and
hydrophobic interactions. Ross and Subramanian have summarized
thermodynamic laws to estimate the above forces as follows: If \( \Delta H > 0, \Delta S > 0 \),
the main force is hydrophobic interaction. If \( \Delta H < 0, \Delta S > 0 \), the main force is electrostatic interactions. If \( \Delta H > 0, \Delta S < 0 \), the main forces are van der Waals and hydrogen bonds [26]. If the temperature changes
little, the reaction enthalpy change is regarded as a constant. In order
to get the detail information of interaction force, the reaction enthalpy
change and entropy change were calculated according to the following
equations [27]:

\[ \ln[K_b/K_{b0}] = (1/T_1 - 1/T_2)/(\Delta H^R)/R \] (3)

\[ \Delta G^\theta = \Delta H^\theta - \Delta S^\theta = -RT\ln K_{b0} \] (4)

The negative of \( \Delta G \) indicates that the interaction process of
RAs interact with HSA are spontaneous. The positive of \( \Delta H \) and \( \Delta S \)
indicate that hydrophobic interaction plays major role. Based on the
hydrophobicity of the amino acid residues of trypsin and the bulk
hydrophobic group benzene ring of RAs, it was tempting to speculate a
direct hydrophobic interaction between the aromatic ring of RAs and
the hydrophobic amino acid residues. The \( \Delta s \) could enter into the
hydrophobic pocket of trypsin. In addition, the hydrogen bonding
between the OH of RAs and some carboxyl of HSA backbone or some
π-π stacking between aromatic ring of RAs and aromatic residues in
the hydrophobic pocket of trypsin. In addition, the hydrogen bonding
between RAs and HSA is the static quenching. The quenching constants
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the hydrophobic amino acid residues. The \( \Delta s \) could enter into the
hydrophobic pocket of trypsin. In addition, the hydrogen bonding
between the OH of RAs and some carboxyl of HSA backbone or some
π-π stacking between aromatic ring of RAs and aromatic residues in
trypsin cannot be excluded.

Synchronous fluorescence spectra studies

Synchronous fluorescence spectra studies are sensitive techniques
to detect the conformation changes. It is typical to study changes of the
molecular microenvironment near the fluorophore groups. When \( \Delta \lambda

Figure 2: (A) Fluorescence spectra of HSA and RED, [RED]=0, 0.80, 1.60, 2.40, 3.20, 4.00 μM; (B) fluorescence spectra of HSA and ACE, [ACE]=0, 0.80, 1.60, 2.40, 3.20, 4.00 μM; (C) Fluorescence spectra of HSA and REA, [REA]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM; (D) fluorescence spectra of HSA and REO; [REO]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM; [HSA]=5 × 10⁻⁶ μM, T=298 K.

Figure 3: Emission quenching curves of HSA with increasing concentration of RAs: (A) RED; (B) ACE; (C) REA; (D) REO. at different temperatures (298 and 310 K).
sets at 20 nm and 80 nm, the synchronous fluorescence spectra give the typical characteristics of tyrosine (Tyr) and tryptophan (Trp) residues [28]. Figure 4 shows the fluorescence intensity of both Tyr and Trp decreased regularly with the addition of RED, ACE, REA and DEO. Both of the fluorescence decrease of Trp residues are obvious greater than Tyr residues. It is concluded the binding sites are all near the Trp residues [29]. Absence of shifts implies that the interaction does not alter notably the polarity around these residues.

(B) synchronous fluorescence spectra of HAS and ACE,
[ACE]=0, 0.80, 1.60, 2.40, 3.20, 4.00 μM;
(C) synchronous fluorescence spectra of HAS and REA,
[REA]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM;
(D) synchronous fluorescence spectra of HAS and REO;
[REO]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM;
[HAS]=5 × 10^{-4} μM, T=298 K.

