In Vitro Investigations of Acetohexamide Binding to Glycated Serum Albumin in the Presence of Fatty Acid

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Abstract: The interaction of drugs with human serum albumin (HSA) is an important element of therapy. Albumin affects the distribution of the drug substance in the body, as well as its pharmacokinetic and pharmacodynamic properties. On the one hand, inflammation and protein glycation, directly associated with many pathological conditions and old age, can cause structural and functional modification of HSA, causing binding disorders. On the other hand, the widespread availability of various dietary supplements that affect the content of fatty acids in the body means that knowledge of the binding activity of transporting proteins, especially in people with chronic diseases, e.g., diabetes, will achieve satisfactory results of the selected therapy. Therefore, the aim of the present study was to evaluate the effect of a mixture of fatty acids (FA) with different saturated and unsaturated acids on the affinity of acetohexamide (AH), a drug with hypoglycaemic activity for glycated albumin, simulating the state of diabetes in the body. Based on fluorescence studies, we can conclude that the presence of both saturated and unsaturated FA disturbs the binding of AH to glycated albumin. Acetohexamide binds more strongly to defatted albumin than to albumin in the presence of fatty acids. The competitive binding of AH and FA to albumin may influence the concentration of free drug fraction and thus its therapeutic effect.

Keywords: fluorescence and UV-Vis spectroscopy; glycation; human serum albumin; saturated and unsaturated fatty acids; acetohexamide

1. Introduction

The pace of life, improper eating habits (energy rich, unbalanced diet), and reduction of physical activity to a total minimum has caused an explosion in the number of diabetes all over the world. Diabetes, a civilization related chronic disease, considered a 21st century epidemic, remains a challenge for modern medicine. It is associated with disorders of the mechanisms regulating blood glucose level. Researchers argue that the complication and progression of the disease are closely related to the toxic effects of glucose and its metabolites, as one of the pathogenic factors. In patients with the increased blood glucose level, the glycation process intensifies, especially glycation of human serum albumin (HSA) [1]. During the persistent state of hyperglycemia, advanced glycation-end products (AGEs) arise, which are responsible for numerous tissue and organ pathologies [2]. The decrease in glycemia level is crucial to the prevention and inhibition of chronic diabetes (macro- and microvascular) disease progression and complications. This goal is achieved by the change of lifestyle and using anti-diabetic drugs. Acetohexamide (1-((p-acetylphenyl)sulfonyl)-3-cyclohexylurea, AH, Scheme 1) is a first-generation sulfonylurea derivative used to treat type 2 diabetes, especially in people whose
blood glucose cannot be controlled by diet. AH is very well absorbed from the gastrointestinal tract and binds to albumin in Sudlow sites I and II [3]. The choice of acetohexamide from many different anti-diabetic drugs was dictated by its physicochemical properties. Its absorbance at wavelengths used to excite the fluorescence of albumin was small enough that allowed high minimize inner filter effect. This is the basic factor limiting the use of the fluorescent technique in drug-protein interactions studies.

Scheme 1. Chemical structure of acetohexamide (AH).

Figure 1 presents AH binding sites to albumin complexed with glucose (GLC-HSA) within IB, IIB, IIA, and IIIA subdomains and amino acid residues with whom the ligand forms hydrogen bonds.

