

occur, resulting in modification of its biological properties.

In recent years, activated phagocytes have been reported to be a major source of reactive oxidants and to play a fundamental role in host defense [6,7]. They contain the heme enzyme myeloperoxidase (MPO) which catalyzes the reaction of chloride ion with hydrogen peroxide ( $H_2O_2$ ) to generate large amounts of hypochlorous acid (HOCl), a powerful oxidizing and chlorinating agent produced by neutrophils [8–10]. Treatment of HSA with HOCl *in vitro* and *in vivo*-generated AOPP have been reported to trigger oxidative bursts in neutrophils as well as in monocytes, thereby appearing to act as true inflammatory mediators [11,12]. The possibility that excessively oxidized albumin in plasma increases the production of ROS by stimulating neutrophils, cannot be excluded. Therefore, very excessively oxidized HSA, could also play an important role as a pro-oxidant in dialysis patients.

We previously demonstrated that oxidative stress in HD patients is manifested by an increase in plasma protein oxidation that is characterized by thiol group oxidation and the formation of carbonyl groups on proteins. We also showed that HSA is the major plasma protein target of oxidative stress in uremia, and that increased levels of carbonyl compounds are correlated with the oxidation of albumin in uremic patients [13]. Therefore, serum albumin is an important protein that has direct protective effects. These effects may be also based on a variety of biological mechanisms. In the present study, we further investigated the role of albumin oxidation and neutrophil oxidative burst in HD patients and compared the results with age- and gender-matched control subjects. An evaluation of the effects of oxidative stress on the structural and functional properties of HSA was also attempted.

## Materials and methods

### Patients

The protocol used in this study was approved by the institutional review board and informed consent was obtained from all subjects. A total of 30 subjects were enrolled: 20 stable HD patients (10 men, 10 women) aged 36–87 years, with a duration of dialysis ranging from 1 to 9 years, and 10 age- and gender-matched healthy control subjects. End-stage renal failure in HD patients was caused by glomerulonephritis ( $n = 5$ ), nephrosclerosis ( $n = 2$ ) or diabetic nephropathy ( $n = 13$ ). At the time of enrollment, all HD patients had been receiving regular bicarbonate hemodialysis therapy (4–5 h/session, three times per week) using high-flux polysulfone hollow-fiber dialyzers. The profiles of healthy controls and HD patients with or without diabetes are summarized in Table 1.

### Purified albumin from healthy control and HD patients

HSA samples were isolated by polyethylene glycol fractionation of blood plasma followed by chromatography on a blue Sepharose CL-6B column (Amersham-Pharmacia, Uppsala, Sweden) [14]. The

Table 1  
Characteristics of the normal and patient groups

	Normal subjects ( $N = 10$ )	HD patients ( $N = 20$ )
Age (years)	$67.8 \pm 1.8$	$62.8 \pm 12.7$
Gender (M/F)	6/4	10/10
Diabetes/nondiabetes	—	13/7
Creatinine (mg/dL)	$0.85 \pm 0.2$	$10.6 \pm 2.6^*$

Values are expressed means  $\pm$  SD.

\*  $P < 0.01$ .

samples were then dialyzed against deionized water for 48 h at 4 °C, followed by lyophilization. The purity of the HSA samples was at least 95%, and the percentage of dimers did not exceed 7%, as evidenced by SDS-PAGE and native-PAGE, respectively. The long-chain fatty acid contents of isolated HSA samples were determined using the copper triethanolamine method [15]. There was no significant change of the long-chain fatty acid contents in purified albumin from healthy controls and HD patients.

### Determination of serum protein and purified albumin oxidation

**Chromatographic analysis of albumin in normal subjects and HD patients.** HSA is a mixture of mercaptalbumin (HMA; reduced form) and nonmercaptalbumin (HNA; oxidized form). HMA contains one highly reactive sulfhydryl group at position 34 (Cys-34), while other serum proteins contain little or none. HNA is comprised of at least three types of molecules. The major HNA component is a mixed disulfide with cysteine or glutathione (HNA-1). The other is a more highly oxidized product than the mixed disulfide, in which the thiol group has been oxidized to the sulfenic (SOH), sulfinic ( $SO_2H$ ), and sulfonic ( $SO_3H$ ) states (HNA-2), the proportions of which are extremely small in extracellular fluids [16,17]. The high-performance liquid chromatography (HPLC) analysis of albumin developed by Sogami et al. permits the clean separation of HSA into HMA and HNA, and is used for the determination of the redox state for various pathophysiologic conditions. An HPLC analysis of serum albumin was performed as described in a previous report [13]. Serum samples were immediately frozen immediately after they were drawn, and were stored at  $-80$  °C until used for HPLC. HPLC was performed using 5  $\mu$ L aliquots of each serum sample and a Shodex Asahipak ES-502N column (Showa Denko, Tokyo, Japan; column temperature:  $35 \pm 0.5$  °C). The HPLC system consisted of an L-6200 intelligent pump equipped with a gradient programmer and an F-1050 fluorescence detector (Jasco, Tokyo, Japan). Elution was performed using a linear gradient with ethanol concentrations increasing from 0% to 5% with the serum dissolved in a mixture of 0.05 mol/L sodium acetate and 0.40 mol/L sodium sulfate (pH 4.85) at a flow rate of 1.0 mL/min. From the HPLC profiles of HSA, the value of each albumin fraction [ $f(HMA)$ ,  $f(HNA-1)$ , and  $f(HNA-2)$ ] was estimated by dividing the area of each fraction by the total area corresponding to HSA.

**Total plasma protein and individual plasma carbonyl contents measurement.** Plasma protein carbonyl content was determined using the method of Climent et al. [18].

### Biological properties of Normal- and HD-HSA

**Radical scavenging ability of Normal- and HD-HSA.** The radical scavenging activity of Normal- and HD-HSA (10  $\mu$ M) was determined from the decrease in the absorbance of 1,1'-diphenyl-2-picrylhydrazyl (DPPH) radicals at 517 nm due to their scavenging of an unpaired electron of the stable DPPH radical in a mixture of 10 mL ethanol, 10 mL of 50 mM 2-(*N*-morpholino)ethanesulfonic acid (Mes) buffer (pH 5.5), and 5 mL of 0.5 mM DPPH in ethanol [19–21].

**Measurement of neutrophil respiratory burst.** Neutrophils were isolated from heparinized peripheral blood of healthy donors using Lymphoprep (Nycomed, Oslo, Norway) density gradient centrifugation. The purity of the PMN preparations routinely exceeded 95%, and cell viability, as determined by propidium iodide staining, was at least 98%. Accumulation of dihydrorhodamine 123 (DRD) in a suspension of neutrophils was measured using a flow cytometer, monitoring the fluorescence at 526 nm [22]. Suspensions of neutrophils ( $1 \times 10^6$  cells) were incubated with 5  $\mu$ M DRD for 15 min at 37 °C in serum-free medium. After DRD incubation, the neutrophil suspension was centrifuged and washed to remove unincorporated probe. The cells were then treated with various concentrations of albumin medium for 1 h at 37 °C, and were then placed on ice. The mean fluorescence intensity of rhodamine (RD) in the cells was determined using a flow cytometer (FACS Calibur; Becton–Dickinson Biosciences, Franklin Lakes, NJ).

#### Structural and functional properties of Normal- and HD-HSA

**CD measurements.** These measurements were performed using a Jasco J-720 type spectropolarimeter (Jasco, Tokyo, Japan) at 25 °C. Far-UV spectra were recorded at a protein concentration of 20  $\mu$ M in 67 mM sodium phosphate buffer (pH 7.4) using 1 mm quartz cells.

**Effective protein hydrophobicity.** The effective hydrophobicity of all the albumins (1  $\mu$ M), dissolved in a 67 mM sodium phosphate buffer (pH 7.4), was probed with 4,4'-dianilino-1,1'-binaphthyl-5,5'-disulfonic acid (bis-ANS) (10  $\mu$ M) at 25 °C. The compound was excited at 394 nm and fluorescence spectra were recorded on a Jasco FP-770 fluorescence spectrometer (Tokyo, Japan) using 1 cm quartz cells.

**Changes in protein net charge.** Changes in the net charge of albumin were evaluated by a modification of the capillary electrophoresis method described by Pande et al. [23]. One milliliter of an HSA sample (2  $\mu$ M) was run in 100 mM borate buffer (pH 8.5 and 20 °C), and the migration time was determined by capillary electrophoresis, model CE990/990-10 from Jasco (Tokyo, Japan).

**Binding properties.** The binding of warfarin (5  $\mu$ M) and ketoprofen (5  $\mu$ M) to purified HSA (10  $\mu$ M) in 67 mM sodium phosphate buffer (pH 7.4 and 25 °C) was examined by ultrafiltration. The unbound ligand fractions were separated using an Amicon MPS-1 micropartition system with YMT ultrafiltration membranes by centrifugation (2000g, 40 min). The concentration of unbound ligand was determined by HPLC [14]. The unbound fraction (%) was calculated as follows:

$$\text{Unbound fraction(\%)} = \frac{\text{ligand concentration in filtered fraction/total ligand concentration(before ultrafiltration)}}{\times 100.}$$

#### Statistics

The statistical significance of collected data was evaluated using the ANOVA analysis followed by Newman–Keuls method for more than 2

means. A value of  $P < 0.05$  was considered to indicate statistical significance. The results are reported as the mean  $\pm$  SD.

## Results

### Determination of serum protein and purified albumin oxidation

The oxidation of a protein typically results in an increase in carbonyl content. This increase is due to the oxidation of Lys, Arg, or Pro residues. Plasma protein carbonyl contents were significantly increased in HD patients, compared with normal subjects. Data for the carbonyl content were as follows: HD patients groups,  $3.12 \pm 1.11$  nmol/mg protein,  $n = 20$ ; controls groups,  $2.1 \pm 0.34$  nmol/mg protein,  $n = 10$  ( $P < 0.01$ ) (Table 2). In a previous study, we determined the redox states of HD patients during oxidative stress, with emphasis on the oxidation of serum albumin [13]. In the present study, using HPLC, we determined the oxidation status of the Cys-34 residues in albumin. Table 2 shows typical HPLC profiles for serum albumin from healthy subjects and HD patients. f(HMA) was substantially decreased and both f(HNA-1) and f(HNA-2) were significantly increased in HD patients compared with healthy subjects ( $P < 0.01$ , Table 2). These results suggest that uremia results in an increase in the oxidized serum albumin levels, via oxidative stress.