**Time-resolved fluorescence spectra**

Time-resolved fluorescence measurements can give information about the conformational heterogeneity of proteins. In order to further substantiate the quenching mechanism of the four RAs to HAS, fluorescence lifetime of the tryptophan of HAS were ascertained in the absence and presence of the four RAs. The fluorescence decay of HAS is reported to be triple-exponential in the literature [30]. The average fluorescence decay lifetime could be calculated according to following equations [31,32]:

\[ f_i = B_i \exp \left( -\frac{t}{\tau_i} \right) \]

\[ < \tau > = \sum_i B_i \tau_i \]

\[ I_{f} = \sum_i f_i \exp (-t/\tau) \]

In above equations, \( \tau \) is the lifetimes of the different components to the total decay, \( < \tau > \) is the average fluorescence decay lifetime, \( f_i \) is a sufficient contribution of each component to total fluorescence intensity and the values of \( f_1 + f_2 + f_3 = 1 \), \( B \) is the relative contributions.

The fluorescence decay of the RAs–HAS system were shown in Figure 5A-5D and Table 3. Experiment data present the free BHb with decay lifetime is 4.76 ns. With addition of RED, ACE, REA and DEO, the average life time changes to 4.85, 4.94, 4.96 and 4.78 ns. The changes display the interaction of HAS with RED, ACE, REA and DEO. The electron transfer between Trp and RED, the same for ACE, REA and DEO, may accompany the lifetime change in the Trp fluorescence decay. It is fairly well possible that the interaction process alters the protein conformation assisting the quenching process, leading to protein denaturation. It is important to note that the observed triple exponential decay reflects the existence of several Trp in the protein, with different decay times, resulting from different environments [33-35].

**Circular Dichroism (CD) studies**

CD spectroscopy is an optical technique to provide information of the secondary and tertiary structures of protein. Figure 6 shows the CD spectra of RAs–HAS. There are two negative bands at 209 and 222 nm and a positive band at 215 nm of HAS. The 209 nm band corresponds to \( \pi-\pi^* \) transition and \( \alpha\)-helix, the 222 nm band due to \( \pi-\pi^* \) transition for both the \( \alpha\)-helix and random coil, and the 215 nm band corresponds to \( \beta\)-sheet.

Table 4 reveals effects of RAs on the percentage of secondary elements using CDNN program. The degrees of increasing in the percentage of \( \alpha\)-helix (RED 26.00%, ACE 25.00%, REA 25.68% and DEO 24.11%, respectively) are observed, the degrees of reduction in the percentage of \( \beta\)-sheet (RED 22.28%, ACE 23.33%, REA 22.96% and DEO 24.38%, respectively) are observed. The results indicate that the binding of RED, ACE, REA and DEO induce some secondary structure changes in HAS and increase of \( \alpha\)-helix content but loss of \( \beta\)-sheet content protein. The decrease in \( \beta\)-sheet content suggests that RED, ACE and DEO are likely to be binding to the amino acid residues of the main polypeptide chain and further cause partial unfolding of HAS.

**Conclusions**

The interactions between RAs and HAS have been investigated by fluorescence and CD spectroscopy. The linearity of modified Stern–Volmer plots indicated that RAs could bind to one class of sites on HAS.
Figure 4: (A) synchronous fluorescence spectra of HSA and RED, [RED]=0, 0.80, 1.60, 2.40, 3.20, 4.00 μM; (B) synchronous fluorescence spectra of HSA and ACE, [ACE]=0, 0.80, 1.60, 2.40, 3.20, 4.00 μM; (C) synchronous fluorescence spectra of HSA and REA, [REA]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM; (D) synchronous fluorescence spectra of HSA and REO, [REO]=0, 0.50, 1.00, 1.50, 2.00, 2.50 μM; [HSA]=5 × 10⁻⁶ μM, T=298 K.

Figure 5: Fluorescence decay profiles of RAs–HSA system in aqueous solution. (A) RED–HSA; (B) ACE–HSA; (C) REA–HSA; (C) REO–HSA.
The thermodynamic parameters of the binding reactions indicated the presences of hydrophobic forces and hydrogen interactions in the RAs–HSA interactions. Additionally, the results showed that the binding of the three RAs to HSA induced small conformational changes in the overall structure of HSA, which was further proved by the quantitative analysis data of the CD spectra. This will provide important insight into the interactions of the physiologically important protein HSA with globally pervasive contaminants RAs.

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