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Native mass spectrometry analysis of conjugated HSA and BSA complexes with various flavonoids[†]

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Mass spectrometry was used to study the binding interaction between serum albumin proteins (BSA and HSA) and flavone dyes, which is known to induce large fluorescence signals for protein detection. By electrospray ionization mass spectrometry (ESI-MS), multiple charged species/states could be produced in ammonium acetate buffer, while preserving the native structures of the proteins. Subsequent introduction of a flavone dye into the buffered solution resulted in an immediate interaction, forming the respective protein–dye conjugates associated by non-covalent interactions. Formation of protein–dye conjugates and molecular mass of the protein species. The resulting data pointed out that the protein–flavone dye maintained a 1:1 ratio in the conjugate, although multiple binding sites for drug molecules are present in albumin proteins.

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Introduction

Mass spectrometry (MS) is a powerful tool that can assist in studying proteins and protein complexes *via* proteomics, which involves the characterization of proteins expressed by cells or organisms.^{1,2} Post-translational modifications such as phosphorylation, ubiquitination, and more can be monitored *via* MS.³⁻¹⁰ Additionally, protein complexes can be confirmed based on the specific residue interactions detected *via* tandem mass spectrometry (MS/MS) fragmentation, a procedure referred to as top down sequencing.^{11–13} Further analysis of complexes can be monitored by using the ion mobility where drift times and fragmentation can be compared, thus providing a two-fold analysis of the complex.^{14,15} These MS techniques can be used to further examine the native structure of a protein, which was first demonstrated by JB Fenn in 1989.¹⁶

In order to achieve native MS, the protein must be prepared in a volatile solution, typically ammonium acetate sulution.¹⁷ It should be noted that the ammonium acetate solution can mimic the protein solvation under physiological conditions, thereby avoiding premature degradation and preserving the quaternary structure of the protein. Such study of biomolecules/biopolymers in their native conformations can yield useful information, including tertiary and quaternary structures, complex stoichiometry, and ligand attachment site(s).^{18–29} As an important class of natural compounds, flavonoids can readily interact with proteins such as albumins, giving strong fluorescent emissions. For example, a study by Pang *et al.* has shown that such protein binding could induce a large fluorescence signal from a flavonoid, which can be used for wash-free SDS-PAGE.²⁰ It has been speculated that the observed interaction is primarily driven by entering a hydrophobic pocket of BSA. However, the structure of albumin is known to have different domains that are associated with multiple binding sites for drug molecules,³⁰ as illustrated in Fig. 1. Site I (or the warfarin site) prefers to bind large heterocyclic



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and negatively charged compounds, while site II is preferred by small aromatic carboxylic acids.^{31,32} Molecular docking studies suggest that the flavonoid 1 likely binds to the site I in HSA.³³ By using pheloretin (a chalcone flavonoid present in apples), Bellocco et al. studied the interaction and binding of flavonoids to HSA.³⁴ Their fluorescence displacement using warfarin (a site selective drug for site I) suggests that the flavonoid is likely bound to the site I (a hydrophobic pocket), as well as that interaction of flavonoids with HSA could modify the protein's conformation.³⁴ A study by Bojić et al. further revealed that flavonoids may not bind at the location where warfarin has high affinity, but rather to different regions within the drug-binding site I.35 Although drug displacement experiments provide information about the dye's binding location, they do not give reliable indication about the protein-ligand ratio, because incomplete displacement often occurs. Herein, we demonstrate that mass spectrometry could be used to examine the interactions between flavonoids and both human and bovine serum albumin (i.e. HSA and BSA, respectively), which could shed some additional light on the protein-ligand ratio in the interactions.

Experimental section

Reagents

Bovine serum albumin (>98.0% purity) and human serum albumin (>99.9% purity) were purchased from Sigma-Aldrich (St Louis, MO); water (optima grade), acetonitrile (optima grade), ammonium acetate (>97% purity), and acetic acid (>99.7% purity) were purchased from Fisher Scientific (Pittsburgh, PA). All commercial reagents were used as received without further purification. Flavone dyes **1–3** were synthesized using the procedures reported previously from one of our laboratories (Scheme 1).^{20,36}

BSA and HSA complexation

BSA and HSA solutions were prepared at 0.2 mg mL⁻¹ in a 20 mM ammonium acetate buffer. Stock solutions of the flavone dyes (10 mM) were prepared in acetonitrile. The flavone dye was added to the BSA or HSA solution at a molar ratio of 3:1. The resulting solution was left at room temperature for 5 minutes to allow for conjugation. Denatured solu-



Scheme 1 Flavonoid structures 1–3 with different substituents.

tions of BSA and HSA were prepared by adjusting the pH of the BSA or HSA solution to below 4.0. The denatured complex solutions were prepared by conjugating the flavone dye and BSA as above and then adjusting the pH to below 4.0.

