Differences in Esterase Activity to Aspirin and p-Nitrophenyl Acetate among Human Serum Albumin Preparations

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Human serum albumin (HSA) has two major ligand-binding sites, sites I and II, and also hydrolyzes some compounds at both sites. In the present study, we investigated differences in esterase activity among HSA preparations, and also the effects of warfarin, indomethacin, and naproxen on the hydrolytic activities of HSA to aspirin and p-nitrophenyl acetate. The esterase activities of HSA to aspirin or p-nitrophenyl acetate were measured from the pseudo-first-order formation rate constant (kobs) of salicylic acid or p-nitrophenol by HSA. Inter-lot variations were observed in the esterase activities of HSA to aspirin and p-nitrophenyl acetate; however, the esterase activity of HSA to aspirin did not correlate with that to p-nitrophenyl acetate. The inhibitory effects of warfarin and indomethacin on the esterase activity of HSA to aspirin were stronger than that of naproxen. In contrast, the inhibitory effect of naproxen on the esterase activity of HSA to p-nitrophenyl acetate was stronger than those of warfarin and indomethacin. These results suggest that the administration of different commercial HSA preparations and the co-administration with site I or II high-affinity binding drugs may change the pharmacokinetic profiles of drugs that are hydrolyzed by HSA.

Key words human serum albumin (HSA); esterase activity; aspirin; p-nitrophenyl acetate; manufacturing lot; drug interaction

The introduction of an ester linkage generally improves the bioavailability of therapeutic agents by increasing passive transport following oral administration. A wide variety of ester-containing drugs and prodrugs are used in clinical practice. Ester-containing prodrugs, such as olmesartan medoxomil, are hydrolyzed, and, thus, activated by HSA. Since polypharmacy is common among the elderly, the drug interactions on the esterase activity of HSA are considered to be important. Differences have been reported in drug-binding data among pharmaceutical-grade albumin and manufacturing lots of bovine serum albumin. Furthermore, lot-to-lot variability in high-molecular-weight aggregates and the glycation of recombinant HSA expressed in Oryza sativa as well as differences in esterase-like activities of HSA among manufacturing lots and suppliers due to cholinesterase contamination have also been reported. However, the effects of drug interactions on hydrolysis by HSA among manufacturing lots have not yet been investigated in sufficient detail.

In the present study, we found differences in esterase activities among the manufacturing lots of HSA tested. We used reactions in which aspirin, as a model compound for site I, and p-nitrophenyl acetate, as a model compound for site II, are hydrolyzed to salicylic acid and p-nitrophenol by HSA, respectively (Fig. 1). We also investigated the effects of indomethacin and naproxen, which are frequently prescribed nonsteroidal anti-inflammatory drugs, and warfarin, which is an anticoagulant agent that has been co-administered with aspirin, on the esterase activity of HSA.

MATERIALS AND METHODS

Materials HSA (A3782) and eserine were purchased from Sigma-Aldrich (MO, U.S.A.). Four different lots (113K7601, 085K7541, 090M7001V, SLBD7204V) of HSA (A3782) were tested. Aspirin, sodium salicylate, p-nitrophenyl acetate, p-nitrophenol, warfarin sodium, indomethacin, naproxen, o-tolucic acid, 5,5′-dithiobis(2-nitrobenzoic acid), sodium dodecyl sulfate (SDS), ethylendiaminetetraacetic acid (EDTA) (pH 8.0) and NEFA C-Test Wako were purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). Indoxyl sulfate and 3-indoleacetic acid were purchased from Nacalai Tesque, Inc. (Kyoto, Japan). All other chemicals and solvents were of analytical grade or higher.

