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Use of entrapment and high-performance affinity chromatography to compare the binding of drugs and site-specific probes with normal and glycated human serum albumin

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Abstract

Protein entrapment and high-performance affinity chromatography were used with zonal elution to examine the changes in binding that occurred for site-specific probes and various sulfonylurea drugs with normal and glycated forms of human serum albumin (HSA). Samples of this protein in a soluble form were physically entrapped within porous silica particles by using glycogen-capped hydrazide-activated silica; these supports were then placed into 1.0 cm × 2.1 mm inner diameter columns. Initial zonal elution studies were performed using (*R*)-warfarin and l-tryptophan as probes for Sudlow sites I and II (i.e., the major drug binding sites of HSA), giving quantitative measures of binding affinities in good agreement with literature values. It was also found for solutes with multisite binding to the same proteins, such as many sulfonylurea drugs, that this method could be used to estimate the global affinity of the solute for the entrapped protein. This entrapment and zonal approach provided retention information with precisions of ±0.1–3.3% (± one standard deviation) and elution within 0.50–3.00 min for solutes with binding affinities of 1×10^4 – 3×10^5 M⁻¹. Each entrapped-protein column was used for many binding studies, which decreased the cost and amount of protein needed per injection (e.g., the equivalent of only 125–145 pmol of immobilized HSA or glycated HSA per injection over 60 sample application cycles). This method can be adapted for use with other proteins and solutes and should be valuable in high-throughput screening or quantitative studies of drug–protein binding or related biointeractions.

Keywords: entrapment, high-performance affinity, chromatography, human serum albumin, glycation, drug-protein binding

Introduction

Diabetes is currently the seventh leading cause of death in the USA. This disease is characterized by an elevated level of glucose in blood and exists in two main forms: type I (juvenile or insulin-dependent) diabetes and type II (non-insulin-dependent or adult onset diabetes) [1]. One side effect of this disease is the glycation, or non-enzymatic addition, of reducing sugars such as glucose to amine groups on proteins [2–4]. One protein in blood that is affected by glycation is human serum albumin (HSA) [2–8]. HSA is not only the most abundant protein in human plasma, but it also serves as an important carrier protein for many drugs, fatty acids, and other solutes [9–12]. This protein has two major binding sites for drugs, which are commonly referred to as Sudlow sites I and II [10–12]. It is known Sudlow sites I and II, or residues near these sites, may be altered as a result of glycation [2–8, 13–15]. It has also been determined that this process could potentially alter the binding of some solutes with HSA [2–4, 16], including several sulfonylurea drugs used to treat type II diabetes [17–21].

The purpose of this study is to examine a new combination of tools for quickly examining the overall binding of HSA to drugs and solutes and to determine how such binding might be affected by glycation. This approach is based on the use of entrapped proteins and high-performance affinity chromatography (HPAC). HPAC is a separation method in which a biologically related binding agent is used as the stationary phase on a high-performance liquid chromatography (HPLC) support such as porous silica [22–25]. This approach has been shown in many past reports to be a useful method for studying and characterizing the binding of solutes with biological agents [22, 26], including the interactions of drugs and other solutes with proteins such as HSA [23–25]. Advantages of HPAC include its speed, precision, ease of automation, and ability to reuse the same protein preparation for many experiments [23, 25].

In most studies with HPAC, proteins and other binding agents are covalently immobilized onto the surface of a support. However, this immobilization process can lead to a decrease or loss of activity for the binding agent owing to multisite attachment, improper orientation, or steric effects [26, 27]. Recently, a noncovalent immobilization technique has been described for the physical entrapment of soluble proteins such as HSA within the pores or near the surface of silica supports [28]. This approach, as illustrated in Figure 1, makes use of large particles of mildly oxidized glycogen that combine with hydrazide-activated silica and entrap soluble proteins on the support while still allowing access of small solutes and binding targets to these proteins. This method has been shown in prior work with frontal analysis and a site-selective probe for Sudlow site I, i.e., (S)-warfarin, to produce immobilized normal HSA

with essentially full activity and binding constants that are in good agreement with values that are seen for soluble HSA [28].