HSA belongs to the group of soluble proteins most abundant in blood, where its concentration reaches 40–50 g/L. HSA constitutes 55–65% of all proteins present in plasma [4]. Due to the low molecular weight (66.5–66.9 kDa) compared to other plasma globulins, HSA is involved in many physiological processes that include the regulation of osmotic pressure, pH control, and transport to many biologically active compounds (e.g., hormones, acids fatty acids, vitamins, and drugs) [5,6]. The transport of ligands, both exogenous and endogenous in a complex with albumin, is manifested by an increase in the solubility of these compounds in serum, a decrease in their toxicity and protection against their oxidation. In the HSA molecule there are two main specific binding sites for exo- and endogenous substances, referred to as site I (located in subdomain IIA) and II (in subdomain IIIA) according to the Sudlow nomenclature [5,7]. Another small drug binding site in HSA is the digitoxin site [8]. Carter et al. described six binding sites for structurally different ligands: two for binding small, heterocyclic or aromatic carboxylic acids (located in IIA and IIIA subdomains), two for binding of long chain fatty acids and small anionic compounds (located in domain I and III in subdomains B) and two sites for the binding of metal ions, subdomain IIIA. The albumin subdomain IA does not have ligand binding properties, despite homology to IIA and IIIA subdomains that bind most drugs [9]. HSA as a carrier protein is the main transporter of saturated and unsaturated fatty acids (FA). In its structure, it has at least seven asymmetric and heterogeneous fatty acid binding sites,
which are also located in the vicinity of most drug binding sites, i.e., in IIA (FA-binding site 7) and IIIA (FA-binding sites 3 and 4) subdomain of macromolecule [7,10,11]. The FA-binding site with the highest affinity is probably in IIIB albumin subdomain (FA-binding site 5) [12]. Under physiological conditions, the albumin molecule binds from 0.1 to 2 moles of FA. During fasting or extreme exercises and in some diseases, i.e., diabetes or obesity, the [FA]/[HSA] molar ratio may increase to 6:1 or higher [5,13]. Bojko et al., based on fluorescence studies, concluded that albumin forms a complex with two fatty acid molecules at [FA]/[albumin] molar ratio between 1:1 and 8:1. [14]. Many studies show that fatty acids compete for the binding site in the albumin molecule or interact with drugs [15]. The binding affinity between FA and HSA molecules mainly depends on the chemical properties of the particular acid, i.e., the chain length, both the presence and position of double bonds [16]. In polytherapy, ligands can compete with each other for a binding site and displace each other from albumin binding. The drug that forms a complex with the protein is a pharmacologically inactive spare pool, it is not subject to biotransformation and elimination. Only free drug fractions are responsible for therapeutic effect.

Binding of a ligand to albumin can change its spatial conformation, which prevents other ligands from binding, or strengthens existing binding. In addition, the binding of the drug to one binding site can change the structure of other binding sites or their number in the albumin molecule [17,18]. The interaction of drugs with HSA is an important element of therapy, because albumin affects the distribution of the drug substance in the body, as well as its pharmacokinetic and pharmacodynamic properties. On the one hand, inflammation, disease processes, or old age can cause structural and functional modification of HSA, causing binding disorders. On the other hand, the widespread availability of various dietary supplements that affect the content of fatty acids in the body, means that knowledge of the binding activity of transporting proteins, especially in people with chronic diseases, e.g., diabetes, will achieve satisfactory results of the selected therapy. This aspect is important during both the determination of drug doses and with respect to the method of their administration, which is particularly important in multi-drug treatments.

Therefore, the purpose of this work is to evaluate the effect of a mixture of fatty acids (FA) with different saturated and unsaturated acids on the affinity of acetohexamide (AH) for glycated albumin, simulating the state of diabetes in the body. Individual fatty acid mixtures corresponded to the FA content in physiological state and in various clinical states, proceeding with an increased concentration of saturated and unsaturated fatty acids.