Esterase Activity Measurement Esterase activity to as-
pirin was measured by modifying the method of Ma et al. \(^{13}\) HSA (final concentration: 200 µM) in 67 mM potassium phosphate buffer (pH 7.4) was pre-incubated at 37°C for 5 min, and aspirin (100 µM) was then added to the pre-incubated mixture at 37°C. The total incubation volume was 0.13 mL. The incubation, which proceeded for 1, 2, 3, or 4 h, was terminated by the addition of 0.5 mL of ice-cold 0.1 M sodium phosphate buffer (pH 2.0)/acetonitrile (20/80) containing 12.5 µM α-toluidic acid as an internal standard to the incubation mixture. The mixture was centrifuged at 18,620× g at 4°C for 5 min. After centrifugation, the supernatant was used in the analysis of salicylic acid (aspirin metabolite) by a HPLC system. These procedures were performed in duplicate. The pseudo-first-order formation rate constant (\(k_{\text{obs}}\)) of salicylic acid was obtained by linear regression from the semi-logarithmic plot of the decay of the concentration of aspirin subtracted from that in the buffer control versus time. Kinetic analysis processed as above except for the final aspirin concentration of 50–400 µM and the incubation time of 2 h.

Esterase activity to p-nitrophenyl acetate was measured by modifying the method of Ikeda et al. \(^{19}\) HSA (final concentration: 50 µM) in 67 mM potassium phosphate buffer (pH 7.4) was pre-incubated at 25°C for 5 min, and p-nitrophenyl acetate (10 µM) was then added to the pre-incubated mixture at 25°C. The reaction of p-nitrophenyl acetate with HSA was followed by spectrophotometrically monitoring the appearance of nitrophenol at 400 nm. \(k_{\text{obs}}\) was obtained by linear regression from the semi-logarithmic plot of the decay of the concentration of p-nitrophenyl acetate versus time. Since the \(k_{\text{obs}}\) of p-nitrophenol is significantly larger than that of the buffer control, that of the buffer control was ignored. Kinetic analysis processed as above except for the final HSA concentration of 30 µM, the final p-nitrophenyl acetate concentration of 12.5–100 µM, total incubation volume of 0.5 mL, incubation time of 60 s, and being terminated by the addition of chilled ethanol (0.5 mL).

**Inhibition Study** The measurement of esterase activity to aspirin was repeated in the presence of the following inhibitors: warfarin, indomethacin, and naproxen at a final concentration of 200 µM. HSA and its inhibitors were dissolved in 67 mM potassium phosphate buffer (pH 7.4). These samples were pre-incubated at 37°C for 5 min before the addition of the substrate, and were then processed as above, with 625 µM α-toluidic acid being used as an internal standard in the inhibition study on indomethacin.

The measurement of esterase activity to p-nitrophenyl acetate was repeated in the presence of the following inhibitors: warfarin, indomethacin, and naproxen at a final concentration of 50 µM, and eserine at a final concentration of 20 µM. HSA and its inhibitors were dissolved in 67 mM potassium phosphate buffer (pH 7.4). These samples were pre-incubated at 25°C for 5 min before the addition of the substrate, and were then processed as above.

**HPLC Analysis** HPLC analysis of salicylic acid was performed according to the method of Fogel et al. \(^{20}\) with some modifications. The HPLC system consisted of a Shimadzu SCL-10A VP system controller, Shimadzu LC-10ADVP pump, Shimadzu SIL-10ADVP auto injector, Shimadzu SPD-10AVP variable wavelength UV detector, and Shimadzu RF-10AXL fluorescence detector (Kyoto, Japan). A column of Wakopak Wakosil 5C18 AR (4.6 mm i.d. × 250 mm; Wako Pure Chemical Industries, Ltd.) was used as the stationary phase. The column temperature was maintained at 40°C. The injected volume was 25 µL. The mobile phase consisted of 0.1 M sodium dihydrogen phosphate pH 2.0/acetonitrile (69.5/30.5) in the inhibitory study on warfarin, but consisted of 0.1 M sodium dihydrogen phosphate pH 2.0/acetonitrile (69/31) in other studies. The flow rate was 0.5 mL/min. The effluent was monitored at 295 nm with a UV detector in the inhibitory study on warfarin and naproxen, and at an excitation wavelength of 296 nm and emission wavelength of 405 nm with a fluorescence detector in the inhibitory study on indomethacin.