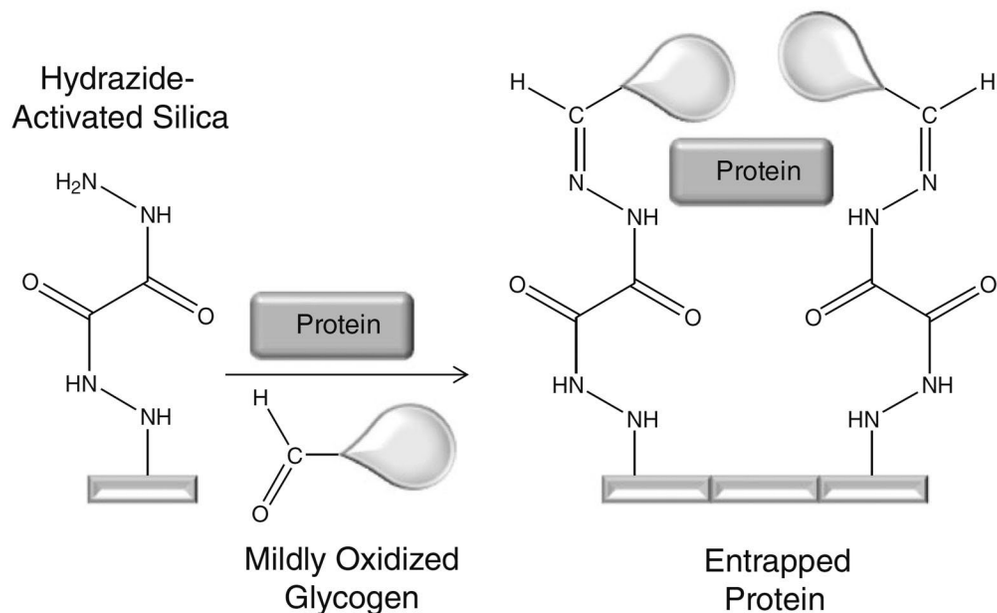
This study will examine the combined use of HPAC columns and protein entrapment to compare the overall binding of representative drugs and probe compounds with normal HSA and glycated HSA. The emphasis will be on the use of such columns in zonal elution studies [23–25] to provide a relatively fast means for examining the binding of the entrapped proteins to injected targets. This method will first be evaluated by using it to compare the binding of normal HSA versus glycated HSA to (*R*)-warfarin and L-tryptophan (i.e., site-selective probes for Sudlow sites I and II, respectively) [10, 29, 30]. This approach will then be extended to studies examining the overall binding of several sulfonylurea drugs to the same protein samples. These results will be compared with those obtained by other methods for similar systems and should provide valuable data regarding the use of protein entrapment and HPAC for comparing the binding properties of normal and modified proteins. The same data should help illustrate the potential for this combined approach in high-throughput screening of drug–protein interactions.

Experimental

Chemicals

p-Periodic acid (periodic acid reagent, or H_5IO_6), glycogen (from bovine liver), HSA (Cohn fraction V, 99% globulin free, 99% fatty acid free), glycated HSA (95% lyophilized, lot no. 115K6108, prepared in vitro and con-

Figure 1. Entrapment of a protein by reaction of hydrazide-activated silica with mildly oxidized glycogen. Details on this approach are provided in the text.



taining 1.8 mol hexose per mole of HSA), (*R*)-warfarin, *L*-tryptophan, acetohexamide, gliclazide, and tolbutamide were purchased from Sigma-Aldrich (St. Louis, MO, USA). Nucleosil Si-300 silica (300-Å pore size, 7- μ m particle diameter) was purchased from Macherey-Nagel (Düren, Germany). Reagents for the micro bicinchoninic acid (BCA) protein assay were from Pierce (Rockford, IL, USA). Other chemicals were of the purest grades available. Econo-Pac 10 DG disposable 10-mL desalting columns were obtained from Bio-Rad Laboratories (Hercules, CA, USA). All solutions were prepared using water from a NANOpure purification system (Barnstead, Dubuque, IA, USA) and filtered through 0.20- μ m GNWP nylon membranes from Millipore (Billerica, MA, USA).

Instrumentation

The chromatographic system consisted of a JASCO (Tokyo, Japan) PU-980i intelligent HPLC isocratic pump, a Rheodyne (Cotati, CA, USA) Advantage PF ten-port valve, and a JASCO UV-975 UV/vis detector. The detection wavelengths were as follows: 304 nm, (*R*)-warfarin; 280 nm, *L*-tryptophan; 205 nm, sodium nitrate; 248 nm, acetohexamide; 226 nm, gliclazide; and 250 nm, tolbutamide. Data were collected using an interface and software from National Instruments (Austin, TX, USA). The temperature of the columns and mobile phases was controlled using a PolyScience (Buffalo Grove, IL, USA) circulating water bath and a water jacket from Alltech (Deerfield, IL, USA). All columns were packed using an HPLC column slurry packer from Alltech. The chromatographic data were analyzed using Peakfit 4.12 (Jandel Scientific Software, San Rafael, CA, USA).

Methods

The overall scheme used for protein entrapment was based on a procedure reported in [28] using normal HSA as a model; the same method was found to entrap glycated HSA in preliminary studies in this prior report. The oxidation of glycogen was performed in pH 5.0 buffer containing 20 mM sodium acetate and 15 mM sodium chloride. A 4.0 mL solution was prepared by dissolving 135 mg of periodic acid and 17 mg of glycogen in this buffer. The resulting solution was covered in aluminum foil, because of the light sensitivity of periodic acid, and was shaken at room temperature for 12–24 h. After oxidation, the glycogen was separated from the remaining periodic acid by using an Econo-Pac 10DG disposable desalting column and pH 5.0, 0.10 M potassium phosphate buffer as the mobile phase. The resulting solution of oxidized glycogen had a concentration of approximately 4.2 mg/mL, a final volume of around 4.0 mL, and approximately 0.5% oxidation of the glucose units in glycogen [28]. This solution was stored in pH 5.0, 0.10 M phosphate buffer at 4 °C.

The normal HSA and glycated HSA supports were prepared under the same immobilization conditions. A 50 mg/mL stock solution of each protein was first prepared in pH 5.0, 0.10 M potassium phosphate buffer. The starting support, Nucleosil Si-300, was converted into a hydrazide-activated form according to a previous procedure [31]. To create the entrapped-protein support, 80 mg of hydrazide-activated silica was combined with 160 μ L of the protein solution. This mixture was sonicated under a vacuum for 15 min to remove any air trapped within the support's pores. A 380- μ L portion of the oxidized glycogen solution was then added, and this mixture was shaken at room temperature for 12 h. During the final hour of the reaction, 200 μ L of a 2 mg/mL oxalic dihydrazide solution in pH 5.0, 0.10 M phosphate buffer was added to the reaction mixture to combine with any remaining aldehyde groups on the glycogen. After immobilization, the support was washed several times with pH 7.4, 0.067 M potassium phosphate buffer. A control support was prepared in the same manner by using only pH 5.0, 0.10 M phosphate buffer in place of the protein solution during the entrapment step [28]. The protein content of each final support was determined in triplicate by using a micro BCA assay, with soluble HSA or glycated HSA being used as the standard and the control support being used as a blank.