ESI-MS experiments

ESI experiments were carried out on a Waters Synapt HDMS quadrupole/time-of-flight (Q/ToF) mass spectrometer (Waters, Milford, MA) equipped with an ESI source. The prepared solutions (see above) were injected directly into the ESI source using a syringe pump at a flow rate of 75 μ L min⁻¹. Instrument parameters for MS analysis were set as follows: capillary voltage, 3.00 kV; sample cone voltage, 45 V; extraction cone voltage, 8.0 V; desolvation gas (nitrogen) flow, 650 L h⁻¹; trap collision energy (CE), 6.0 eV; transfer CE, 4.0 eV; argon flow through trap and transfer cells, 1.5 mL min⁻¹; source temperature, 40 °C; desolvation temperature, 125 °C. MassLynx (version 4.1) program was used for data analysis.

Results and discussion

Flavonoid compounds 1-3 (Scheme 1) were synthesized and their interactions with either HSA or BSA were examined using UV-vis and fluorescence spectroscopy and native mass spectrometry (MS). Previously, it has been reported that flavonoids can selectively bind to BSA over other proteins in aqueous media, which was accompanied with a strong fluorescence turn on.²⁰ The flavonoid-albumin interactions between HSA and flavonoids were reexamined in ammonium acetate buffer, as a volatile buffer is necessary for preserving the native structure of a protein. As shown in Fig. 2, addition of HSA induced a large fluorescence from the flavonoid 1. In addition, the emission also exhibited significant bathochromic shifts as the solvent polarity increased (ESI Fig. S1[†]), along with a sharp decrease in emission intensity. The observed large fluorescence response indicated that the flavonoid-albumin interaction was preserved in ammonium acetate buffer, which enabled the MS study.



Fig. 2 Fluorescence response of flavonoid 1 (20 μ M) in 20 mM ammonium acetate buffer solution, upon addition of HSA.



Fig. 3 ESI-MS spectrum of native HSA. lons with +15-+18 charges are present, with +17 being the predominant charge state. This charge distribution is indicative of HSA in its native form.

Initial MS investigation was carried out by acquiring the native spectra of both BSA and HSA, which were prepared in 20 mM ammonium acetate in order to preserve the native

Table 1	Experimental vs.	calculated m/2	z values for	native HSA
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HSA native				
Charge state	Measured <i>m</i> / <i>z</i>	Calculated <i>m</i> / <i>z</i>	% error	[M + xK + yH]
15	4464	4463	0.02	$13K^{+} + 2H^{+}$
16	4185	4187	0.05	$14K^{+} + 2H^{+}$
17	3939	3940	0.03	$14K^{+} + 3H^{+}$
18	3718	3722	0.11	$13K^{+} + 4H^{+}$

state. The ESI-MS spectrum of HSA contained ions with +15 to +18 charges (Fig. 3 and Table 1). Based on the MassLynx software of the mass spectrometer, the observed m/z values corresponded to native HSA ionized by a combination of K⁺ and H⁺ cations (Table 1). Among the charge distribution from the native HSA (*i.e.* ions with +15 to +18 charges), the ion with the +17 charge was the most predominant species (Fig. 3), which is consistent with what has been reported in the literature.³⁷

Similarly, native BSA produced a spectrum with charged species ranging from +15 to +19, with the +17 ion being the predominant state.