Further, to investigate the extent of alterations in the biological properties of HSA in HD patients, HSA of HD patients (HD-HSA), and normal subjects (Normal-HSA) were purified on a blue Sepharose CL-6B column. The albumin carbonyl contents of the purified fraction were significantly increased in HD patients. Data for the carbonyl content of purified HSA were as follows: HD-HSA,  $2.86 \pm 0.45$  nmol/mg protein,  $n = 20$ ; Normal-HSA,  $2.13 \pm 0.14$  nmol/mg protein,  $n = 10$ ,  $P < 0.01$  (Table 2). HD-HSA samples also showed a markedly increased HNA ratio, compared with Normal-HSA ( $P < 0.01$ , Table 2). In addition, the HNA/HMA ratio of purified HSA was closely correlated with the HNA/HMA of plasma ( $R = 0.952$ ,  $P < 0.01$ ;

Table 2  
Determination of serum protein and purified albumin oxidation

	Serum protein		Purified HSA	
	Normal subjects	HD patients	Normal-HSA	HD-HSA
Carbonyl contents (nmol/mg protein)	$2.10 \pm 0.34$	$3.12 \pm 1.11^*$	$2.13 \pm 0.14$	$2.86 \pm 0.45^{**}$
f(HMA) (%)	$53.6 \pm 6.4$	$40.4 \pm 8.7^*$	$46.6 \pm 4.9$	$32.7 \pm 6.8^{**}$
f(HNA-1) (%)	$38.7 \pm 6.3$	$49.7 \pm 8.0^*$	$42.0 \pm 4.4$	$54.4 \pm 7.1^{**}$
f(HNA-2) (%)	$7.7 \pm 0.9$	$9.9 \pm 1.4^*$	$11.4 \pm 0.3$	$12.9 \pm 0.7^{**}$

Values are expressed means  $\pm$  SD.

\*  $P < 0.01$  as compared with Normal subjects.

\*\*  $P < 0.01$  as compared with Normal-HSA.

data not shown). These results suggest that the state of purified HSA accurately reflects the redox state of albumin in the blood.

#### Physiological properties of Normal- and HD-HSA

##### Radical scavenging ability of Normal- and HD-HSA

The effects of Normal- and HD-HSA were examined by monitoring the time-dependent change in the absorbance of DPPH radicals. As shown in Fig. 1, Normal-HSA scavenged DPPH radicals, and its ability to scavenge proceeded more rapidly than HD-HSA. HD-HSA scavenged about 15% of DPPH radicals even after 20 min. In contrast, the effect of Normal-HSA proceeded more rapidly, about 20% of the DPPH radicals were scavenged within 20 min. This result shows that the radical scavenging activity of Normal-HSA was significantly greater than that of HD-HSA.

##### Neutrophil respiratory burst

It was recently reported that in vitro-oxidized albumin up-regulates ROS generation in neutrophil suspensions [4]. In order to directly assess whether purified HSA had the ability to induce oxidative stress in a neutrophil suspension, we used the DRD method and a FACS analysis. The levels of ROS were higher for HD-HSA than for Normal-HSA (Fig. 2) ( $P < 0.05$ ). These results show that HD-HSA was not able to decrease the antioxidant activity, but also to trigger the oxidative burst of human neutrophils.

#### Structural and functional properties of Normal- and HD-HSA

##### Structural properties of Normal- and HD-HSA

The structural properties of Normal- and HD-HSA were examined by a variety of methods. Thus, Fig. 3A

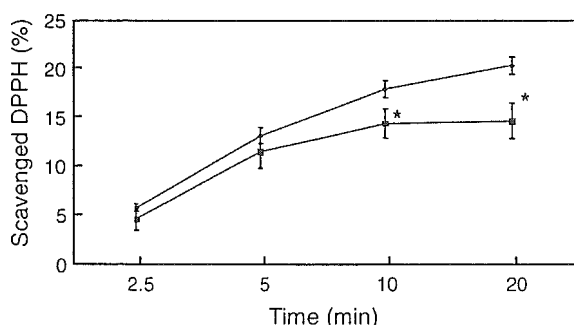


Fig. 1. Radical scavenging ability of Normal- and HD-HSA. Normal- and HD-HSA at a final concentration of 20  $\mu$ M were added to a suspension of ethanol, in which DPPH radicals were incorporated (final concentration of 20  $\mu$ M), in 175 mM KCl and 10 mM Mes buffer (pH 7.4) at 25  $^{\circ}$ C. Changes in the amount of DPPH radicals in the suspension were monitored as a decrease in absorbance at 517 nm. Changes in DPPH radicals scavenged relative to total DPPH radicals with time are shown. \* $P < 0.05$ , compared with Normal-HSA.

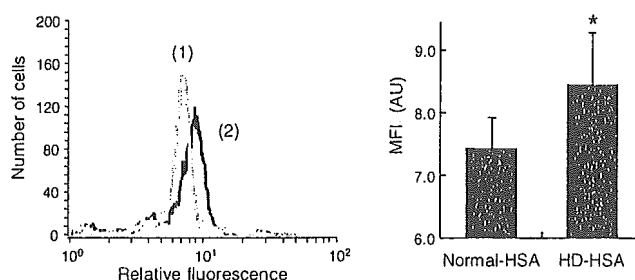


Fig. 2. ROS production by neutrophils incubated with purified HSA. (1) Normal-HSA, (2) HD-HSA. \* $P < 0.05$ , compared with Normal-HSA.

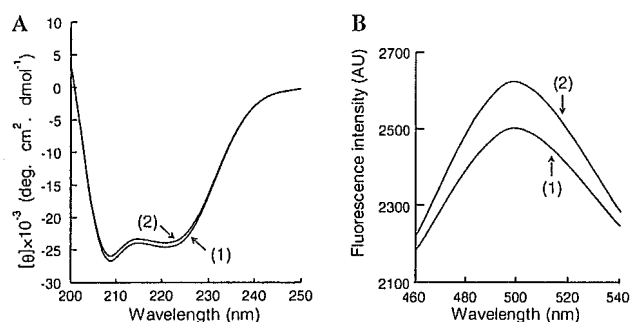


Fig. 3. (A) Purified HSA far-UV CD spectra of normal subjects and HD patients. (1) Normal-HSA, (2) HD-HSA. The spectra are the average of three determinations. (B) Effect of uremia on the fluorescence of purified HSA-bound bis-ANS. (1) Normal-HSA, (2) HD-HSA. The spectra are the average of three determinations.

shows Far-UV CD spectra. As can be seen in Fig. 3A, the characteristics of the CD spectrum of HD-HSA were slightly different from that obtained for Normal-HSA. The effect of oxidation on the exposure of hydrophobic areas was also examined by using the fluorescence probe bis-ANS. The results obtained (Fig. 3B) indicate that the conformation of HD-HSA in particular, involves an increase in accessible hydrophobic regions, compared with Normal-HSA. The net charges of HD- and Normal-HSA were investigated by determining their migration times in capillary electrophoresis. The migration time for HD-HSA ( $25.65 \pm 0.52$  min) was increased slightly compared with that of Normal-HSA ( $24.84 \pm 0.46$  min) ( $P < 0.01$ ). These results suggest that HD-HSA also underwent conformational changes, hydrophobic regions were more exposed and negative charge on the molecule was increased.

##### Binding property of Normal- and HD-HSA

The unique ligand binding properties of HSA can, to a great extent, be explained by the presence of two major binding sites, Site I and Site II [24], which are located within specialized cavities in subdomains IIA and IIIA, respectively [25]. The potential effect of oxidation on these sites was examined by using warfarin and ketopro-

fen as representative ligands. As seen from Table 3, the high-affinity binding of warfarin, which takes place at Site I [26] was little changed, but the high affinity binding of the Site II-ligand ketoprofen [27] was significantly reduced in HD-HSA, compared with Normal-HSA.

## Discussion

Apart from the low-molecular weight antioxidants, the antioxidant activity in human blood plasma relies mainly on proteins such as albumin [28]. In plasma, the free thiol group is quantitatively the most important scavenger of oxidants. Furthermore, the formation of carbonyl groups in plasma albumin, as previously demonstrated by Himmelfarb and Mcmonagle [28], involves in basic amino acid residues as well as Cys-34 with free thiol group, which is also oxidized. In the previous study, using a Western blot immunoassay, we demonstrated that the carbonylation of albumin accounts for nearly all of the excess plasma protein oxidation in HD patients [13]. We also found that the f(HMA) value in albumin from HD patients was less than 80% of the f(HMA) value in albumin from healthy controls, as indicated by HPLC analysis (Table 2). Since albumin is the most abundant plasma protein, it could play a major role as an antioxidant in plasma at least by thiol oxidation and carbonyl formation. In this context, we expected that the characterization of oxidation status of serum albumin would provide, not only useful information regarding the redox state of the human body, but also alterations in the conformation and function of HSA which may result in modifications of its biological properties.

In HD patients with increased oxidative stress, the oxidative modification of plasma proteins especially albumin appear to be more extensive and may be one of the pathological conditions associated with the widespread vascular complications that are frequently observed in HD patients. To investigate the extent of the alterations in the biological properties of HSA in HD patients, HSA from HD patients (HD-HSA) and normal subjects (Normal-HSA) were purified with blue Sepharose CL-6B column chromatography. The retention time and chromatogram of the purified albumin and plasma albumin were found to be highly correlated (Table 2). Thus, the HNA/HMA ratio of purified HSA can be used as an index for the HNA/HMA ratio of plasma ( $R = 0.952$ ,  $P < 0.01$ ). The high correlation between the HNA/HMA value of plasma and purified HSA suggest that the state of purified HSA used throughout the experiments accurately reflects the redox state of albumin in the blood.