All supports were downward slurry-packed into separate 1.0 cm \times 2.1 mm diameter stainless steel columns at 27.6 MPa (4,000 psi) using pH 7.4, 0.067 M potassium phosphate buffer as the packing solution. The columns were stored at 4 °C in this packing solution. The sample solutions for the chromatographic studies were prepared in pH 7.4, 0.067 M potassium phosphate buffer, which was also used as the mobile phase. Stock solutions with a concentration of 30 μ M (*R*)-warfarin, or 100 μ M *L*-tryptophan, acetohexamide, gliclazide, or tolbutamide were prepared in the same pH 7.4 buffer and diluted with this pH 7.4 buffer to give 10 μ M working samples for these analytes; these sample concentrations were found to be sufficiently low compared with the column's overall binding capacity to provide linear elution conditions during the retention factor measurements.

A 20- μ L volume of each 10 μ M sample solution was injected in triplicate onto the entrapped-protein columns and the control column at 37 °C. Flow rates of 0.1–1.0 mL/min were used to examine the binding of each analyte to the columns. The void time of each column was determined by making triplicate 20- μ L injections of 25 μ M sodium nitrate in the presence of pH 7.4, 0.067 M potassium phosphate buffer at all the flow rates tested. Sodium nitrate has routinely been used in the past as a void volume marker for columns containing covalently immobilized HSA and glycated HSA, as well as with columns containing hydrazide-activated silica, and has been found to give good agreement with calcu-

lated void volumes based on the known packing densities and porosities of the corresponding supports [19–21, 23, 25, 28–31]. The void time of the system was determined by repeating the injections of sodium nitrate but using a zero-volume union in place of the column. A fit to an exponentially modified Gaussian curve was used to determine the central moment of each peak for the calculation of retention factors.

Results and discussion

General properties of entrapped HSA and glycated HSA columns

The entrapment supports were found through a BCA protein assay to contain $37 (\pm 1)$ mg HSA per gram of silica, or $32 (\pm 2)$ mg glycated HSA per gram of silica, where the values listed in parentheses throughout this article represent a range of \pm one standard deviation. These supports were packed into $1.0 \text{ cm} \times 2.1 \text{ mm}$ inner diameter columns, resulting in a total protein content in terms of moles (m_{Ltotal}) of 8.7 nmol or 7.5 nmol (i.e., 500 μg or 580 μg) for the HSA and glycated HSA (note these values were estimated by using a packing density of 0.45 g/cm^3 for the support material, as reported by the manufacturer, and a molar mass for HSA and glycated HSA of 66.5 kDa). The columns were found to be stable and give reproducible retention factors over the course of more than 60 sample injection cycles and approximately 4 months of operation. Because each entrapped protein sample was used for many binding studies, this resulted in a decrease in the cost and overall amount of protein needed per injection. For instance, the equivalent of only 125–145 pmol (8.3–9.7 μg) of immobilized HSA or glycated HSA was used per injection when employed over the course of 60 application cycles.

These columns were initially evaluated by using zonal elution as a rapid means for obtaining estimates of binding affinities for injected probe compounds or solutes with the entrapped proteins. Figure 2 provides some typical chromatograms that were obtained for (*R*)-warfarin and *L*-tryptophan on these columns. When compared with the control column containing no protein, both the normal HSA column and the glycated HSA column showed markedly increased retention for both probe compounds. (*R*)-warfarin, which has an association equilibrium constant of roughly 2×10^5 – $3 \times 10^5 \text{ M}^{-1}$ for normal HSA [29], was eluted within 5–6 min at 0.5 mL/min from both types of entrapped-protein columns and within 2.5–3 min from the same columns at 1.0 mL/min. *L*-tryptophan, which has a tenfold lower association equilibrium constant of approximately 1×10^4 – $2 \times 10^4 \text{ M}^{-1}$ for normal HSA [30], was eluted in only 1.0–1.5 min at 0.5 mL/min from both types of entrapped-protein columns and within 0.50–0.75 min at 1.0 mL/min.