In order to verify that the characteristics of HSA and BSA in Fig. 3 and 4 correspond to their respective native forms, the



Fig. 4 ESI-MS spectrum of native BSA. lons with +15-+19 charges are present, with +17 being the predominant charge state. Such charge distribution indicates that BSA is in its native form.

mass spectra in their denatured forms were also acquired. Our initial effort to obtain denatured BSA was by using sodium dodecyl sulfate (SDS). However, the SDS-denatured BSA gave a weak MS signal intensity, as denaturing requires the use of excessive SDS (*e.g.* a 50:1 ratio for SDS:BSA),²¹ which caused ionization suppression. Therefore, a pH driven denaturation method was adopted in this study, as BSA molecules can change their folding conformations, *i.e.* deviate from their native structure, when the pH is beyond the 4.0–8.0 range.^{38–40} Thus, acetic acid was added to the 20 mM ammonium acetate-BSA solution until the pH was below 4.0, which caused the protein to denature and expose more residues to the solvent. The newly exposed residues allowed the protein to accept additional charges, as compared to the native state, thereby increasing the charge state of the protein. The ESI-MS spectrum of denatured BSA showed ions with +30 to +48 charges (Fig. 5). Such increase in the number of charges is expected for denatured BSA due to more residues being exposed when the protein is unfolded. In other words, the ESI-MS spectrum of denatured BSA contained ions of distinctively lower m/z (*e.g.* m/z1400–2300, Fig. 5), as a consequence of the higher charge states, which are in sharp contrast to those observed from native BSA (Fig. 4).



Fig. 5 (a) The enhanced spectrum of BSA denatured by acetic acid. Charge states ranging from +30 to +48 are present; the higher charge states indicate an unfolded structure as more residues are present to accept charges m/z values for selective peaks of different charge states are shown on the top. (b) Full spectrum of denatured BSA in upper right corner with a box indicating the enhanced region.



Fig. 6 ESI-MS spectra of native BSA (bottom) and BSA bound to the flavone dye 1 (top). Shifts in the *m/z* values for the BSA-flavone peaks can be observed, as well as a shift in the predominant charge state for native uncomplexed BSA vs. native complexed BSA.

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Fig. 7 Expanded view of the m/z 3850–4350 region in the ESI-MS spectra of native BSA (bottom) and BSA bound to the flavone dye 1 (top). A shift in the predominant charge state can be observed from native unbound BSA to complexed BSA. The m/z values at 3958 and 4206 are indicated by arrows in the BSA-flavone spectrum. For comparison, dashed lines at m/z 3976 and 4225 mark the potential positions for the BSA-(flavone)₂ complex.

BSA-flavone dye conjugation

After establishing a signature mass spectrum for native and denatured BSA, a flavone dye was introduced into the BSA solution to probe the BSA–flavone interaction. The BSA solution was allowed to react with flavone dye **1** for various time frames ranging from thirty seconds to thirty minutes. The time interval of five minutes was chosen for analysis, as it produced the most intense signal (this trend was also observed in the fluorescence spectral studies). The sample obtained after the reaction for over thirty minutes indicated that no additional dye molecules were added to BSA after the five minutes time interval. Therefore, the resulting conjugated solution, obtained after mixing BSA and the flavone dye for five minutes, was injected to determine the stoichiometry of the protein–dye complex.

The ESI-MS spectrum of the BSA–flavone solution showed an increase in the m/z values of all charge states, as compared to the native BSA, indicating that complexation occurred between BSA and flavone (Fig. 6). In addition to the increase in the m/z values, the predominant charge state from +17 for native BSA was changed to +16 for the BSA complexed to the flavone dye (Fig. 7), corroborating the change in the elemental composition after conjugation. As shown in Table 2, BSA was ionized by K⁺ incorporation when subjected to ESI, presumably because the sample analyzed had been treated with potassium salt. The ESI-MS spectrum of the complex revealed that complexation also caused partial potassium/proton exchange. The m/z values of different charge states of the conjugate were reconciled if one dye molecule was added to BSA and three K⁺

BSA native				
Charge state	Measured <i>m</i> / <i>z</i>	Calculated <i>m</i> / <i>z</i>	% Error	
15	4475	4470	0.11	
16	4193	4193	0.00	
17	3943	3949	0.15	
18	3726	3731	0.13	
19	3530	3537	0.20	

Table 2 Experimental vs. measured m/z values for native BSA. K⁺

charges were used to obtain the calculated m/z values

ions were replaced with H^+ . The calculated m/z values for such stoichiometry and charges agree with these measured values, as presented in Table 3.