We first focused on the antioxidant activity of albumin because oxidative stress is thought to play a significant role in the pathogenesis of many diseases,

including atherosclerosis [29,30]. There is now ample evidence to suggest that albumin, as the main circulating protein, is a quantitatively important antioxidant in the blood and extracellular fluids [31,32]. Our present studies are in full agreement with this view, Normal-HSA offered significant protection against free radicals, compared to HD-HSA (Fig. 1). These antioxidant effects were found to be concentration-dependent (data not shown), consistent with a beneficial effect of high albumin levels in humans. This in turn suggests that excessively oxidized albumin, which acts as a pro-oxidant, may increase cardiovascular complications in HD patients. In these patients recurrent blood interactions with bioincompatible dialysis membranes trigger neutrophil activation and the subsequent generation of highly reactive oxygen species (ROS) including  $O_2^-$  and its derivatives ( $H_2O_2$ ,  $OH^\cdot$ , and  $^1O_2$ ) via the nicotinamide adenine dinucleotide phosphate (NADPH) oxidase complex and OCl via a MPO-dependent reaction between chloride and  $H_2O_2$  [4]. To test the hypothesis that the increased oxidative stress in blood of HD patients is caused by oxidized albumin, we examined the effects of incubating neutrophils with purified albumin from Normal- or HD-HSA. Purified albumin from HD patients (HD-HSA) induced oxidative stress via a neutrophil respiratory burst (Fig. 2). In previous studies, excess AOPP-modified albumin, was found to be associated with a high level of respiratory burst [12], but the samples used in those studies were produced by chemical modification in vitro. In the present studies, HD-HSA, isolated from uremia patients was found to have been modified in vivo. Treatment of neutrophils with purified albumin from HD patients induced oxidative stress. Therefore, HD-HSA was, found not only to have a decreased antioxidant activity, but also able to trigger the oxidative burst of human neutrophils, compared with Normal-HSA.

These results suggest that HSA is continuously exposed to oxidative stress, so much so that alterations in its biological properties that could result in the conformational, functional changes of HSA occur. Therefore, the significance of conformational changes of purified HSA on its functional properties were examined. Slight decreases in  $\alpha$ -helical content accompanied by tertiary conformational changes were observed for HD-HSA (data not shown). These changes, which could

Table 3  
Binding of warfarin and ketoprofen to Normal-HSA and HD-HSA

	Free fraction (%)	
	Warfarin	Ketoprofen
Normal-HSA	19.6 ± 1.9	5.97 ± 0.75
HD-HSA	23.3 ± 2.0	9.37 ± 1.36*

Values are expressed means ± SD.

\*  $P < 0.05$ .

be observed by near-UV CD resulted in an increased exposure of hydrophobic regions of the protein (Fig. 3). Changes in net charge on the HSA surface were also observed. These results suggest that the increase of negative charge reflects oxidation of basic amino acid residues in albumin. In plasma, all amino acids in the protein are susceptible to oxidative modification by oxidants such as hydroxyl radicals and hypochlorous acid. Among them, amino acids such as cysteine, histidine, lysine, and arginine are more vulnerable to oxidation [33]. Modification of these residues results in conformational changes in cases of uremia.

The high-affinity binding of warfarin and ketoprofen was studied in order to determine whether the drug binding properties of HSA had been affected by uremia. In this experiment, the drug binding properties of HD-HSA were found to be reduced. The decreased ketoprofen binding to HD-HSA is most probably caused by conformational changes involving Site II in subdomain IIIA [24]. The decreased binding to HD-HSA might be also caused by the oxidation of <sup>410</sup>Arg. Ahmed et al. [34] recently suggested that <sup>410</sup>Arg is a target site for oxidative stress. Using an anti-carboxyl methyl-Lysine and an anti-hydro-imidazolone antibody to identify advanced glycation endproducts, we also found that HD-HSA could be recognized by this antibody. This indicates that Lys and Arg residues in HD-HSA were modified (data not shown), causing structural changes due to oxidative stress in HD patients. In addition, this alteration may further aggravate the pre-existing elevated concentration of unbound free drugs in HD patients. Therefore, the structural alteration of HSA in uremia results in the loss of some important properties of HSA.

In conclusion, the present study has shown that the oxidative modification of HSA in HD patients may lead to alterations in the conformation as well as the biological properties of HSA. The present study further propose that, in addition to its serum concentration, the quality of HSA molecules may be not only a crucial factor affecting its protective effects but also a risk factor as a pro-oxidant in HD patients. The present study provides evidence to suggest that isolated HSA in hemodialyzed patients has, not only reduced structural and functional properties, but also acts as a mediator of the neutrophil activation state associated with chronic uremia.

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## Characterization of benzodiazepine binding site on human $\alpha_1$ -acid glycoprotein using flunitrazepam as a photolabeling agent

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### Abstract

The binding of flunitrazepam (FNZP) by human  $\alpha_1$ -acid glycoprotein (hAGP) and the relationships between the extent of drug binding and desialylation and the genetic variants of hAGP were examined. The photolabeling specificity of [<sup>3</sup>H]FNZP was confirmed by findings in which other hAGP-binding ligands inhibited the formation of covalent bonds between [<sup>3</sup>H]FNZP and hAGP. The photolabeling of asialo-hAGP suggested that sialic acid does not involve in the binding of [<sup>3</sup>H]FNZP. No difference in the labeling could be found between the F1\*S variants and A variant. Similarly, FNZP did not show a difference in binding affinity to the two genetic variants of hAGP. Sequence analysis of the photolabeled peptide indicated a sequence corresponding to Tyr91-Arg105 of hAGP.

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**Keywords:** Human alpha 1-acid glycoprotein; Flunitrazepam; Binding site; Topology analysis; Photoaffinity labeling

### 1. Introduction

Human  $\alpha_1$ -acid glycoprotein (hAGP), an acute phase protein in blood, consists of 183 amino acid residues and five N-linked oligosaccharides, with a molecular weight of 41–43 kDa [1]. It is negatively charged due to the presence of sialic acids in its glycan chains [2]. hAGP can be produced as three main genetic variants, the A variant and the F1 and S variants, which are encoded by two different genes [3]. The proportion of each variant are approximately F1 50%, S 35% and A 15% [4]. The relative occurrence of the three main phenotypes of native hAGP composition in the population was found to be about 50% for F1+S+A, 35% for F1+A and 15% for S+A [5]. There is a difference of at least 22

amino acid residues between the F1\*S and A variants while the F1 and S forms differ by only a few residues [3,6]. F1\*S and A variants also differ in the number of Met residues.

hAGP's basal level is approximately 20  $\mu$ mol/L, but it can vary from 5- to a 10-fold range in response to stress, infection, or the effects of neoplasm in evocation of an inflammatory response [7]. In addition, changes in the expression of genetic variants of hAGP could also occur apart from its glycosylation pattern depending on the type of inflammation [8]. The biological function of this protein is not clear, though it has been reported to have an anti-inflammatory and immunomodulating role as well as protective effects [9].

In the clinical practice, hAGP is a valuable diagnostic and prognostic tool. For example, increased hAGP levels associated with advanced tumors altered the pharmacokinetics of Imatinib (STI571), a tyrosine kinase inhibitor, in leukemia patients [10]. hAGP also appears to be an independent predictor of response and a major objective prognostic factor of survival in patients with non-small cell

*Abbreviations:* hAGP, Human  $\alpha_1$ -acid glycoprotein; FNZP, Flunitrazepam; PSL, photo stimulated luminescence

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lung cancer treated with docetaxel chemotherapy [11]. In addition, hAGP binds and transports a number of endogenous and exogenous compounds including various basic and neutral drugs influencing their pharmacokinetics and pharmacodynamics [12].

Benzodiazepines are clinically important central nervous system depressants with anxiolytic, sedative, antiepileptic and muscle relaxant therapeutic actions. The major binding protein in plasma for benzodiazepines has been identified as serum albumin. However, hAGP had been shown to bind benzodiazepines with comparable affinity by Maruyama et al. [13]. In a previous study, we have shown that diazepam was able to inhibit the photoincorporation of FNZP to hAGP but not to albumin, suggesting that certain benzodiazepines bind more readily to hAGP instead of albumin and may involve different binding mechanism [14]. During acute phase reaction, albumin synthesis is decreased while that of hAGP may increase. Thus, the pharmacokinetics of drugs that are bound by both of these two proteins may be altered [15].

Due to the clinical significance of hAGP binding of drugs, especially during acute phase reactions, the topology of its binding site has been investigated with a variety of methods including chemical modification, quantitative structure activity relationship (QSAR) methods and molecular modeling [16–18]. In a study of the binding of nine diazepam to hAGP by means of fluorescence and CD (circular dichroism) spectroscopies at our laboratory, we found that benzodiazepines have one high affinity binding site [13,19].

X-ray crystallographic analysis and nuclear magnetic resonance are the major modern direct techniques to elucidate the structures of macromolecules and their complexes at atomic resolution [20,21]. However, the application of these methods to hAGP–drug complexes is not feasible at present due to the great difficulty in the crystallization of this serum glycoprotein which comprises a mixture of variants as well as glycoforms, and also lacking of a suitable expression system that is able to produce a recombinant hAGP with correct physiologically related glycans.

Photoaffinity labeling is one of the most powerful techniques, which can be used to study a ligand binding protein [22,23]. In a recent study, we have shown that [<sup>3</sup>H]FNZP (Fig. 1) was able to form covalent bond upon photoirradiation to hAGP which can be inhibited by diazepam [14]. This paper reports further attempt using [<sup>3</sup>H]FNZP as a photolabeling agent to elucidate the topography of binding site amino acid sequence of hAGP and to shed light on the possible binding mechanisms.

## 2. Materials and methods

### 2.1. Materials

[<sup>3</sup>H]FNZP (71 Ci/mmol) was purchased from PerkinElmer Life Sciences. hAGP (purified from cohn fraction

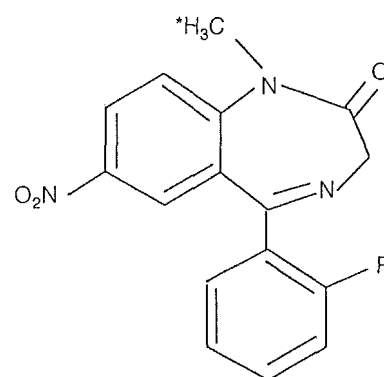


Fig. 1. Chemical structure of [<sup>3</sup>H]FNZP. \*: <sup>3</sup>H-labeled position.

VI) was purchased from Sigma Chemical Co. Sequencing grade modified trypsin was purchased from Promega Co. (USA). Cyanogen bromide (CNBr), dithiothreitol, trifluoroacetic acid, ammonium bicarbonate, and imidazole were purchased from Nacal Tesque (Kyoto, Japan). 4-Vinylpyridine was purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). Neuraminidase, resorcinol, propranolol, lidocaine, verapamil, disopyramide, dipyridamole, testosterone, indomethacin, and salicylic acid were purchased from Sigma. *N*-glycosidase F recombinant (PNGase F) was purchased from Roche (Germany). Progesterone was from Nihon Schering. FNZP, diazepam, fludiazepam (Sumitomo Pharmaceuticals), chlorpromazine, imipramine (Mitsubishi Wellpharma), warfarin (Eisai), and phenylbutazone (Ciba-Geigy Japan) were donated by the respective companies. Acenocoumarin was a gift from Dr. Janssen from the Faculty of Pharmacy, Utrecht University, the Netherlands. All other chemicals and solvents were of analytical grade.