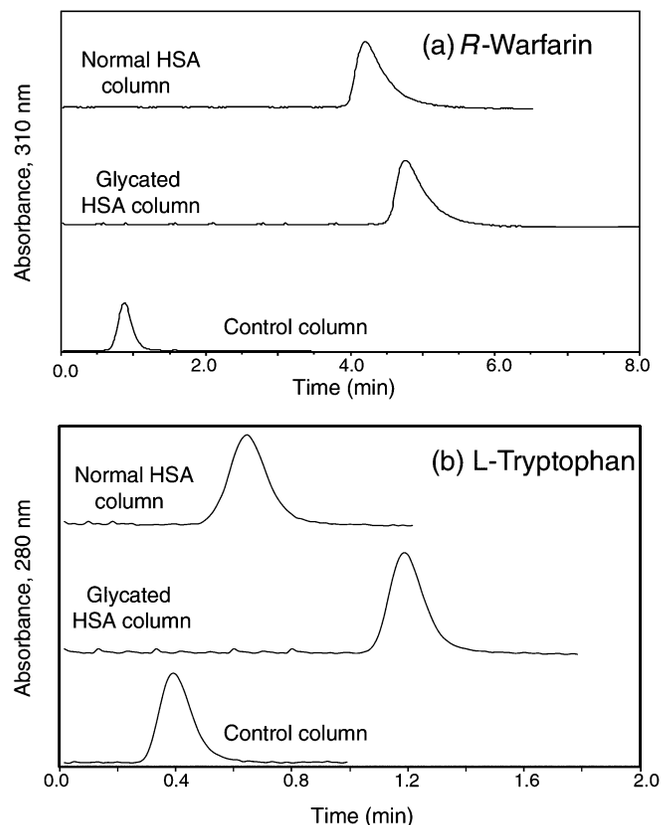


Figure 2. Chromatograms obtained for the injection of a (*R*)-warfarin or b *L*-tryptophan as a site-selective probe onto a control column containing no entrapped protein or columns containing entrapped normal human serum albumin (HSA) or glycated HSA. These results were obtained at 0.50 mL/min. Other conditions are given in the text.

Reproducible retention factors and stable binding characteristics were obtained on these columns for triplicate injections obtained over five flow rates ranging from 0.1 to 1.0 mL/min, with retention factors showing only random variations of $\pm 4\%$ overall for (*R*)-warfarin and $\pm 11\%$ for *L*-tryptophan. The precisions of the retention factors measured at individual flow rates for the same solutes ranged from ± 0.4 to $\pm 1.1\%$ and from ± 0.6 to $\pm 3.3\%$, respectively, with the precision increasing as slower flow rates and longer residence times were used.

Binding of (*R*)-warfarin with entrapped HSA and glycated HSA

The next part of this study used HPLC and the columns containing entrapped samples of normal HSA or glycated HSA to evaluate the binding of these columns to (*R*)-warfarin as a probe for Sudlow site I (i.e., the warfarin–azapropazone site) [10, 29, 32]. This drug (see Figure S1 for the structure) has been previously shown to bind to both normal and glycated HSA in a site-selective manner. The association equilibrium constants at pH 7.4

and 37 °C for these interactions have been estimated by various methods for normal HSA [2, 29, 30, 32–34] and by frontal analysis for in vitro glycosylated HSA [2], thus providing reference values for this current study for the evaluation of the entrapped-protein columns.

In the zonal elution approach that was used in this study, the specific retention factor (k) was determined for an injected drug or solute on a given protein column after correcting for the effect of any nonspecific interactions with the support on the total retention, e.g., a correction for (*R*)-warfarin which was only 2.5–2.6% of the total retention seen on the entrapped protein columns; this approach was adequate for the current study, but a more thorough analysis of the nonspecific interactions could be obtained in future work through frontal analysis. The use of the specific retention factor provided a relatively fast and easy way of measuring and comparing the level of binding for each injected solute on the entrapped-protein columns. For more detailed studies of binding, the resulting value of k due to the entrapped protein could also be directly related to the global affinity constant (nK_a') of the solute with the immobilized protein, as represented by the general expression in Equation 1 for any group of independent binding sites [23, 25]:

$$\text{Single-site or multisite binding: } k = \frac{(nK_a' m_{Ltotal})}{V_M} \quad (1)$$

where m_{Ltotal} again represents the total number of moles of protein, and V_M is the column void volume [23].

It is important to note in Equation 1 that nK_a' is a number-weighted sum of the association equilibrium constants for all binding sites of the solute on the immobilized protein, as indicated by Equation 2 for a system with multiple, independent binding sites [23, 25]:

$$nK_a' = \sum (n_i K_{ai}) \quad (2)$$

In Equation 2, n_i represents the relative number of moles of binding site i for a given solute per mole of protein, and K_{ai} is the association equilibrium constant for the same site and solute. If only a single type of binding site for a drug or solute is present and essentially all of the immobilized protein is present in an active form (i.e., as could occur for an entrapped but noncovalently immobilized protein), Equation 1 can be rewritten as Equation 3:

$$\text{Single-site binding: } k = \frac{(K_a m_{Ltotal})}{V_M} \quad (3)$$

In this specific case, $n = 1$ and nK_a' is now given by K_a , the association equilibrium constant for the solute at its binding site on the protein.

Table 1 summarizes the specific retention factors that were measured for (*R*)-warfarin on the columns containing entrapped normal HSA and glycosylated HSA. To correct for differences in the protein content of these supports and to allow a direct comparison of their retention values, the specific retention factors were normalized by dividing them by each support's total protein content [35, 36]. When these normalized retention

Table 1. Retention factors and global affinity constants measured for (*R*)-warfarin and L-tryptophan on columns containing entrapped normal human serum albumin (HSA) or glycosylated HSA

Type of HSA and solute	Specific retention factor, k^a	Normalized retention factor, $k/(\text{protein content})^b$	Global affinity, $nK_a' (M^{-1})^c$
Normal HSA			
(<i>R</i>)-warfarin	84.7 (\pm 0.5)	2.29 (\pm 0.06)	$2.9 (\pm 0.1) \times 10^5$
L-tryptophan	5.7 (\pm 0.2)	0.15 (\pm 0.01)	$1.8 (\pm 0.1) \times 10^4$
Glycosylated HSA			
(<i>R</i>)-warfarin	78.5 (\pm 0.8)	2.45 (\pm 0.08)	$3.1 (\pm 0.1) \times 10^5$
L-tryptophan	14.22 (\pm 0.03)	0.44 (\pm 0.01)	$5.5 (\pm 0.2) \times 10^4$

The values in parentheses represent a range of \pm one standard deviation, as based on triplicate measurements made over five flow rates ($n = 15$). As indicated in the text, the same range of flow rates was used to initially evaluate the reproducibility and stability of these columns over a range of operating conditions but gave no significant differences in the measured retention factors.