Table 3 Experimental vs. calculated m/z values for BSA complexed to the flavone dye ${\bf 1}$

BSA-flavone				
Charge state	Measured m/z	Calculated m/z^a	[% error]	[M + xK + yH]
15	4482	4482	0.00	$11K^{+} + 3H^{+}$
16	4206	4205	0.02	$13K^{+} + 3H^{+}$
17	3958	3960	0.05	$14K^{+} + 3H^{+}$
18	3743	3742	0.03	$15K^{+} + 3H^{+}$

^{*a*} For a BSA-flavone binding ratio of 1 : 1





Fig. 8 ESI-MS spectrum of the denatured BSA-flavone 1 complex. No peaks are present between m/z 4000 and m/z 4500, which are present in native BSA (Fig. 6). The peaks at around m/z 1500 are consistent with the denatured BSA.

To investigate whether the native conformation was necessary for complexation, the BSA and dye were allowed to conjugate, then acetic acid was added to bring the pH < 4 to denature the BSA complex. After addition of acetic acid, the ESI-MS spectrum became identical to that of the denatured BSA, indicating that the complex did not survive after denaturation (Fig. 8). The flavone dye was also treated with acetic acid, in order to confirm that acidic conditions were not damaging its structure (ESI Fig. S2[†]). The spectrum of the flavone dye before and after treatment with acetic acid were identical under the same conditions, demonstrating that no structural changes occur on the flavone dye at pH < 4. It should be noticed that the denatured BSA-flavone solution did not exhibit the m/z value and predominant charge state shifts, which were observed in the native BSA-flavone sample. The finding thus pointed out that only the BSA in the native conformation can complex the flavone dye.

As an interesting feature observed during the ESI-MS study, the predominant charge of +17 for native BSA was shifted to +16 when BSA was conjugated to a flavonoid dye (Fig. 6). It appeared that the ability of BSA to accommodate positive charges was perturbed when the hydrophobic flavone 1 was bound to the protein, making the +16 charge state more popular. Similarly, the +15 charge state of the BSA-flavone complex was increased in abundance at the expense of the +16 charge state in native BSA (Fig. 6). This result corroborates that one flavone dye was bound to BSA upon the interaction, as incorporating one molecule of flavone 1 increased the protein's hydrophobic property, thereby decreasing the protein's ability to accommodate positive charges by +1. The ESI-MS spectrum also provides evidence that the flavonoid dyes bind BSA to form a complex in a 1:1 ratio. This conclusion is further supported by estimating the molecular mass of 1 (ESI

Table S2†). From the calculated MW of the BSA-flavone 1 complexes with different charges, the average mass increase in the complexes with the +15, +16, +17 and +18 charges was calculated to be 332 ± 86 Da, which agrees reasonably well with the MW of flavone 1 (MW = 309). The data thus consistently indicated that only one flavone dye was incorporated in a BSA, forming a 1:1 complex.

As shown in Fig. 6, the BSA-flavone complex gave a set of new peaks with characteristic m/z values and charged states. Upon including one flavone in the protein-dye



Fig. 9 ESI-MS spectra of native HSA (bottom) and HSA bound to the flavone dye 1 (top). Shifts in the m/z values for the HSA-flavone peaks can be observed, as well as a shift in the predominant charge state for native uncomplexed HSA vs. native complexed HSA.



Fig. 10 ESI-MS spectra of native HSA (bottom spectrum) and HSA complexed to dye 2 [366 Da] (top spectrum). The *m*/*z* values along with the predominant charge state shift in the spectra of the complex (a and b) indicate a successful complexation.

complex, each charge state exhibited an increasing m/zvalue, in comparison with that from the BSA protein only. This can be seen more clearly in Fig. 7, where the spectrum is presented in an expanded region for the +16 and +17 charge states. One fundamental question is whether the detected BSA-flavone peak could include more than one flavone dye. If the BSA-flavone complex includes two flavone 1 molecules, one would expect the further mass increase in the resulting complex. For example, the mass for the +17 charge state of the "BSA-flavone 1" complex would be at "3958 + (309/17) = 3976", while the mass for the +16 charge state would be at "4206 + (309/16) = 4225." However, no obvious peak was detected at either m/z 3976 or 4225 in the spectrum of the BSA-flavone 1 complex (Fig. 7). This observation strongly suggests that both charge states gave consistent results, i.e. inclusion of one "flavone 1" molecule in the complex.