### 2.2. Photoaffinity labeling of hAGP

hAGP (50 μM) was incubated with [<sup>3</sup>H]FNZP (25 μM) in 100 μL of 20 mM Tris–HCl, pH 7.4, in a 1.5 mL Eppendorf tube at room temperature in the dark for 60 min. The incubation mixture was then placed on ice and irradiated for 30 min by a 100 W black light/blue lamp (Ultra-Violet Products, Inc., San Gabriel, CA, USA) at a distance of 10 cm. After irradiation, the photolabeled hAGP was precipitated by adding 1 mL of acetone, followed by centrifugation at 15 × 1000 rpm for 10 min. The pellet was washed with 1 mL of ethanol and centrifuged a second time.

### 2.3. Cyanogen bromide cleavage

For cyanogen bromide cleavage reactions, the pyridylethylated pellet was then dissolved in 100 μL of cyanogen bromide (CNBr) in 70% formic acid (CNBr:Met=400:1) and incubated under N<sub>2</sub> for 24 h in the dark at room temperature. One milliliter of milli-Q water was added at the end of the CNBr cleavage and the mixture was lyophilized.



#### 2.4. SDS-PAGE and electroblotting

Photolabeled hAGP was analyzed on SDS-PAGE using a 10% polyacrylamide gel according to Laemmli and a sampling buffer (10 mM Tris-HCl [pH 7.6], 1% [w/v] SDS, 20 mM dithiothreitol, 4 mM ethylenediaminetetraacetic acid and 2% [w/v] sucrose) [24]. The concentration of the protein was determined by Bradford assay with bovine serum albumin as standard [25]. The gel after electrophoresis was electrophoretically transferred onto a polyvinylidene difluoride membrane in a transfer buffer (25 mM Tris, 193 mM glycine, 10% methanol) by using a semidry blotting assembly. The blotted membrane was stained with Coomassie Brilliant Blue R250, followed by drying completely in air.

#### 2.5. Autoradiographic analysis

For autoradiographic analysis, the dried PVDF membrane was then placed in contact with an imaging plate (BAS III, Fuji Photo film Co.) in a cassette (BAS cassette 2040) at room temperature for 48 h. The imaging plate was scanned and analyzed using a Bio-Imaging Analyzer BAS 5000 model (Fuji Photo Film Co.) and analyzed using the Fuji Film Science Lab 98 software, L Process V1.6. The incorporation of radioactivity into individual fragments was quantified by using the software Image Gauge V3.1 (Fuji Film).

#### 2.6. Reductive pyridylethylation and deglycosylation of hAGP

After the photolabeled hAGP was precipitated by acetone, 100  $\mu$ L of the buffer was added to the protein precipitate. 10  $\mu$ L of 1% sodium dodecyl sulfate (SDS) and 1 M (2-mercaptoethanol) were added to this solution, before reduction were performed at 100 °C at 10 min. For deglycosylation of hAGP, 10  $\mu$ L of 10% *n*-octanoyl-*N*-methylglucamide (MEGA-8), 50  $\mu$ L of deionized water and 2 units of PNGase F were added to the reduction solution, and this solution was incubated for 24 h. After the addition of 1  $\mu$ L of 4-vinylpyridine, the mixture was further incubated under N<sub>2</sub> for 30 min at room temperature in the dark and dialyzed for desalination.

#### 2.7. Separation of hAGP Genetic Variants

The hAGP genetic variants were separated using the method of Herve et al. [26]. An iminodiacetate (IDA) Sepharose gel loaded with copper (II) ions and equilibrated in buffer 1 (20 mM sodium phosphate buffer, pH 7.0, containing 0.5 M sodium chloride) was packed into a glass column (2.0  $\times$  30.0 cm L, Pharmacia LKB). Commercial hAGP (70 mg dissolved in 1.0 mL of buffer 1) was applied to the column at a flow rate of 1.0 mL/min. Fractions

(10 mL) were collected, and their respective absorbance values were determined spectrometrically at 280 nm. As soon as the F1\*S variant flowed through the column, elution buffer 2 (buffer 1 plus 20 mM imidazole) was applied to the column to remove the bound A variants. The peak fractions of each eluent were collected, concentrated on YM 10 membrane filter (Amicon, Danvers, MA), dialyzed against deionized water, and lyophilized. The purities of the isolated hAGP preparations were determined by isoelectric focusing (IEF) followed by incubation at 37 °C for 24 h with 1 unit of neuraminidase [27].

#### 2.8. Desialylation of hAGP

hAGP was desialylated enzymatically, using the methodology described by Primoic and McNamara [27], using an immobilized neuraminidase obtained from *Clostridium perfringens*. The sialic acid content was determined by the thiobarbituric acid method. The product was dialyzed against deionized water and the dialysate was lyophilized. Approximately 95% of the sialic acid was removed, leaving an average of one sialic acid residue per protein molecule. The molecular weight of desialylated hAGP was therefore 40,000.

#### 2.9. Determination of binding constant

Aliquots of various ratios of FNZP (5–40  $\mu$ M)–hAGP (10  $\mu$ M) were transferred to amicon MPS-1 micropartition system with YMT ultrafiltration membrane, centrifuged at 2000 rpm for 40 min at 25 °C. The free concentration of FNZP was determined by HPLC in a system consisting of a Hitachi 655A-11 pump and Hitachi 655A at 250 nm. A column of LichroCART 250-4, Lichrospher, RP-Select B (5  $\mu$ m, 4 mm i.d.  $\times$  250 mm; Kanto Chemical) was used. The mobile phases consisted of 0.02 M KH<sub>2</sub>PO<sub>4</sub> buffer (pH 2.5)–acetonitrile (6:4, v/v).

Binding parameters were determined by fitting the experimental data to the following Scatchard equation using a non-linear squares program (MULTI program) [28].

$$\frac{r}{D_f} = nK - rK$$

where *n* is the number of binding sites, *K* the binding constant and *D<sub>f</sub>* the free drug concentration, with *r* denoting the moles of bound ligand per mole of total protein.

#### 2.10. Tryptic digestion and purification of photolabeled hAGP peptide fragments

Tryptic digestion was done in 50 mM NH<sub>4</sub>HCO<sub>3</sub> (pH 7.8). After deglycosylation, deglycosylated hAGP was incubated with trypsin for 5 h at 37 °C. The ratio of a trypsin:hAGP is 1:20 (w/w). Tryptic peptides were sepa-

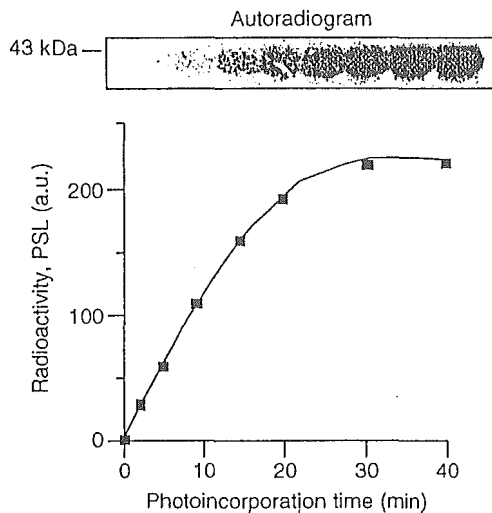


Fig. 2. Time course of  $[^3\text{H}]\text{FNZP}$  photoincorporation.  $50\ \mu\text{M}$  hAGP was incubated with  $25\ \mu\text{M}$   $[^3\text{H}]\text{FNZP}$  for 60 min prior to photoirradiation. Aliquots of  $100\ \mu\text{L}$  was taken from the mixture solution at each time point as stated in the figure during irradiation until 30 min. All samples were separated with 10% SDS-PAGE and electroblotted onto a PVDF membrane for autoradiographic analysis. PSL: photo stimulated luminescence.

rated by reverse-phase  $\text{C}_{18}$  HPLC using an aqueous acetonitrile gradient in the presence of 0.1% TFA. The separated peptides fractionated every 30 s and  $200\ \mu\text{L}$  of each fraction was added to 2.5 mL of scintillation cocktail, and the radioactivity was determined using a LSC-500 liquid scintillation counter (Aloka, Tokyo, Japan). Then, the fraction that has the highest radioactivity was collected and was evaporated on a speedvac evaporator until the volume of the sample reached about  $50\ \mu\text{L}$  in the Eppendorf tube.

### 2.11. Capillary HPLC separation and sequencing

After evaporating,  $10\ \mu\text{L}$  of the sample was then injected onto the ABI 173 A MicroBlotter Capillary HPLC System, Perkin Elmer. The sample was manipulated following the manufacturer's instructions (User's Manual, 90,3836 Rev A, November 1995, Perkin Elmer). Meanwhile, the blotted membrane from the capillary HPLC separation was in contact with an imaging plate for 48 h prior to autoradiographic analysis. The PVDF membrane was positioned with the chromatogram of a peptide map from the ABI 173 A MicroBlotter Capillary HPLC System. Portions of the PVDF membrane were excised for sequencing on an Applied Biosystems Procise Sequencer with reference to the autoradiogram.

### 2.12. Statistical analysis

All data are presented as means  $\pm$  S.D. Statistical analysis of difference was determined by one-way ANOVA followed by the modified Fisher's least squares difference method.

## 3. Results

### 3.1. Photolabeling of $[^3\text{H}]\text{FNZP}$ to hAGP

The autoradiogram in Fig. 2 shows that a band of radiolabeled protein appeared only upon the photoirradiation of hAGP with  $[^3\text{H}]\text{FNZP}$  in a time-dependent manner. The radioactivity band indicated covalent incorporation of  $[^3\text{H}]\text{FNZP}$  to hAGP via photoirradiation. The absence of a radioactive band in the sample without irradiation indicated that no covalent attachment of  $[^3\text{H}]\text{FNZP}$  to hAGP occurred in the dark. A 30-min exposure to light was sufficient for the covalent attachment of  $[^3\text{H}]\text{FNZP}$  to hAGP (Fig. 2).