- The specific retention factor was obtained by taking the difference between the overall retention factor on an entrapped-protein column and the retention factor due to nonspecific interactions measured for the same solute on the control column, with the latter value being $2.13 (\pm 0.05)$ for (*R*)-warfarin and $0.77 (\pm 0.03)$ for L-tryptophan.
- The normalized retention factors were calculated by dividing the specific retention factors by the measured total protein content for each given support, using values of $37 (\pm 1)$ mg HSA per gram or $32 (\pm 2)$ mg glycosylated HSA per gram.
- The value of nK_a' was estimated by using Equation 1 along with the specific retention factor (k) for each solute, the measured void volume (V_M), and the number of moles of protein in the column (m_{Ltotal}). The value used for V_M was 29 μ L and m_{Ltotal} was 8.7 nmol or 7.5 nmol for the normal HSA or glycosylated HSA columns, respectively; details on the determination of V_M and m_{Ltotal} are provided in the text.

factors were compared, no significant difference at the 95% confidence level (i.e., the confidence level used for all comparisons in this study) was seen in the values for (*R*)-warfarin on the columns containing normal HSA and glycosylated HSA. These results indicated the level of glycosylation present in this case did not have an observable effect on the binding of (*R*)-warfarin with HSA at Sudlow site I. These results are consistent with previous data and conclusions obtained when frontal analysis was used to compare covalently immobilized normal HSA with *in vitro* glycosylated HSA that had levels of modification for the glycosylated HSA that were similar or equivalent to those used in the current study [2].

Using the fact that (*R*)-warfarin has only one major binding site on normal HSA and glycosylated HSA, and that most of these entrapped proteins should have been active, we used the retention data to further estimate and compare the association equilibrium constants for this interaction. These results are also provided in Table 1. The K_a of $2.9 (\pm 0.1) \times 10^5$ that was calculated from the zonal elution data for (*R*)-warfarin with normal HSA was in good agreement with previously reported values of 2.1×10^5 – 3.3×10^5 M⁻¹ for this interaction under the same temperature conditions [2, 29, 30, 32–34]. The K_a of $3.1 (\pm 0.1) \times 10^5$ M⁻¹ obtained for (*R*)-warfarin with the glycosylated HSA was also consistent with prior estimates of $2.3 (\pm 0.3) \times 10^5$ –to $2.7 (\pm 0.3) \times 10^5$ M⁻¹ that have been made by frontal analysis for similar preparations of this protein [2]. In addition, the calculated K_a values did not show any significant difference between the normal HSA and glycosylated HSA columns, as noted in the prior studies [2].

Binding of L-tryptophan with entrapped HSA and glycosylated HSA

The same columns as described in the previous section were examined for their binding to L-tryptophan as a probe for Sudlow site II (i.e., the indole-benzodiazepine site) [10, 30, 37]. Like (*R*)-warfarin, this solute (see Figure S1 for the structure) is known to bind to both normal and glycosylated HSA in a site-selective manner [2, 10, 30]. The association equilibrium constant for the binding of L-tryptophan with normal HSA has also been determined by various techniques at 37 °C [30, 34, 37, 38], and the binding of this solute under the same conditions with *in vitro* glycosylated HSA has been previously examined by frontal analysis [2], again providing values that could be used to evaluate the performance of the entrapped-protein columns.

The results from the zonal elution studies that were obtained for L-tryptophan are included in Table 1. Unlike the data acquired for (*R*)-warfarin, the normalized retention factors that were calculated for L-tryptophan showed a large change when going from normal HSA to glycosylated HSA, even after a correction had been made for differences in the total protein content of the

columns. The normalized retention factor for L-tryptophan on the entrapped glycosylated HSA column was roughly three times higher than that for L-tryptophan on the column containing normal HSA. This difference was significant at the 95% confidence level. The value of nK_a' between these two columns increased by the same factor and was again significant at the 95% confidence level. These results clearly indicated that the glycosylation of HSA did affect the binding of L-tryptophan to Sudlow site II, as was concluded in a previous study [2].

Given the fact that L-tryptophan has only one major binding site on HSA, the global affinity measured for this solute was assumed to represent the association equilibrium constant for this interaction. The average K_a of $1.8 (\pm 0.1) \times 10^4$ M⁻¹ that was estimated by this approach for L-tryptophan with the entrapped normal HSA was in good agreement with previous values of 1.1×10^4 – 2.4×10^4 M⁻¹ that have been determined by other methods for this interaction at 37 °C [2, 30, 34, 37–39]. The K_a of $5.5 (\pm 0.2) \times 10^4$ M⁻¹ measured for L-tryptophan with the entrapped glycosylated HSA also fit within the range of 5.2×10^4 – 6.4×10^4 M⁻¹ that has previously been measured by frontal analysis at 37 °C with *in vitro* glycosylated HSA preparations having similar levels of modification [2].