2. Results and Discussion

In the present work, the quenching of albumin fluorophores fluorescence method has been used. This method is a sensitive tool for studying conformational changes in macromolecules caused by low molecular compounds or/and various processes, including glycation. The main goal of the study was to analyze the interaction of acetohexamide (AH), a drug with hypoglycaemic activity and a sulfonylurea derivative of the first generation, with glycated human serum albumin without (af)gHSA and in the presence of fatty acids ((af)gHSAphys, (af)gHSA4S, (af)gHSA8S, (af)gHSA4US, (af)gHSA8US). According to Stryer theory, the basic condition for the energy transfer to the drug molecule without radiations is not more than 10 nm distance between human albumin fluorophores (tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs)) and the drug chromophore, which results in the quenching of macromolecule fluorescence [19]. A comparison of the course of fluorescence quenching curves at excitation wavelength $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm was aimed in order to demonstrate the participation of tryptophanyl residue (Trp-214) or/and tyrosyl residues (Tyrs) in drug-albumin interaction. The differences in the course of quenching fluorescence curves excited at $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm indicate participation in the interaction of both Trp-214 residue located in subdomain IIA and tyrosyl residues located in IIA (Tyr-263), IB (Tyr-138, Tyr-140, Tyr-148, Tyr-150, Tyr-161), IIB (Tyr-319, Tyr-332, Tyr-334, Tyr-341, Tyr-353, Tyr-370), and IIIA (Tyr-401, Tyr-411, Tyr-452, Tyr-497) subdoamins [20,21]. The mechanism of drug interaction with albumin can be determined on the basis of Stern–Volmer curves. The linear $F_0/F = f(C_{drug})$ relationship indicates
a dynamic or static mechanism of macromolecule fluorescence quenching in the environment of subdomains containing amino acid residues that are involved in the formation of the drug–albumin complex. The positive/negative deviation from the straight line of the Stern–Volmer curve indicates the occurrence of both dynamic and static quenching [22]. Estimation of the distance between the drug molecule and excited albumin fluorophore is possible from the Lehrer modified Stern–Volmer curve (\(F_0/\Delta F = f(1/[C_{drug}])\)). Based on this dependence, the Stern–Volmer constant (\(K_{SV}\)) and the fraction of the initial fluorescence of albumin maximally available for the drug (\(f_a\)) are determined.

An increase in the distance between the albumin fluorophore and the drug molecule is manifested by an observed effect may be associated with the quenching fluorescence of excited fluorophores (tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs)) of albumin (af)gHSA (Figure 2a) and (af)gHSAphys (Figure 2b) by acetohexamide, which was found in no more than 10 nm proximity [19].

2.1. The Interaction of Acetohexamide with Human Serum Albumin in the Absence of Fatty Acids (AH-(af)HSA) and the Influence of Physiological Composition of Fatty Acids on AH-(af)HSA Complex (AH-(af)HSAphys)

As shown in Figure 2, an increase in acetohexamide (AH) concentration in the AH-(af)gHSA and AH-(af)gHSAphys systems causes a gradual decrease in macromolecule fluorescence intensity. The observed effect may be associated with the quenching fluorescence of excited fluorophores (tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs)) of albumin (af)gHSA (Figure 2a) and (af)gHSAphys (Figure 2b) by acetohexamide, which was found in no more than 10 nm proximity [19].