**Determination of Indoxyl Sulfate and 3-Indoleacetic Acid Concentrations** Four percent HSA solutions were mixed with twice volumes of acetonitrile. The mixture was centrifuged at 18,620× g at 4°C for 5 min. After centrifugation, the supernatant was used in the analysis of indoxyl sulfate and 3-indoleacetic acid by the above-mentioned HPLC system. HPLC analysis of indoxyl sulfate was performed according to the method of de Loor et al. \(^{22}\) and Banoglu et al. \(^{23}\) with some modifications. The mobile phase consisted of 25 mM sodium dihydrogen phosphate pH 2.0/acetonitrile (90/10). The flow rate was 0.7 mL/min. The effluent was monitored at 280 nm with a UV detector, and at an excitation wavelength of 280 nm and emission wavelength of 390 nm with a fluorescence detector. HPLC analysis of 3-indoleacetic acid was performed according to the method of Martinez et al. \(^{24}\) and Kelen et al. \(^{25}\) with some modifications. The mobile phase consisted of 0.1 M sodium dihydrogen phosphate pH 2.0/acetonitrile (69/31). The flow rate was 0.7 mL/min. The effluent was monitored at 280 nm with a UV detector, and at an excitation wavelength of 280 nm and emission wavelength of 340 nm with a fluorescence detector.

**Determination of Non-esterified Fatty Acid (NEFA) Concentrations** The NEFA concentrations contained in 4% HSA solutions were determined using the NEFA C-Test Wako from two different lots.

**Determination of Sulfhydryl Concentrations** The sulfhydryl concentrations in HSA solutions were determined according to the method of Takabayashi et al. \(^{26}\) DTNB (5,5’-dithiobis(2-nitrobenzoic acid), 0.1 mL of 4 mg/mL solu-
tion in 50 mM phosphate buffer containing 2 mM EDTA (pH 8.0) was mixed with 0.4 mL of 2.5% SDS and 0.5 mL of 5 mg/mL HSA in 50 mM phosphate buffer (pH 8.0). The absorbance was measured against reagent blank at 412 nm after 30 min at room temperature. The sulphydryl concentration was calculated using the molar absorption coefficient of 13600 M$^{-1}$ cm$^{-1}$.

**Determination of HSA Concentrations** The concentrations of HSA were determined spectrophotometrically at 279 nm ($\varepsilon = 36000$ M$^{-1}$ cm$^{-1}$).27

**Determination of Glycated Albumin in HSA** Glycated albumin values were measured using an enzymatic method in the laboratory at SRH, Inc. (Tokyo, Japan).

**Statistical Analysis** Data are expressed as mean values and the standard deviation. The Student's $t$-test was used to compare data obtained in 2 groups. An ANOVA was performed to test the significance of differences among groups. Significance regarding differences in the means among groups was determined by Tukey's test or Dunnett's pairwise $t$-test. Pearson's correlation coefficient test was used to test the relationship between two groups. Significance was defined as $p$<0.05.

**RESULTS**

Differences in esterase activity to aspirin among 4 different lots of HSA preparations, and effects of warfarin, indomethacin and naproxen on the esterase activities of those HSA preparations are summarized in Fig. 2. Inter-lot variations of esterase activity to aspirin were observed in the control group, and the esterase activities of 090M7001V and SLBD7204V of HSA preparations were higher than those of 113K7601 and 085K7541. The esterase activities of 090M7001V and SLBD7204V compared with 113K7601 were significantly decreased by the addition of warfarin and indomethacin. The inhibition of esterase activity to aspirin by warfarin and indomethacin was stronger than that by naproxen.