Binding of sulfonylurea drugs to entrapped HSA and glycosylated HSA

The final stage of this work used the entrapped HSA and glycosylated HSA columns to screen and compare the binding of various sulfonylurea drugs to these proteins. The sulfonylurea drugs considered in this study were acetohexamide, gliclazide, and tolbutamide (see the structures in Figure 3). Unlike (*R*)-warfarin and L-tryptophan, these sulfonylurea drugs are known to bind to both Sudlow site I and Sudlow site II of normal HSA and glycosylated HSA, as well having a set of weaker interaction sites with these proteins [2, 17, 19–21]. Thus, for these drugs the use of retention factors to describe protein interactions would be expected to reflect the contributions of multiple binding regions, as indicated by Equations 1 and 2, rather than a single binding site, as described by Equation 3.

Some typical chromatograms obtained in these studies are shown in Figure 4. As noted earlier with (*R*)-warfarin and L-tryptophan, each of the sulfonylurea drugs exhibited a large difference in retention between the entrapped-protein columns and the control column. These drugs were eluted within 1.1–5.0 min at 0.5 mL/min and within 0.6–2.5 min at 1.0 mL/min from the columns containing the entrapped proteins. The measured retention factors had good reproducibility over flow rates ranging from at least 0.1 to 1.0 mL/min, with values varying by only $\pm 3\%$, $\pm 6\%$, or $\pm 8\%$ for acetohexamide, gliclazide, and tolbutamide, respectively. The retention factors measured at individual flow rates for the same drugs ranged from ± 0.1 to $\pm 0.6\%$, from ± 0.1 to $\pm 1.7\%$,

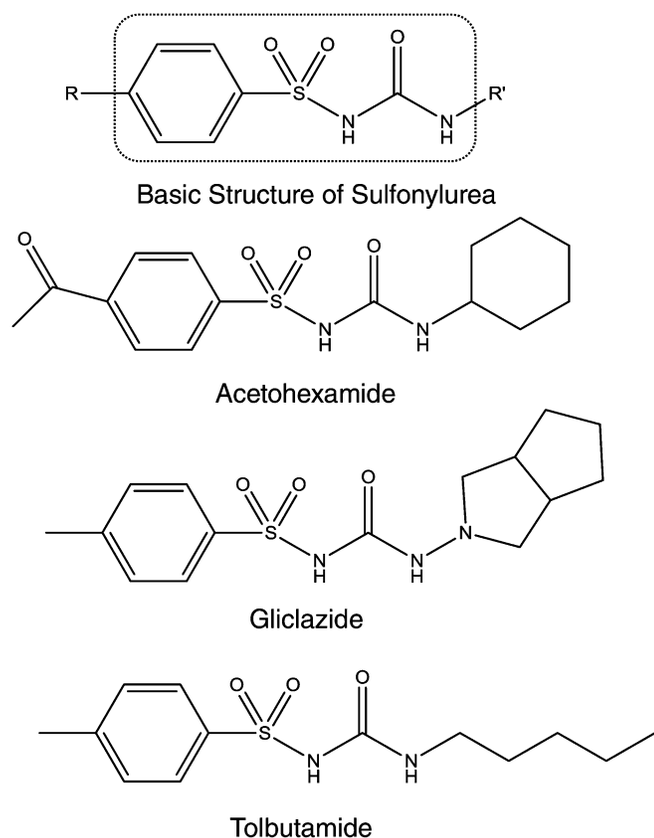


Figure 3. Basic structure of a sulfonylurea drug and the structures of the specific sulfonylurea drugs that were examined in this study.

and from ± 0.2 to $\pm 0.7\%$, respectively, and tended to improve as the flow rate was decreased and the residence time was increased.

Table 2 provides the specific retention factors that were measured for each of the sulfonylurea drugs on the normal HSA and glycosylated HSA columns. Normalized retention factors that corrected for differences in the protein content of these columns were also calculated (see the footnote in Table 2). These sulfonylurea drugs showed an increase of 55–77% in the normalized retention factors for the glycosylated HSA column compared with the normal HSA column. This increase in the normalized retention factor was in agreement with previously reported trends noted for these sulfonylureas when covalently immobilized samples of normal HSA and a preparation of *in vitro* glycosylated HSA similar to that used in the current study were used [18]. These trends also agreed with those seen in separate ultrafiltration studies using physiological levels of normal HSA and *in vitro* glycosylated HSA [20, 21, 40].

These retention data were further used to estimate the global affinities of each sulfonylurea drug with the entrapped samples of HSA and glycosylated HSA, as given in Table 2. In this case, it was known from prior studies that all of these drugs had two sets of binding sites on HSA

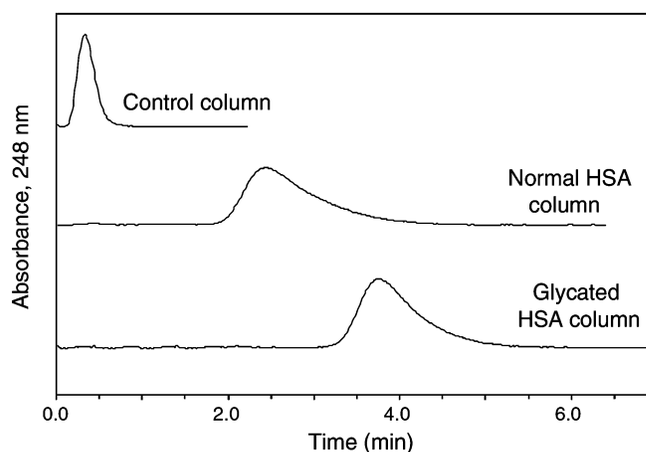


Figure 4. Chromatograms obtained for acetohexamide on a control column containing no entrapped protein or on columns containing entrapped normal HSA or glycosylated HSA. These results were obtained at 0.50 mL/min. Other conditions are given in the text.