The red-shift of HSA fluorescence spectrum with the increase of ligand concentration the
afHSA cavity within IIA (Trp-214, Tyrs) or placed was not compact and the segment of polypeptide changed its conformation to a more extended
state after the addition of the asinesin. The emission of indole Trp-214 may be blue-shifted if the group
theasinesin–HSA system, and the structure of the hydrophobic subdomain where a tryptophan was
authors explained that tryptophan residue was brought to a more hydrophilic environment in the
phys fluorescence of free-drug albumin (af)gHSA system at excitation wavelength
aromatic rings of the acetohexamide molecule and aromatic amino acid rings of the hydrophobic
of AH is associated with a decrease in polarity (increase in hydrophobicity) of albumin fluorophores
GLZ-gHSA
a second generation sulfonylurea derivative, also caused a blue-shift of glycated (gHSA
phys)
towards shorter waves (blue-shift) by 4 nm relative to the spectrum (af)gHSA and (af)gHSA
max = 327–323 nm) (Figure 2a, in the main view) has been observed. The hypsochromic shift indicates a decrease in the polar nature, and thus the formation of a hydrophobic environment around the tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs) of (af)gHSA due to the interaction of AH with albumin. As in our previous work, the presence of gliclazide (GLZ), a hydrophilic environment of albumin fluorophore (Trp-214). Similar observations were noticed by
Figure 2. Emission fluorescence spectra of (a) (af)gHSA and (b) (af)gHSAphys at 5 × 10⁻⁶ mol L⁻¹ (0)
concentration in the presence of AH at 5 × 10⁻⁶ mol L⁻¹ (1) – 5 × 10⁻⁵ mol L⁻¹ (10) concentrations,
λex = 275 nm (in the main view) and λex = 295 nm (in the insert); the arrows indicate the shift
direction of the maximum fluorescence of (af)gHSA and (af)gHSAphys (Δλmax) with the increase of AH
centration; t = 37 °C.
In addition, in the AH-(af)gHSA system, after excitation at λex = 275 nm, the shift in the albumin
fluorescence emission band towards shorter waves (blue-shift) by 4 nm relative to the spectrum of the drug-free albumin (Δλmax = 327–323 nm) (Figure 2a, in the main view) has been observed. The hypsochromic shift indicates a decrease in the polar nature, and thus the formation of a hydrophobic environment around the tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs) of (af)gHSA due to the interaction of AH with albumin. As in our previous work, the presence of gliclazide (GLZ), a hydrophilic environment of albumin fluorophore (Trp-214). Similar observations were noticed by
Feng et al. [25] during the fluorescence analysis of theasinesin binding to human serum albumin
(HSA). The red-shift of HSA fluorescence spectrum with the increase of ligand concentration the
authors explained that tryptophan residue was brought to a more hydrophilic environment in the
theasinesin–HSA system, and the structure of the hydrophobic subdomain where a tryptophan was
placed was not compact and the segment of polypeptide changed its conformation to a more extended
state after the addition of theasinesin. The emission of indole Trp-214 may be blue-shifted if the group
is buried within a native protein, and its emission may shift to longer wavelengths (red-shift) when protein is unfolded [20].