### 3.2. Inhibitory effects of various drugs on binding of $[^3\text{H}]\text{FNZP}$ to hAGP

Previous studies indicated that hAGP has one common drug binding site, which appears to be wide and flexible, comprising basic drugs, acidic drugs and steroid hormone binding regions [29]. In the photoinhibition experiment, all competitors that are known to bind hAGP, except phenylbutazone, successfully inhibited the photoincorporation of FNZP to an extent of more than 50%. In contrast, salicylic acid, which is not a binding ligand of hAGP, did not inhibit the photoincorporation (Fig. 3).

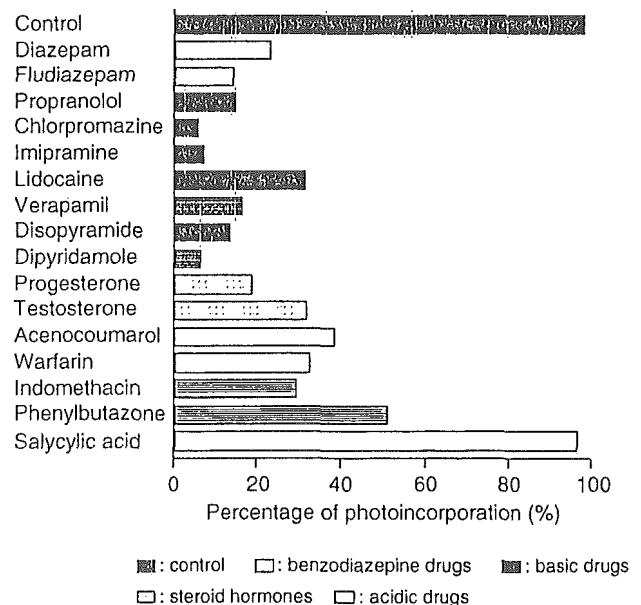


Fig. 3. Photoincorporation percentage of hAGP with  $[^3\text{H}]\text{FNZP}$  in the presence of various inhibitors. hAGP ( $50\ \mu\text{M}$ ) was incubated with  $[^3\text{H}]\text{FNZP}$  ( $25\ \mu\text{M}$ ), in the presence of competitors ( $250\ \mu\text{M}$ ) prior to photolysis. The photolabeled hAGP was cleaved with CNBr, separated with a 10% gel SDS-PAGE and electroblotted onto a PVDF membrane before being subjected to autoradiographic analysis. Radioactivity of the 34 kDa band was compared among different competitors.

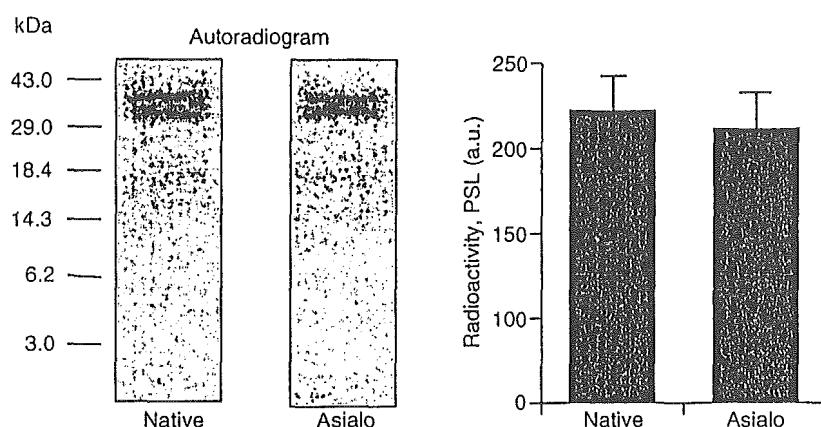


Fig. 4. Cyanogen bromide fragments of native and asialo hAGP photolabeled with [ $^3\text{H}$ ]FNZP. 50  $\mu\text{M}$  hAGP (native or asialo hAGP) was incubated with 25  $\mu\text{M}$  [ $^3\text{H}$ ]FNZP for 60 min prior to photoirradiation, and after photoirradiation, cleaved with CNBr and separated with tricine gel electrophoresis and electroblotted onto a PVDF membrane for autoradiographic analysis.

### 3.3. Autoradiogram of CNBr fragments of native and asialo hAGP photolabeled with [ $^3\text{H}$ ]FNZP

hAGP possesses five N-glycan chains with sialic acids as the terminal group. The carbohydrate content of hAGP is about 45% (w/w) of molecular weight, including 14 sialic acid residues per molecule [6,30]. Sialic acid residues have been reported to influence the binding of several drugs to hAGP [31]. Nevertheless, photolabeling experimental results showed no significant difference in the extent of photoincorporation of FNZP into asialo-hAGP and native hAGP (Fig. 4). This indicated that sialic acids do not play an important role in the binding interaction of FNZP to hAGP.

### 3.4. Autoradiogram of CNBr fragments hAGP variants photolabeled with [ $^3\text{H}$ ]FNZP

The A variant and the F1\*S variant differ in their amino acid sequence by at least 22 residues (Fig. 5) [3,6,32]. Large differences in the binding of various drugs have been reported for the F1\*S vis-a-vis the A variant [17]. The most abundant F1\*S variant possesses only one Met at position

111, whereas the A variant possesses two Met at positions 111 and 156. The electrophoretic pattern of the CNBr fragment of native hAGP will thus show a band of about 34 kDa and three bands of smaller molecular weights (about 9.8, 6.0 and 3.8 kDa) (Fig. 6). The peptide band of about 34 kDa corresponds to residues 1–111, and three bands of about 9.8, 6.0 and 3.8 kDa correspond to residues 112–183 and 112–156, and 157–183, respectively. Since the 34-kDa peptide of hAGP photolabeled with [ $^3\text{H}$ ]FNZP showed the highest radioactivity, the covalent bond formation site of FNZP is located within the sequence from the N-terminus to Met111. In addition, no significant difference could be found for the extent of photoincorporation of FNZP among the two genetic variants (F1\*S and A). Furthermore, there is no significant difference in the  $K_a$  values of FNZP to native hAGP, F1\*S and A (Table 1, Fig. 7).

### 3.5. Amino acid sequence of the photolabeled tryptic peptides

The tryptic digestion of the sample was separated and simultaneously blotted onto a strip of PVDF membrane

F1*S	QIPLCANLVP	10	VPITNATLDQ	20	ITGKWFYIAS	30	AFRNEEYNKS	40
A	QIPLCANLVP		VPITNATLDR		ITGKWFYIAS		AFRNEEYNKS	
	VQEIQATFFY	50	FTPNKTEDTI	60	FLREYQTRQD	70	QCQYNTTYLN	80
	VQEIQATFFY		FTPNKTEDTI		FLREYQTRQN		QCQYNTSSYLN	
	VQRENGTISR	90	YVGGQEHFAH	100	LLLRDRTKY	110	MLAFDVNDEK	120
	VQRENGTVSR		YEGGREHVAH		NLELRDRTKL		MFGSYLDDEK	
	NWGLSVYADK	130	PETTKEQLGE	140	FYEALDCLR	150	PKSDVVYTDW	160
	NWGLSFYADK		PETTKEQLGE		FYEALDCLCI		PRSDVMYTDW	
	KKDKCEPLEK	170	QHEKERKQEE	180	GES			
	KKDKCEPLEK		QHEKERKQEE		GES			

Fig. 5. Amino acid sequence of hAGP variants. Differences in the amino acid sequence between the F1\*S and A variant are underlined.

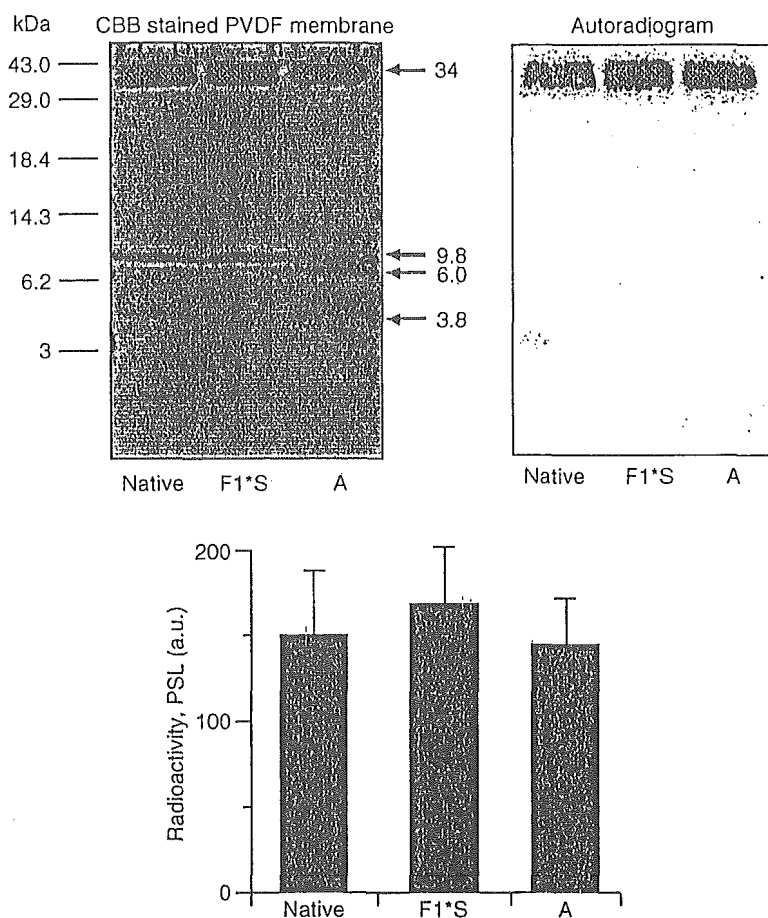


Fig. 6. Cyanogen bromide fragments of native and hAGP variants photolabeled with [ $^3\text{H}$ ]FNZP. 50  $\mu\text{M}$  hAGP (native, F1\*S or A) was incubated with 25  $\mu\text{M}$  [ $^3\text{H}$ ]FNZP for 60 min prior to photoirradiation, and after photoirradiation, cleaved with CNBr and separated with tricine gel electrophoresis and electroblotted onto a PVDF membrane for autoradiographic analysis.

with an ABI 173 A MicroBlotter Capillary HPLC System. Autoradiographic analysis of the PVDF membrane indicated that the radioactive spot corresponded to the observed peak at about 40 min (Fig. 8a, b). Edman sequencing performed on the tryptic peptide of the hAGP photolabeled with [ $^3\text{H}$ ]FNZP revealed an amino acid sequence of YVGGQEHFA (Fig. 9), corresponding to the Tyr91-Arg105 peptide sequence on F1\*S variant of hAGP digested with trypsin. The YEGGREHVA sequence (91–99 in variant A) was not detected along with the YVGGQEHFA (F1\*S variant) sequence, could be due to the presence of an arginine residue for variant A at position 95, thereby producing much shorter tryptic peptides, which were not eluted at a similar retention time as the peptide of F1\*S variant containing the YEGGREHVA sequence. In addition, the lower abundance of

variant A (15%) in the AGP sample might have contributed to the observation too [4].