(i.e., moderate-to-high affinity sites and weak binding regions), with most of this binding being attributed to one or two moderate-to-high affinity sites [17, 19–21]. Thus, the global affinities shown in Table 2 would be predicted from Equation 2 to represent a number-weighted sum of the association equilibrium constants for these sites; note that zonal elution competition studies with probes such as (*R*)-warfarin and *l*-tryptophan could also be employed to obtain information on the affinities at specific binding sites [17, 19–21, 23]. The relative trends shown in Table 2 for these results were the same as noted for the normalized retention factors, as would be expected from Equation 1 because these two sets of values were directly proportional to each other.

The global affinities given in Table 2 compared well with those predicted by Eqs. 1 and 2 when calculating such results using previous binding capacities and association equilibrium constants obtained by frontal analysis, or a combination of these results with site-selective binding constants that were obtained by zonal elution at the higher-affinity sites for sulfonylurea drugs with covalently immobilized samples of normal HSA and *in vitro* glycosylated HSA [17, 19–21]. In each case, most of the calculated global affinity (i.e., 92–99% for acetohexamide, 82–92% for gliclazide, and 92–98% for tolbutamide) was due to the contributions of the moderate-to-high binding regions for these drugs with HSA (i.e., Sudlow sites I and II).

Because the zonal elution approach used in this study employed direct measurements of retention to estimate global affinities, the precision of this approach was much better than when combining association equilibrium constants and binding capacity results to calculate global affinities. For instance, the global affinities obtained by using data from prior frontal analysis and zonal elution studies [17, 19–21] had typical final preci-

sions of ± 20 -50%, as based on error propagation. In contrast, the global affinities in Table 2 that were measured directly by zonal elution for the entrapped-protein columns had precisions of ± 3 -10%.

Another potential advantage of using entrapped proteins to measure global affinities was that little or no loss of activity should have occurred during the immobilization process [26, 28]. This feature is essential if zonal elution data are to be used directly for determining binding constants [23, 25] and has made this approach generally impractical for use with covalent immobilization techniques because the loss of protein activity is common during such coupling methods [23, 25, 26]. For instance, it was estimated from previous frontal analysis data [17, 19-21] that a combined loss in activity of 35-60% for acetohexamide, 45-59% for tolbutamide, and 75-80% for gliclazide occurred at Sudlow sites I and II when normal HSA and glycosylated HSA that had been covalently immobilized by the Schiff base method were used [17, 19-21]. On the basis of prior work with (*R*)-warfarin and l-tryptophan, it is expected that a similar or even larger decrease in the activity for HSA might occur when other covalent immobilization methods are used [35, 36].

Concluding remarks

This report examined the use of entrapment and HPLC-based zonal elution studies to examine the changes in binding and overall affinity that occurred for site-specific probes and various sulfonylurea drugs with normal HSA versus glycosylated HSA. The columns created by entrapment gave retention factors with precisions of ± 0.1 -3.3% over the course of more than 4

months of operation. The compounds retained on the entrapped-protein columns had elution times of less than 2.5-3.0 min at 1.0 mL/min for solutes with affinities as high as 2×10^5 - 3×10^5 M⁻¹, e.g., as represented by (*R*)-warfarin on the normal HSA and glycosylated HSA columns, and less than 0.50-0.75 min for solutes with affinities down to 1×10^4 - 2×10^4 M⁻¹, e.g., as seen for l-tryptophan on the normal HSA column. The ability to use these columns over many experiments produced a system that required the equivalent of less than 125-145 pmol of immobilized protein per injection over 60 application cycles. All of these characteristics are attractive for using such an approach in biointeraction studies and in the high-throughput screening of drug-protein binding.

The ability of this method to provide a rapid, quantitative measure of binding affinity was also examined. In the case of solutes having a single major binding site on HSA, this approach could be used to determine the association equilibrium constant for the injected solutes with the entrapped proteins. For instance, the binding constants determined by this approach for (*R*)-warfarin at Sudlow site I and for l-tryptophan at Sudlow site II were in good agreement with results reported with other techniques for normal HSA or glycosylated HSA [2, 29, 30, 32-34, 37-39]. For solutes with multisite binding, this method could be used to estimate the global affinity of the solute for the entrapped protein. This was illustrated by using this approach to examine and compare the global affinities of various sulfonylurea drugs with normal HSA and glycosylated HSA. The values obtained again showed good agreement with trends and results that were predicted by other techniques and with use of similar protein preparations [17-21, 40].

Table 2. Retention factors and global affinity constants measured for sulfonylurea drugs on columns containing entrapped normal HSA or glycosylated HSA

Type of HSA and drug	Specific retention factor (k)	Measured global affinity, nK_a' (M ⁻¹) ^a	Estimated global affinity, nK_a' (M ⁻¹) ^b
Normal HSA			
Acetohexamide	46.5 (± 0.1)	$1.6 (\pm 0.1) \times 10^5$	1.6×10^5 - 1.7×10^5
Gliclazide	17.1 (± 0.2)	$5.8 (\pm 0.2) \times 10^4$	5.2×10^4 - 7.9×10^4
Tolbutamide	30.8 (± 0.1)	$1.0 (\pm 0.1) \times 10^5$	1.0×10^5 - 1.1×10^5
Glycosylated HSA			
Acetohexamide	62.7 (± 0.1)	$2.5 (\pm 0.1) \times 10^5$	1.6×10^5 - 1.8×10^5
Gliclazide	23.4 (± 0.1)	$9.1 (\pm 0.3) \times 10^4$	6×10^4 - 12×10^4
Tolbutamide	47.0 (± 0.2)	$1.9 (\pm 0.1) \times 10^5$	1.1×10^5 - 1.5×10^5