Based on the obtained data from the emission fluorescence spectra, the quenching curves of (af)gHSA and (af)gHSA$_{phys}$ ($5 \times 10^{-6}$ mol·L$^{-1}$) fluorescence in the presence of AH ($5 \times 10^{-6}$ mol·L$^{-1}$ – $5 \times 10^{-5}$ mol·L$^{-1}$), $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm, have been drawn (Figure 3).

![Fluorescence quenching curves](image)

**Figure 3.** Fluorescence quenching of (af)gHSA and (af)gHSA$_{phys}$ ($5 \times 10^{-6}$ mol·L$^{-1}$) complexed with AH ($5 \times 10^{-6}$ mol·L$^{-1}$ – $5 \times 10^{-5}$ mol·L$^{-1}$) (a) $\lambda_{ex} = 275$ nm, (b) $\lambda_{ex} = 295$ nm and fluorescence quenching of (c) AH-(af)gHSA and (d) AH-(af)gHSA$_{phys}$ system for $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm; the error bars are smaller than the symbols.

The course of albumin fluorescence quenching curves illustrates the reduction in fluorescence intensity of human serum albumin without (af)gHSA and with (af)gHSA$_{phys}$ fatty acids with the increase of acetoheaxamide concentration in AH-albumin system (Figure 3a–d). The presence of fatty acids at physiological concentration affects the ability of acetohexsamide to quench albumin fluorescence. For (af)gHSA a stronger fluorescence quenching than for (af)gHSA$_{phys}$, at both excitation wavelengths $\lambda_{ex} = 275$ nm (Figure 3a) and $\lambda_{ex} = 295$ nm (Figure 3b), has been observed. This phenomenon indicates a greater ability of acetoheaxamide to absorb energy from defatted albumin than in the presence of fatty acids, that change macromolecule. For ligand: albumin molar ratio 10:1, the quenching of the internal (af)gHSA and (af)gHSA$_{phys}$ fluorescence by AH equals to 30.20% and 10.50%, respectively ($\lambda_{ex} = 275$ nm), while $\lambda_{ex} = 295$ nm equals to 31.70% and 4.60%, respectively. The comparison of fluorescence quenching curves of (af)gHSA and (af)gHSA$_{phys}$ in the presence of acetohexamide at the excitation wavelength $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm, allowed to indicate the fluorophores (Trp-214 and/or Tyrs) involved in the interaction with AH. Nearly identical course of albumin fluorescence quenching curves in AH-(af)gHSA (Figure 3c) system, at $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm...
(4.60% difference in quenching of the intrinsic albumin fluorescence), indicates the contribution of a Trp-214 residue (located in IIA subdomain of albumin) or its environment and a negligible contribution of Tyrs residues in the interaction of AH with (af)gHSA in the environment of binding site. A significant difference (56.60%) in quenching of (af)gHSAphys fluorescence registered for $\lambda_{ex} = 275$ nm and 295 nm (Figure 3d) indicates the participation in AH interaction with (af)gHSAphys, not only Trp-214, but also Tyrs residues. A stronger fluorescence quenching in AH-(af)gHSAphys system at excitation wavelength $\lambda_{ex} = 275$ nm than 295 nm may indicate a much easier access of AH to 17 tyrosyl residues (Tyr-30, -84, -138, -140, -148, -150, -161, -263, -319, -332, -334, -341, -353, -370, -401, -411, -497) located in IB, IIB, IIA, and IIIA [26] subdomains than to the tryptophanyl residue. It is noteworthy that the fluorescence quenching technique is not sufficient to indicate which Tyrs moieties are involved in AH binding. Literature data indicate that in human albumin structure there are two main specific binding sites for endo- and exogenous substances such as acetohexamide. These binding sites defined by Sudlow et al. as site I (in subdomain IIA, where Trp-214, Tyr-263 and His-240 are located) and site II (in subdomain IIIA, where Tyr-401, Arg-410 and Tyr-411 are located) [5,7].

Based on the data obtained from (af)gHSA and (af)gHSAphys in the presence of AH, the Stern–Volmer curves have been plotted, $\lambda_{ex} = 275$ nm (Figure 4a) and $\lambda_{ex} = 295$ nm (Figure 4b). The dashed lines indicate a model rectilinear course of Stern–Volmer dependence ($F_0/F = f([C_{AH}])$).

![Figure 4](image.jpg)

**Figure 4.** The Stern-Volmer curves for AH-(af)gHSA and AH-(af)gHSAphys. (a) $\lambda_{ex} = 275$ nm, (b) $\lambda_{ex} = 295$ nm; the error bars are smaller than the symbols.

Stern–Volmer curves plotted for glycated, defatted serum albumin in the presence of acetohexamide (AH-(af)gHSA system) show a different course from the curves plotted for AH-(af)gHSAphys system both at excitation $\lambda_{ex} = 275$ nm (Figure 4a), and $\lambda_{ex} = 295$ nm (Figure 4b). A stronger fluorescence quenching ($F_0/F$) at whole range of acetohexamide concentrations have been occurred for albumin (af)gHSA compared to the $F_0/F$ value obtained for glycated, fatted albumin (af)gHSAphys. Negative deviation from the rectilinear relationship $F_0/F = f([C_{AH}])$ for AH-(af)gHSA system at $\lambda_{ex} = 275$ nm (Figure 4a) and $\lambda_{ex} = 295$ nm (Figure 4b), indicates dynamic (collision) and static (formation of a stable ligand-albumin complex in the ground state) quenching fluorescence of glycated, defatted albumin by acetohexamide. Literature data indicate that the negative deviation from the rectilinear course of the Stern–Volmer curve is explained by occurring initially more easily available binding sites for the drug, and after saturating them, those more difficult to access in the macromolecule [27,28].

The rectilinear course of Stern–Volmer plots ($F_0/F = f([C_{AH}])$ relationship) obtained for the AH-(af)gHSAphys system may indicate dynamic or static fluorescence quenching mechanism. When the dynamic mechanism takes place, there is a collision of the acetohexamide molecule with albumin (af)gHSAphys fluorophores in excitation state. The static mechanism leads to a decrease in the intensity...
of emitted fluorescence when acetohexamide binds to the fluorophore molecule in the non-excited state, reducing the population of excited fluorophores [27].