#### 4. Discussion

FNZP has been successfully used as a photoaffinity labeling agent in topology analysis of the GABA receptors

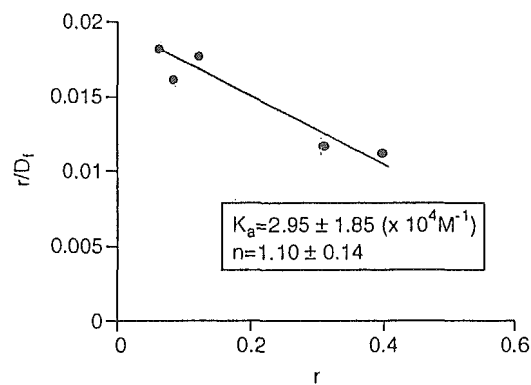


Fig. 7. Typical Scatchard plot of FNZP to hAGP. Binding experiments were performed at pH 7.4 and 25  $^{\circ}\text{C}$ .

Table 1  
Binding parameter of FNZP to hAGP variants at pH 7.4 and 25  $^{\circ}\text{C}$

hAGP	$K_a$ ( $\times 10^4 \text{ M}^{-1}$ )	$n$
Native hAGP	$2.95 \pm 1.85$	$1.10 \pm 0.14$
F1*S	$2.92 \pm 2.29$	$1.06 \pm 0.14$
A	$2.67 \pm 1.24$	$0.95 \pm 0.26$

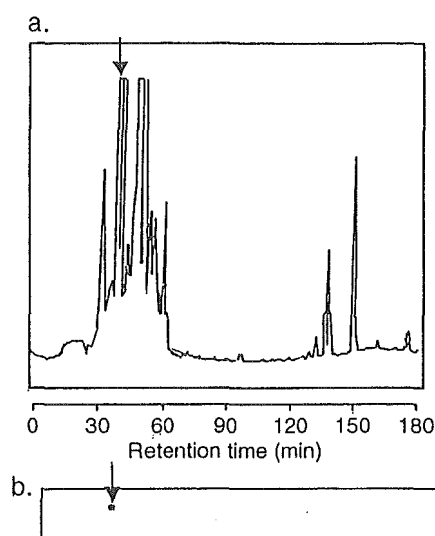


Fig. 8. Chromatogram of capillary HPLC and autoradiogram of blotted PVDF membrane. The major radioactive peptides were collected and concentrated with a speedvac evaporator. The concentrated sample from previous reversed-HPLC analysis was separated and simultaneously blotted onto a strip of PVDF membrane with an ABI 173 A MicroBlotter Capillary HPLC System. The blotted membrane from the capillary HPLC separation was in contact with an imaging plate for 48 h prior to autoradiography analysis. (a) peptides detected at UV-absorption (210 nm); (b) autoradiogram of blotted PVDF membrane.

[33,34]. In order to elucidate the binding cavity topology of hAGP, we used FNZP as a photolabeling probe. The covalent bond formation site of FNZP is located within the sequence from the N-terminus to Met111 (Fig. 6). Previous studies indicated that hAGP has one common drug binding site, which appears to be wide and flexible, comprising basic drugs, acidic drugs and steroid hormone binding regions [29]. Differences in the binding of drugs to this two main genetic variants have been reported: Disopyramide, imipramine and methadone bind selectively to the A variant, whereas dipyrindamole, quinidine and mifepristone bind preferentially to the F1\*S variant. Progesterone, propranolol and chlorpromazine show no selectivity in binding [17]. No significant difference in the  $K_a$  values of FNZP to native hAGP, F1\*S variant and A variant could be found among these three species of proteins (Table 1). In agreement with the binding experiment results, no significant difference could be found for the extent of photoincorporation of FNZP among these two species of proteins (Fig. 6). These results suggest that FNZP indiscriminately binds to F1\*S and A variants. In addition, in the photoinhibition experiment, all drugs that are known to bind hAGP, except phenylbutazone, successfully inhibited the photoincorporation of FNZP to an extent of more than 50%. In contrast, salicylic acid used as a negative control which is not a binding ligand of hAGP did not inhibit the photoincorporation (Fig. 3).

hAGP possesses five N-glycan chains, which have di-, tri-, and/or tetraantennary structures, with sialic acids as the terminal group. The glycan structures show microheteroge-

neity under physiological conditions [35], and the partially desialylated hAGP is known to exist in plasma of patients with liver disease [36]. Sialic acid residues have been reported to influence the binding of several drugs to hAGP [31]. For example, the binding affinity of chlorpromazine was significantly decreased upon the elimination of sialic acid residues [37], whereas those of several phenothiazine neuroleptics, except chlorpromazine, decreased moderately [38]. In contrast, there are also reports that indicate the noninvolvement of sialic acid in the binding of drugs such as dicoumarol [39,40]. Our photolabeling experimental results showed no significant difference in the extent of the photoincorporation of FNZP into asialo-hAGP and native hAGP (Fig. 4). This indicated that sialic acids do not play an important role in the binding interaction of FNZP to hAGP.

Edman sequencing of the peptide photolabeled with [ $^3\text{H}$ ]FNZP revealed an amino acid sequence of YVGG-QEHFA (Fig. 9), corresponding to the Tyr91-Arg105 peptide sequence on hAGP digested with trypsin. This is in agreement with our previous finding shown in Fig. 3. Since the His102 residue of alpha subunit of the GABA ( $\gamma$ -aminobutyric acid) receptor has been identified as the photolabeled site of [ $^3\text{H}$ ]FNZP [34], we speculate that His97 and His100 of hAGP may be the amino acid residues forming a covalent bond with FNZP. Confirmation experiments are currently underway.

Maruyama et al. reported that the binding parameters ( $nK$ ) of the benzodiazepine:HSA systems are slightly higher than those of the benzodiazepine:AGP systems [13]. Since benzodiazepines bind to both albumin and AGP at comparable affinity ( $K_a=10^4$ – $10^5$ ), binding to AGP in the acute phase is expected to have higher clinical significance because hAGP concentration increases 3- to 10-fold during infection, diseases with inflammatory component as well as psychiatric illness. In addition, the plasma albumin level decreases during acute phases [41]. This could significantly alter the free plasma concentrations of the ligand and contribute to its variable kinetic behavior [42].

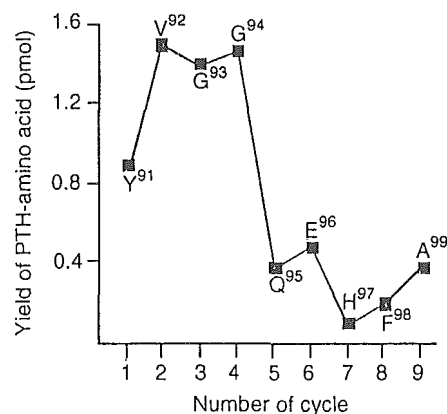


Fig. 9. N-terminal amino acid sequence analysis by the Edman degradation method. PTH: phenylthiohydantoin.

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# Cooperative effect of hydrophobic and electrostatic forces on alcohol-induced $\alpha$ -helix formation of $\alpha_1$ -acid glycoprotein

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**Abstract**  $\alpha_1$ -Acid glycoprotein (AGP) is a serum glycoprotein that mainly binds basic drugs. Previous reports have shown that AGP converts from a  $\beta$ -sheet to an  $\alpha$ -helix upon interaction with biomembranes. In the current studies, we found that alkanols, diols, and halogenols all induce this conformational change. Increased length and bulkiness of the hydrocarbon group and the presence of a halogen atom promoted this conversion, whereas the presence of a hydroxyl group inhibited it. Moreover, the effect was dependent on the hydrophobic and electrostatic properties of the alcohols. These results indicate that, in a membrane environment, hydrophobic and electrostatic factors cooperatively induce the transition of AGP from a  $\beta$ -sheet to an  $\alpha$ -helix.

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**Keywords:**  $\alpha_1$ -Acid glycoprotein; Alcohol;  $\alpha$ -Helix formation; Conformational transition

## 1. Introduction

$\alpha_1$ -Acid glycoprotein (AGP), a member of the lipocalin family, is a polypeptide with two internal disulfide bonds and five carbohydrate chains that account for ~40% of its total mass (36 kDa) [1,2]. It is a major binding protein for neutral and basic ligands [3–5]. Although the three-dimensional structure and biological functions are still unknown, circular dichroism (CD) measurements [6] and molecular modeling [7] have revealed that this protein is mostly made up of  $\beta$ -sheets in aqueous solution.

It is widely accepted that membrane transport of a drug depends on the free drug concentration in solution. However, because this hypothesis does not fully explain the uptake mechanism for some AGP-binding drugs, a protein-mediated uptake system has been proposed [8–11]. In such a system, structural changes in the protein due to interaction with membrane surfaces decrease the drug-binding capacity. We previ-

ously reported that the interaction between AGP and model biomembranes (reverse micelles and liposomes) results in a unique conformational transition ( $\beta$ -sheet to  $\alpha$ -helix) and a decrease in ligand-binding capacity [12,13]. Other studies of AGP interacting with vesicles [14] and liposomes [15] also support the conclusion that AGP interacts with the membrane in circulation.

A part of this interaction is due to the fact that the membrane potential decreases the local pH relative to the bulk solution [16]. The difference in pH has been experimentally determined to be 1.6 pH units, and the calculated value reaches 2.7 pH units [17]. Indeed, there are several reports that proteins, including other lipocalins, undergo structural and functional changes under mild acidic conditions on the membrane surface [18–21]. Also, Bychkova et al. [22] reported that cytochrome *c*, a mitochondrial protein, has a molten globule state in the presence of 40% methanol (MeOH) at pH 4.0, a condition that mimics the acidic conditions and low dielectric constant of the membrane surface environment.