The values in parentheses represent a range of \pm one standard deviation, as based on triplicate measurements made over five flow rates ($n = 15$). The specific retention factors and values for nK_a' were determined in the same manner as described in Table 1. In obtaining the specific retention factors, we found the retention factor measured for nonspecific interactions by these drugs on the control column were 0.42 (± 0.04) for acetohexamide, 0.95 (± 0.10) for gliclazide, and 0.39 (± 0.04) for tolbutamide.

a. The nK_a' values measured for the normal HSA column were determined by using the following normalized retention factors: 1.26 (± 0.03) for acetohexamide, 0.46 (± 0.01) for gliclazide, and 0.83 (± 0.02) for tolbutamide. The nK_a' values for the glycosylated HSA column were found by using the following normalized retention factors: 1.96 (± 0.06) for acetohexamide, 0.73 (± 0.02) for gliclazide, and 1.47 (± 0.05) for tolbutamide.

b. These values were calculated by using frontal analysis and zonal elution data from [17, 19-21], as described in the text.

These results indicate the entrapped-protein columns could be used directly with HPAC and zonal elution to estimate and compare binding constants or global affinities for normal and modified proteins such as HSA and glycosylated HSA. Obtaining these quantitative estimates of affinities was made possible by the fact the entrapped protein was immobilized in a form that retained all or most of its original activity [28]. This feature, in turn, allowed the value of K_a or nK_a' to be determined from the measured retention factors of solutes if the protein content of the support was also known. This direct approach based on zonal elution was not possible in the past when covalent immobilization methods were used because of the loss of activity that occurs for some of the immobilized protein and the possible presence of different degrees of inactivation at separate regions on an immobilized protein [26]. Instead, retention measurements have been used in the past to make only relative comparisons of activity [18, 23, 35, 36], or more complicated and time-consuming methods based on frontal analysis or zonal elution plus competition experiments have been required to obtain binding constants that can be measured in a manner independent of the actual amount of active binding sites [2, 17, 19–21, 23–25].

The combined speed and ease of using zonal elution with the entrapped forms of proteins should make this approach attractive for clinical and pharmaceutical applications that may benefit from high-throughput screening or quantitative studies of binding between proteins and small soluble targets. For instance, the shifts in normalized retention factors and affinities that were measured when entrapped normal HSA and glycosylated HSA were used are of potential interest in examination of how glycosylation may affect the protein binding of sulfonamide drugs in blood [17–21]. Such information, in turn, may help clinicians to avoid issues with using either too low or too high of a dose for these drugs when treating diabetes, which could lead to the undesired side effects of inadequate glucose control or hypoglycemia, respectively [19, 41]. The same general methods as used in this study could be adapted for other proteins and solutes or for comparing different protein populations and biointeraction systems of interest in clinical, pharmaceutical, or biomedical research.

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Analytical and Bioanalytical Chemistry

Electronic Supplementary Material

Use of entrapment and high-performance affinity chromatography to compare the binding of drugs and site-specific probes with normal and glycated human serum albumin

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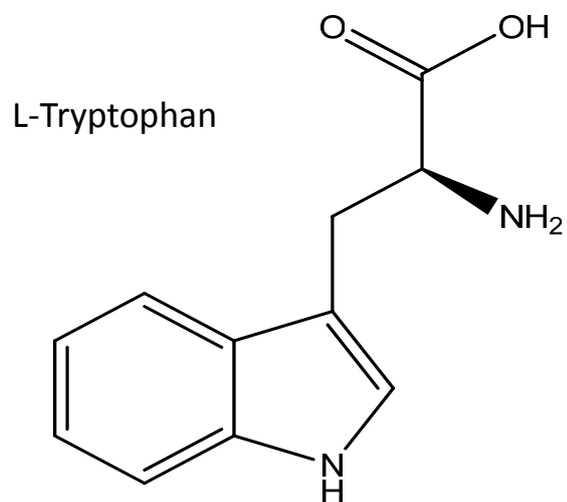
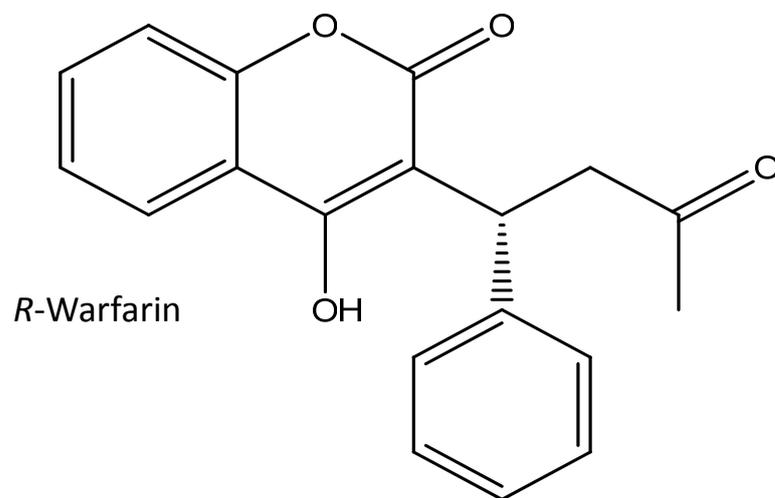


Figure S1. Structures of *R*-warfarin and L-tryptophan