From the $F_0/F = f[C_{AH}]$ relationship for a system with a linear course of the Stern–Volmer curve (AH-(af)gHSAphys system), the Stern–Volmer constants $K_{SV}$, the biomolecular quenching rate constants $k_q$ ($k_q = K_{SV}/\tau_0$) and maximum available fluorescence fraction of all albumin $f_a$ fluorophores were determined. Quenching parameters ($K_{SV}$, $k_q = K_{SV}/\tau_0$) and $f_a$ for a system with non-linear Stern–Volmer relationship (AH-(af)gHSA system) were determined from $F_0/\Delta F = f(1/[C_{AH}])$ relationship represented by Stern–Volmer equation modified by Lehrer [29]. The obtained results are given in Table 1.

Table 1. Stern-Volmer constants $K_{SV}$ (mol$^{-1}$L$^{-1}$), biomolecular quenching rate constants $k_q$ (mol$^{-1}$Ls$^{-1}$) and maximum available fluorescence fraction of all albumin fluorophores $f_a$ calculated for the AH-(af)gHSA and AH-(af)gHSAphys systems; $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm.

| $\lambda_{ex}$ = 275 nm | $K_{SV}$ ± RSD $^\dagger$ × 10$^4$ (mol$^{-1}$L$^{-1}$) | $^*$ $k_q$ ± RSD $^*$ × 10$^{12}$ (mol$^{-1}$Ls$^{-1}$) | $f_a$ ± RSD $^\dagger$ |
|------------------------|-----------------------------------------------|-----------------------------------------------|-------------------|
| AH-(af)gHSA            | 4.11 ± 0.04                                    | 6.85 ± 0.06                                    | 0.43 ± 0.01       |
| AH-(af)gHSAphys        | 0.23 ± 0.01                                    | 0.38 ± 0.02                                    | 1.00 ± 0.01       |

| $\lambda_{ex}$ = 295 nm | $K_{SV}$ ± RSD $^\dagger$ × 10$^4$ (mol$^{-1}$L$^{-1}$) | $^*$ $k_q$ ± RSD $^*$ × 10$^{12}$ (mol$^{-1}$Ls$^{-1}$) | $f_a$ ± RSD $^\dagger$ |
|------------------------|-----------------------------------------------|-----------------------------------------------|-------------------|
| AH-(af)gHSA            | 3.10 ± 0.05                                    | 5.17 ± 0.08                                    | 0.50 ± 0.01       |
| AH-(af)gHSAphys        | 0.10 ± 0.01                                    | 0.17 ± 0.02                                    | 1.00 ± 0.01       |

$^\dagger$ relative standard deviation; $^*$ calculated using: $k_q = \frac{k_{SV}}{\tau_0}$, where: $\tau_0 = 6.0 \times 10^{-9}$ s [30]—the average fluorescence lifetime of albumin without quencher.

Higher values of $K_{SV}$ and $k_q$ constant obtained for the AH-(af)gHSA system compared to $K_{SV}$ and $k_q$ obtained for AH-(af)gHSAphys system (Table 1) indicate the location of acetohexamide molecules closer to the fluorophores of glycated, defatted albumin (af)gHSA than fluorophores of fatty acid physiological mixture albumin ((af)gHSAphys). Exciting albumin (af)gHSA and (af)gHSAphys at wavelength $\lambda_{ex} = 275$ nm, fluorescence emission was observed not only from the tryptophanyl residue, but also from tyrosyl residues and higher values of $K_{SV}$ constants were obtained than at $\lambda_{ex} = 295$ nm excitation wavelength. This demonstrates the contribution of (af)gHSA and (af)gHSAphys tyrosyl residues in AH-albumin interaction. The IIA human serum albumin subdomain contains one tryptophanyl residue (Trp-214) and one tyrosyl residue (Tyr-263), which would indicate that in acetohexamide binding to albumin (af)gHSA and (af)gHSAphys other subdomains, where tyrosyl residues are located (IB, IIB and IIIA subdomains), are also involved. Our observations are consistent with the results obtained by Joseph et al. [3] who, using the high-performance affinity chromatography (HPAC) technique, indicated the IIA and IIIA subdomains, i.e., the I and II binding sites, respectively, as the main binding site of acetohexamide in the glycated human structure serum albumin. The order of fluorescence quenching rate constants $k_q$ (10$^{12}$ or 10$^{11}$) determined for AH-(af)gHSA and AH-(af)gHSAphys system clearly indicates a static fluorescence quenching mechanism in the acetohexamide-albumin system (Table 1). According to Lakowicz, when the maximum value of $k_q$ constant in the aqueous solution is 1 × 10$^{10}$ [M$^{-1}$s$^{-1}$], the dynamic fluorescence quenching mechanism occurs [20]. Since the values of $K_{SV}$ constant and $f_a$ fraction are inversely proportional, the decrease in $K_{SV}$ is accompanied by an increase in $f_a$. The presence of fatty acids reduces the strength of AH-(af)gHSAphys fluorophores interaction with the increase the availability ($f_a$) of AH to tryptophanyl and tyrosyl (af)gHSAphys residues.