There are other reports that a protein with high propensity for  $\alpha$ -helix formation can easily convert to the  $\alpha$ -helix form at pH 2.0 in the presence of various alcohols [23–26]. In addition, Kodicek et al. [27] reported that AGP heated in the presence of methanol (MeOH) forms a similar  $\alpha$ -helix structure. These reports suggested that this effect was due to the hydrophobic force of alcohol and this force might induce the  $\alpha$ -helix formation in AGP. On the other hand, the factors in such conformational transition of AGP are not elucidated.

In the present study, we used various alcohols (alkanols, diols, and halogenols) to investigate the mechanism of the  $\alpha$ -helix formation in AGP and use of these alcohols allowed examination of how negative charge on the membrane surface and hydrophobic interaction inside the membrane affects the  $\alpha$ -helix formation in AGP.

## 2. Materials and methods

### 2.1. Materials

AGP (Cohn Fraction VI) was purchased from Sigma Chemical Co. (St. Louis, MO, USA). All alcohols were purchased from Nacalai Tesque (Kyoto, Japan). All other chemicals and solvents were of analytical grade.

### 2.2. Measurement of CD spectra

CD spectra were recorded in a 1-mm path length cell with a JASCO J-720 spectropolarimeter using 10  $\mu$ M AGP in 20 mM sodium acetate buffer, pH 4.0. The data were expressed as the molar residue ellipticity [ $\theta$ ] at 222 nm, which is an index of  $\alpha$ -helix content.

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**Abbreviations:** The abbreviations used for the alcohols are summarized in Table 1;  $\alpha_1$ -acid glycoprotein; CD, circular dichroism; ASA, solvent-accessible surface area; ASA (T), total ASA; rASA (H), relative ASA of hydrophobic region; rASA (N), relative ASA of negative charge region; rASA (P), relative ASA of positive charge region; HE<sub>50</sub>, concentration of alcohol needed to form 50%  $\alpha$ -helix structure



### 2.3. Calculation of $m$ values

To analyze the effects of alcohols on AGP, we assumed a two-state transition between the  $\beta$ -sheet and the  $\alpha$ -helix states. The equilibrium constant for the conformational transition,  $K$ , is defined as  $K = [H]/[B]$ , where  $[H]$  and  $[B]$  are the concentrations of the  $\alpha$ -helix and  $\beta$ -sheet states, respectively. The free energy change ( $\Delta G_f$ ) for formation of the helical state is calculated according to following equation:

$$\Delta G_f = -RT \ln K,$$

where  $R$  is the gas constant, and  $T$  is the temperature in Kelvin. We assumed a linear dependence of  $\Delta G_f$  upon the alcohol concentration ([alcohol]):

$$\Delta G_f = \Delta G_0 - m[\text{alcohol}],$$

where  $\Delta G_0$  is the  $\Delta G_f$  value in the absence of alcohol, and  $m$  is a measure of the dependence of  $\Delta G_f$  on the concentration of alcohol.

Least squares curve fitting was done using the MULTI program [28]. The solvent-accessible surface areas (total ASA [ASA (T)], hydrophobic ASA [ASA (H)], negative charge ASA [ASA (N)], and positive charge ASA [ASA (P)]) of alcohols were estimated using the Molecular Operating Environment (Chemical Computing Group Inc., Canada). ASA (H)/ASA (T), ASA (N)/ASA (T) and ASA (P)/ASA (T) were represented as relative ASA (H) [rASA (H)], relative ASA (N) [rASA (N)] and relative ASA (P) [rASA (P)], respectively (see Table 1).

## 3. Results

### 3.1. Effects of alkanols on $\alpha$ -helix formation in AGP

Based on a previous report that 40% methanol can induce  $\alpha$ -helix formation at pH 4.0 [22], we investigated the effect of hydrocarbon groups in alcohols on the structure of AGP (Fig. 1A). The  $[\theta]$  value at 222 nm, an index of  $\alpha$ -helix content, showed saturation at lower concentrations as the alkyl chain length was increased. This demonstrated that the capacity to induce  $\alpha$ -helix formation was increased as the hydrocarbon chain length was increased. However, there did not seem to be a difference in the effects of various alkanols on the  $\alpha$ -helix content of AGP. Furthermore, 1-propanol, which has a straight chain, was more effective in promoting  $\alpha$ -helix formation than 2-propanol, which has a branched chain.

### 3.2. Effects of diols on $\alpha$ -helix formation in AGP

Next, to determine the role of the hydroxyl group, we examined the effects of several diols on the  $\alpha$ -helix content. As in the case of the alkanols, transition to an  $\alpha$ -helix structure was ob-

served at lower concentrations as the chain length was increased (Fig. 1B). However, the effect was weaker than in the case of the alkanols. This result suggests that the hydroxyl group of the alcohol only plays a role in their dissolution in water and that the induction of  $\alpha$ -helix formation is due to the hydrocarbon moiety.

### 3.3. Effects of halogenols on $\alpha$ -helix formation in AGP

In addition, we examined the effect of halogen moieties by using several halogenols (Fig. 1C). The  $[\theta]$  value at 222 nm in 2-chloroethanol (ClEtOH) showed the transition and saturation at lower concentrations than for the alkanols and diols, and caused a remarkable increase in the  $\alpha$ -helix content. In contrast, the effect of 2-fluoroethanol (FEtOH) was weaker than that of ethanol. Bromoethanol could not be used because of phase separation. Alternatively, 2,2,2-trifluoroethanol (TFE) and 1,1,1,3,3,3-hexafluoro-2-ethanol (HFIP) was used. It was observed that these alcohols strongly promote an  $\alpha$ -helix formation in AGP. These results imply that the presence of multiple F atoms increases the effectiveness markedly, although the F atom itself is not so potent and halogen atoms markedly increase the ability of the solvent to induce  $\alpha$ -helix formation in AGP.

### 3.4. Effects of hydrophobic force and negative charge on $\alpha$ -helix formation in AGP

Based on the findings with various alcohols, it appeared likely that the hydrophobicity and negative charge of the alcohol participate in the induction of  $\alpha$ -helix formation in AGP. Thus, we estimated rASA (H), rASA (P), and rASA (N), respectively, which do not depend on the molecular weight or ASA (T), to evaluate the properties of the alcohols. The effects of the alcohols on AGP were evaluated using with the  $m$  value and the  $\alpha$ -helix effect<sub>50</sub> (HE<sub>50</sub>), which is the concentration (%) resulting in a 50% transition from a  $\beta$ -sheet to an  $\alpha$ -helix. For all kinds of alcohols, HE<sub>50</sub> appears to more accurately describe the effects of the alcohol than the  $m$  value because the  $m$  value depends on maximum  $\alpha$ -helix content. In Fig. 2, correlations between HE<sub>50</sub> and rASA were presented. In alkanols, a good correlation was observed between the HE<sub>50</sub> and rASA (H) ( $r = 0.8849$ ), suggesting that hydrophobic interactions are involved in  $\alpha$ -helix formation in AGP at pH

Table 1  
The  $m$ , HE<sub>50</sub>, rASA (H), rASA (N) and rASA (P) values for various alcohols

	Abbreviation used	$m$ (J mol <sup>-1</sup> M <sup>-1</sup> )	HE <sub>50</sub>	rASA (H)	rASA (N)	rASA (P)
<i>Alkanols</i>						
Methanol	MeOH	573.7	35.1	0.41	0.17	0.42
Ethanol	EtOH	751.1	28.8	0.67	0.11	0.21
1-Propanol	1-PrOH	952.2	21.8	0.72	0.10	0.17
2-Propanol	2-PrOH	789.8	25.7	0.78	0.09	0.12
<i>Diols</i>						
1,2-Ethandiol	1,2-Et(OH) <sub>2</sub>	328.9	67.4	0.42	0.21	0.38
1,2-Propanediol	1,2-Pr(OH) <sub>2</sub>	381.2	54.5	0.58	0.17	0.25
1,4-Butanediol	1,4-Bu(OH) <sub>2</sub>	393.2	49.3	0.64	0.12	0.24
2,3-Butanediol	2,3-Bu(OH) <sub>2</sub>	469.5	51.3	0.68	0.13	0.18
<i>Halogenols</i>						
2-Fluoroethanol	FEtOH	625.7	37.1	0.43	0.28	0.28
2-Chloroethanol	ClEtOH	397.4	26.6	0.38	0.42	0.20
2,2,2-Trifluoroethanol	TFE	545.3	23.1	0.20	0.60	0.20
1,1,1,3,3,3-Hexafluoro-2-propanol	HFIP	1004.9	5.8	0.07	0.79	0.12