Differences in esterase activity to $p$-nitrophenyl acetate among the 4 lots of HSA preparations, and effects of warfarin, indomethacin and naproxen on the esterase activities of those HSA preparations are summarized in Fig. 3. Marked inter-lot variations were observed in the control group. Particularly, the esterase activity of 113K7601 was low. The inhibitory effect of naproxen on the esterase activity of HSA to $p$-nitrophenyl acetate was much stronger than those of warfarin and indomethacin. Although inter-lot variations in each inhibitor group were similar to those in the control, naproxen strongly inhibited esterase activity of 085K7541, 090M7001V and SLBD7204V compared with 113K7601.

Kinetic parameters for hydrolysis of aspirin by HSA are summarized in Table 1. Since esterase activity of HSA to aspirin was examined between two groups, the study of kinetic parameters for hydrolysis of aspirin was performed with lots of 085K7541 and 090M7001V, which were low glycated albumin value. As for kinetic parameters for hydrolysis of aspirin, there were no significant differences between 085K7541 and 090M7001V.

Kinetic parameters for hydrolysis of $p$-nitrophenyl acetate by HSA are summarized in Table 2. The $V_{\text{max}}$ value of 113K7601 was slightly higher than that of other lots, and the $K_m$ value of 113K7601 was much higher than that of other lots. The $V_{\text{max}}/K_m$ (apparent intrinsic clearance) value of 113K7601 was smaller than those of other lots.

None of the esterase activities of any of the HSA tested to $p$-nitrophenyl acetate were inhibited by 20 $\mu$g eserine, which is a well-known cholinesterase inhibitor (Table 3).

The glycitated percentages of HSA, NEFA contents and sulphydryl contents contained in the HSA solutions from different manufacturing lots of HSA preparations are summarized in Table 4. Glycated albumin values contained in 113K7601 and SLBD7204V of HSA preparations were higher than those of 085K7541 and 090M7001V. Although less than 0.01 mol NEFA/mol HSA contents of 113K7601 and 085K7541 of HSA preparations were detected, about 0.1–0.2 mol NEFA/mol HSA contents of 090M7001V and SLBD7204V were observed. Sulphydryl contents in the HSA preparations of 090M7001V and SLBD7204V were also higher than in those of 113K7601 and 085K7541, as similar to NEFA contents. Contents of indoxyl sulfate and 3-indoleacetic acid in all lots of HSA preparations were less than 0.01 mol/mol HSA (data not shown).

**DISCUSSION**

Esterase activity in different manufacturing lots of HSA preparations to aspirin and $p$-nitrophenol acetate varied widely. However, esterase activity to aspirin in each manufacturing lot did not correlate with that to $p$-nitrophenol acetate. These results suggest the site specificity of esterase activity.

The inhibition of esterase activity to aspirin by warfarin and indomethacin was stronger than that by naproxen, whereas the inhibition of esterase activity to $p$-nitrophenyl acetate by warfarin and indomethacin was weaker than that by naproxen. We confirmed that substrates binding to site I (i.e., warfarin and indomethacin) and site II (i.e., naproxen) of HSA inhibited hydrolysis in sites I and II, respectively.

Since differences were observed in esterase activity among manufacturing lots, further investigations on the HSA preparations used in the present study were performed. The effects of cholinesterase contamination on the esterase activity of HSA from different manufacturing lots were determined because Chapuis et al.19 previously reported that the esterase activity of serum albumin may be due to cholinesterase contamination in some commercially available HSA preparations. They also found significant differences in esterase activity toward moxisylyte among the commercially available types and lots of HSA; however, the A3782 type of HSA, which was used in the present study, displayed no esterase activity toward moxisylyte. We examined the esterase activity of HSA solutions to $p$-nitrophenyl acetate with or without eserine. None of the HSA solutions tested exhibited the inhibitory effects of eserine on the esterase activity of the HSA preparations, which is consistent with previous findings.