A similar feature was also observed when **1** was used to interact with HSA. The predominant peak in the +17 charge state for native HSA was shifted to +16 when HSA was conjugated to a flavonoid dye **1** (Fig. 9). As a consequence of incorporating a molecule of flavone **1**, the charge states (+15, +16, +17 and +18) of native HSA were decreased by **1** (to +14, +15, +16 and +17, respectively). These characteristics indicated the formation of the complex HSA-**1** with a **1** : **1** ratio.

Table 4Measured vs. expected m/z values for flavone dyes binding at
a 1:1 stoichiometric ratio with HSA

HSA-flavone 2 complex				
Charge state	Measured <i>m</i> / <i>z</i>	Calculated <i>m</i> / <i>z</i>	[% error]	Charges [M + xK + yH]
15	4482	4482	0.00	$11K^{+} + 4H^{+}$
16	4200	4200	0.00	$10K^{+} + 6H^{+}$
17	3959	3960	0.03	$13K^{+} + 4H^{+}$
18	3748	3748	0.00	$17K^{+} + 1H^{+}$

Complexes of HSA with other flavones

In order to explore the general trend of the protein's interaction with flavones, HSA was also used to examine its binding character with other flavone compounds such as 2 and 3, whose structures bear different substituents. Thus, flavone dyes 2 and 3 were individually left to conjugate with HSA and the resulting solutions were analyzed by ESI-MS. As shown in Fig. 10, the spectrum revealed a predominant charge state shift from +17 state for HSA to +16 state for HSA-dye 2, indicating that complexation has occurred. The m/z values in these spectra also showed that complexation was accompanied with a potassium/ proton exchange (Table 4), as observed with dye 1.

When assuming an HSA to flavone ratio of 1:1, the calculated m/z value matched well with the experimentally measured m/z values for all charge states (Table 4). The data further revealed that the average value of the measured molecular mass for HSA-flavone dye 2 complex was 66 798 Da (ESI Table S4†), and the average measured molecular mass for native HSA was 66 422 Da (ESI Table S3†). The mass increase by forming the HSA-flavone 2 complex was estimated to be 356 (= 66 798–66 442), which closely matched the molecular mass (MW = 366) for flavone 2. These results pointed out that the resulting complex of HSA-flavone 2 had a 1:1 stoichiometry. Similar results were also observed from HSA-flavone 3 with comparable low % error (ESI Fig. S3 & Table S5†), which maintained a 1:1 stoichiometric ratio.

Conclusion

Conjugation of albumin proteins (HSA and BSA) with flavone dyes (compounds 1–3; Scheme 1) is known to induce a large fluorescence signal for molecular imaging applications. Using ESI-MS, this study demonstrates a useful strategy to characterize the structure of the protein–flavone complexes that are formed by weak non-covalent interactions. The study is based

on the assumption that ESI (a soft ionization method) will allow the generation of multiple charges on protein structures without causing fragmentation, while protein's native conformations can be preserved in an ammonium acetate buffer. In the buffer solution, ESI-MS of the native forms of either HSA or BSA detected several well-separated protein species that exhibited different m/z values and charge states (ranging from +15 to +18). Inclusion of a flavone dye (such as 1) caused a spectral shift toward a higher m/z value, as well as a shift of the predominant charge state toward a lower value (e.g. ranging from the corresponding +14 to +17 charge state). The predominant charge state and m/z shift were only observed when BSA and HSA were in their native state, indicating that a non-covalent interaction is occurring between the dye and proteins. The observed shifts in m/z values and reduced charge states could be attributed to the inclusion of a relative hydrophobic flavone into the proteins' drug binding site, which attenuates the protein's ability to stabilize the multiple charges. Analysis of the ESI-MS data from the resulting protein-flavone conjugates consistently pointed to the 1:1 ratio for the protein-to-flavone dye, despite the multiple drug binding sites present in the HSA or BSA structure (Fig. 1). Therefore, the ESI-MS study here provides direct evidence for the non-covalent binding interaction between the albumin proteins and the flavone dyes examined. Since the flavones have been shown to be an important class of fluorescent dyes for their selective interaction with HSA, the study provides a valuable method to guide the further development of this class of materials for molecular imaging applications.

Author contributions

N. A. performed the MS studies, L. M. performed synthesis and fluorescence analysis. N. A. and L. M. wrote the initial draft, C. W. and Y. P. finalized the paper with input from all authors.

Conflicts of interest

The authors declare that they have no competing financial interests.

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