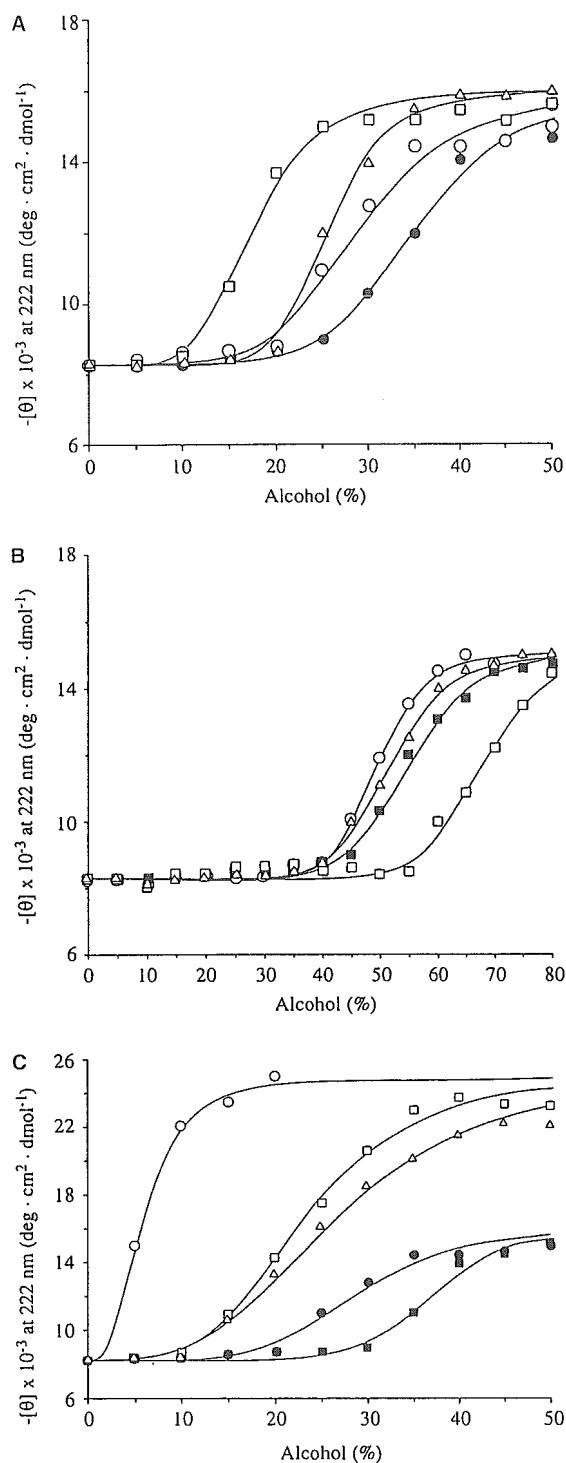


Fig. 1. Effects of alkanols (A), diols (B), and halogenols (C) on  $\alpha$ -helix formation in AGP. The  $\alpha$ -helix content of AGP was monitored at pH 4.0 by measuring the ellipticity at 222 nm. Abbreviations are as follows: in panel A, MeOH ( $\bullet$ ), EtOH ( $\circ$ ), 1-PrOH ( $\square$ ), and 2-PrOH ( $\Delta$ ); in panel B, Et(OH)<sub>2</sub> ( $\square$ ), 1,2-Pr(OH)<sub>2</sub> ( $\blacksquare$ ), 2,3-Bu(OH)<sub>2</sub> ( $\Delta$ ), and 1,4-Bu(OH)<sub>2</sub> ( $\circ$ ); and in panel C, EtOH ( $\bullet$ ), FEtOH ( $\blacksquare$ ), ClEtOH ( $\Delta$ ), TFE ( $\square$ ), and HFIP ( $\circ$ ). The lines were drawn using the MULTI program [28].

4.0 (Fig. 2A). For diols, rASA (H) showed significant correlations with the HE<sub>50</sub> ( $r = 0.9685$ ,  $P < 0.05$ ). This result also indicates that hydrocarbon group contributes to  $\alpha$ -helix for-

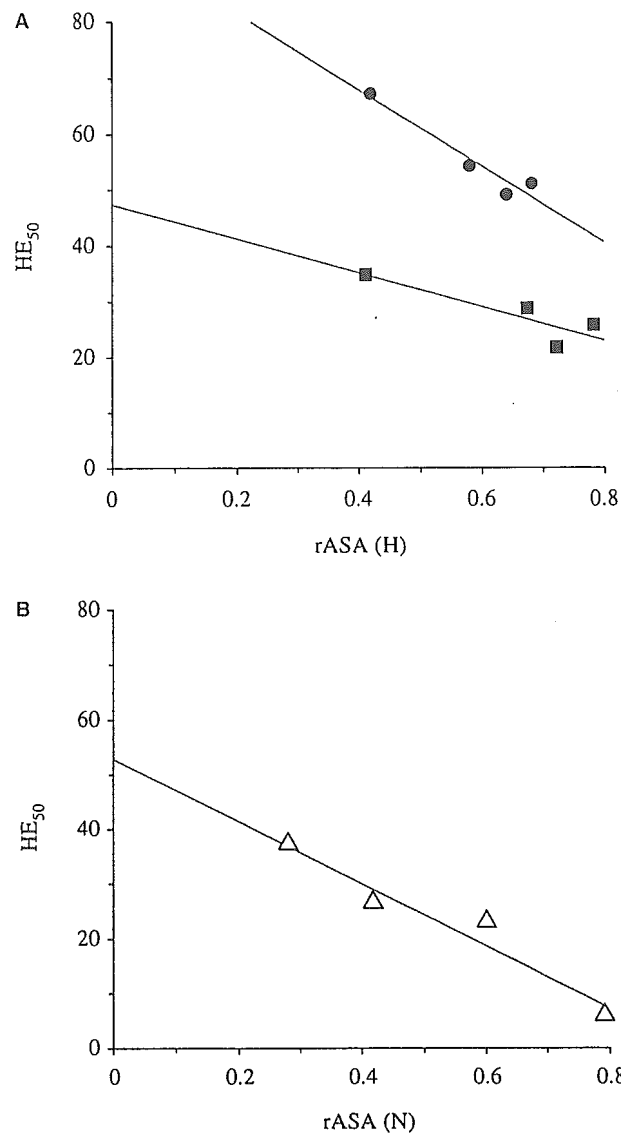


Fig. 2. Correlations between the values of HE<sub>50</sub> and rASA. A and B show the correlations between HE<sub>50</sub> and rASA (H) or rASA (N), respectively. Each symbol represented as follow: alkanol ( $\bullet$ ), diol ( $\blacksquare$ ) and halogenol ( $\Delta$ ).

mation in AGP, although the effect itself was weaker than that of alkanols (Fig. 2A). On the other hand, for halogenols we found a significant correlation between HE<sub>50</sub> and rASA (N) ( $r = 0.9702$ ,  $P < 0.05$ ), which explains the strong effect of the negative charge on  $\alpha$ -helix formation in AGP (Fig. 2B).

### 3.5. Effect of halogen ( $Cl^-$ ) on $\alpha$ -helix formation in AGP

The significant correlation between HE<sub>50</sub> and rASA (N) in halogenols indicates that halogen is an important factor in the induction of  $\alpha$ -helix formation in AGP. Previously, we reported that NaCl (1 M) induced an  $\alpha$ -helix structure at pH 2.0 [12]. To examine the effect of the halogen,  $Cl^-$ , on the  $\alpha$ -helix formation, CD spectra was monitored in the presence of HCl and NaCl (0–1 M) at pH 2.0 (Fig. 3). Interestingly, addition of HCl induced the  $\alpha$ -helix structure, although AGP was in an unfolded state at pH 2.0. Moreover, NaCl also showed a significant ability to induce  $\alpha$ -helix formation. These results suggested that halogen is a key factor in the induction of  $\alpha$ -helix

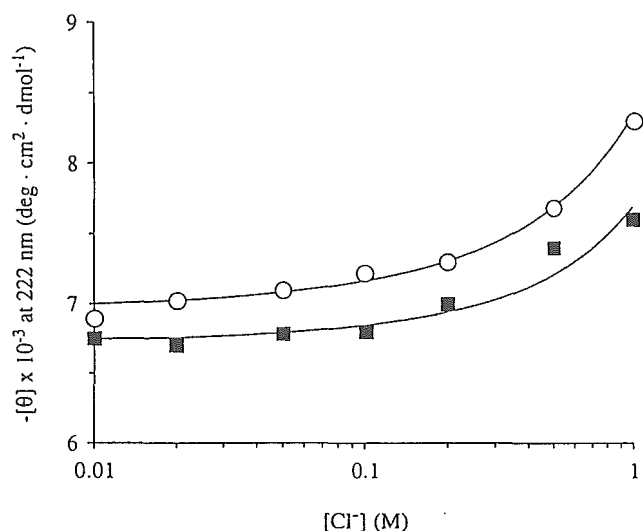


Fig. 3. Effect of halogen ( $\text{Cl}^-$ ) on  $\alpha$ -helix formation in AGP. The  $\alpha$ -helix content of AGP was monitored at pH 2.0 by measuring the ellipticity at 222 nm. Abbreviations are as follows: ○, NaCl; and ■, HCl. The lines were drawn using the MULTI program [28].

formation by halogenols and that negative charge acts cooperatively with hydrophobicity to induce  $\alpha$ -helix structure formation.

#### 4. Discussion

Previously, we reported that AGP interacts with biomembrane models, including liposomes and reverse micelles, and that this interaction is followed by conformational transition from a  $\beta$ -sheet form to an  $\alpha$ -helix form [12,13]. This conformational transition may be linked not only to ligand-binding capacity but also to intracellular biological activity [29–33], but which factors participate in such a conformational transition has not been determined. Generally, it has been thought that following factors are related to the interaction between proteins and lipid membranes: (1) negative charge of the membrane surface; (2) a mild acidic environment on the membrane surface and (3) a hydrophobic membrane interior [17]. However, it is difficult to evaluate the role of these factors in the conformational transition using liposomes and reverse micelles separately.

In the present study, to clarify the mechanism by which AGP undergoes a transition from an  $\alpha$ -helix to a  $\beta$ -sheet structure, we examined the effects of various alcohols. Both Alkanol and diol showed larger values of  $r\text{ASA}(\text{H})$  than  $r\text{ASA}(\text{N})$  and  $r\text{ASA}(\text{P})$  and the abilities of alkanols and diols to induce formation of an  $\alpha$ -helix structure were dependent on the length of their hydrocarbon groups. These  $\text{HE}_{50}$  values correlated well with the  $r\text{ASA}(\text{H})$ . These results indicate that hydrophobic force is a key factor for inducing and stabilizing the  $\alpha$ -helix structure. The effects of diols were weaker than other alcohols because the hydroxyl group weakened the hydrophobicity. On the other hand, the  $\text{HE}_{50}$  values of halogenols correlated significantly with the  $r\text{ASA}(\text{N})$ , implying that the  $\alpha$ -helix formation in the presence of alcohols is due to the cooperative action of hydrophobic and electrostatic forces. The strong effect of halogenols may be due to their high  $r\text{ASA}(\text{N})$ . This hypothesis

is also supported by the finding that HCl and NaCl induced  $\alpha$ -helix formation in AGP even at pH 2.0. Under acidic conditions, it appears that hydrophobic interaction between the hydrocarbon group of the alcohol and the exposed hydrophobic region of AGP induces formation of the  $\alpha$ -helix structure. Kodicek et al. [27] reported that AGP formed a similar  $\alpha$ -helix structure in the presence of MeOH at high temperature. We also found that the conformational structure of AGP perturbed at high temperature (data not shown). In addition, suppression of the electrostatic repulsion of positive charges in AGP by the negative charge of the halogen might enhance the hydrophobic interaction. Although HCl and NaCl also induced  $\alpha$ -helix formation, the effect was weaker than that of alcohols. These results indicate that there is a slight increase in the extent of  $\alpha$ -helix conformation when AGP interacts with the membrane surface, and then the peptide inserts into the membrane interior, where it forms an  $\alpha$ -helix-rich structure.

In the present study, we found that hydrophobic and electrostatic forces cooperatively promote  $\alpha$ -helix formation in AGP. The biological function of AGP is not clear, but some reports suggest that it has intracellular activities [29–33]. Given that AGP is a plasma protein, the conformational transition to an  $\alpha$ -helix structure may be critical for its activity.

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