Since indoxyl sulfate and 3-indoleacetic acid, which are endogenous substances binding to site II of HSA,28 were not detected in HPLC analysis and a tested HSA preparation was of high purity ($\geq 99\%$), it is hardly likely that only HSA of 113K7601 was contaminated with some substance binding to HSA. The $K_m$ value of 113K6701 for hydrolysis of $p$-nitrophenol acetate by HSA was higher than those of other lots. However, esterase activity of 113K6701 to $p$-nitrophenol acetate was inhibited by naproxen, warfarin and indomethacin in common with those of other lots. These findings may suggest structural changes around the region of esterase activity.
Sulfhydryl contents were measured for the purpose of detecting post-translational modification on 34Cys of HSA by oxidative stress concerned with human mercaptalbumin and human nonmercaptalbumin. As with the report of Minami et al., sulfhydryl contents of 090M7001V and SLBD7204V were higher than those of 113K7601 and 085K7541. Coupling between fatty acid binding and sulfhydryl oxidation was also observed as with bovine serum albumin, although the hydrolisis for p-nitrophenyl acetate was not decreased unlike the report of Anraku et al.

Fatty acid binding to HSA induces conformational changes, and the presence of up to 3 mol fatty acids/mol HSA or 8 mol fatty acids/mol HSA has been shown to enhance the binding of warfarin or aspirin to HSA in subdomain IIA, respectively. The binding of 5-dimethylamino-naphthalene-1-sulfonamide (DNSA) to site I was also increased in a linear fashion by addition of up to 2 mol fatty acid/mol HSA to commercial albumin preparations. Since low levels of fatty acids were detected in the HSA of 090M7001V and SLBD7204V, we attributed the strong esterase activity of the HSA of 090M7001V and SLBD7204V to aspirin to enhanced aspirin binding to HSA. The esterase activities of the HSA of 090M7001V and SLBD7204V to aspirin were markedly inhibited by warfarin and indomethacin. The affinity or percentage protein binding for aspirin to HSA is known to be lower than that of salicylic acid, and the affinity of salicylic acid to HSA is lower than that of warfarin and indomethacin. Hence, the inhibition by warfarin and indomethacin following enhanced warfarin and indomethacin binding to HSA over aspirin binding may decrease esterase activity toward aspirin. No relationship was observed between the glycation of albumin and esterase activity to aspirin and p-nitrophenyl acetate. Conflicting findings have been reported on the affinity of glycated albumin to site I or II high-affinity binding drugs. Methylglyoxal-derived hydroimidazolone is also a major glycation adduct of albumin and esterase activity of...
HSA modified by methylglyoxal to \( p \)-nitrophenyl acetate decreased.\(^{33}\) Since the extent of modification, however, is small, the effect of methylglyoxal-derived hydroimidazolone can be hardly considered the cause. Differences between the post-translational modifications of HSA prepared experimentally and structural changes containing the post-translational modifications formed in raw material and preparation procedures of HSA preparation may cause different effects on physiological function of HSA between these some previous studies and the present study. Further experiments are required to define the relationship between physiological function and heterogeneity of HSA preparation.

The present study demonstrated differences in esterase activities in sites I and II among the manufacturing lots of HSA preparations. Our results also suggest that drugs that bind to site I or II with high affinity inhibit the hydrolysis of drugs by HSA, with the extent of inhibition differing among the manufacturing lots of HSA. Differences in the manufacturing lots of commercial HSA preparations may affect the pharmacokinetics of drugs that are hydrolyzed by HSA, and hydrolytic activity may be altered by the presence of high-affinity binding drugs to HSA. Further studies are needed in order to examine the relationship between esterase activity and the conformation of HSA and also determine the effects of drug interactions between various drugs on the esterase activity of HSA.

**Conflict of Interest** The authors declare no conflict of interest.

**REFERENCES**