To determine the nature of the interaction of acetohexamide with (af)gHSA and (af)gHSAphys, saturation curves (binding isotherms) were plotted in the AH-(af)gHSA and AH-(af)gHSAphys systems, $\lambda_{ex} = 275$ nm (Figure 5a) and $\lambda_{ex} = 295$ nm (Figure 5b).
with the hydrophobic fragments of its surface [23]. However, the shape of the binding isotherms determined to investigate the possible cooperation of acetohexamide binding to the macromolecule. For a specific class of binding sites was also obtained. Moreover, Hill interaction factors ($n_H$) were determined. The number of acetohexamide molecules forming a complex with one molecule of both (af)gHSA and (af)gHSA phys at equilibrium concentration, $\log[r/(1-r)] = n_H \log[L_f]$) equations, the association constants $K_a$, which determine the stability of formed drug-albumin complex, were determined. The number of acetohexamide molecules forming a complex with one molecule of both (af)gHSA and (af)gHSA phys (Figure 5a,b). Specific binding is characterized by high affinity and low binding capacity, while non-specific binding with low affinity and unlimited drug binding capacity [23].

Since acetohexamide saturates the binding sites of defatted ((af)gHSA) and fatted ((af)gHSA phys) albumin, using the Klotz (the independent variable is the inverse of the concentration of free ligand fraction, $1/r = f(1/[L_f])$) and Hill (the independent variable is the logarithm of ligand free fraction concentration, $\log[r/(1-r)] = f(\log[L_f])$) equations, the association constants $K_a$, which determine the stability of formed drug-albumin complex, were determined. The number of acetohexamide molecules forming a complex with one molecule of both (af)gHSA and (af)gHSA phys at equilibrium state ($n$) for a specific class of binding sites was also obtained. Moreover, Hill interaction factors ($n_H$) were determined to investigate the possible cooperation of acetohexamide binding to the macromolecule. The data are presented in Table 2.

**Figure 5.** Binding isotherms of (af)gHSA and (af)gHSA phys at $5 \times 10^{-6}$ mol·L$^{-1}$ concentration with AH at $5 \times 10^{-6} - 5 \times 10^{-5}$ mol·L$^{-1}$; (a) $\lambda_{ex} = 275$ nm, (b) $\lambda_{ex} = 295$ nm; the error bars are smaller than the symbols.

The shape of the binding isotherms ($r = f([L_f])$) obtained for (af)gHSA albumin indicates the mixed (specific and non-specific) nature of the interaction, which means that acetohexamide binds not only to its specific binding sites in defatted albumin molecule, but also non-specifically interacts with the hydrophobic fragments of its surface [23]. However, the shape of the binding isotherms for albumin saturated with fatty acids indicates the occurrence of only the non-specific nature of acetohexamide binding to (af)gHSA phys (Figure 5a,b). Specific binding is characterized by high affinity and low binding capacity, while non-specific binding with low affinity and unlimited drug binding capacity [23].

Since acetohexamide saturates the binding sites of defatted ((af)gHSA) and fatted ((af)gHSA phys) albumin, using the Klotz (the independent variable is the inverse of the concentration of free ligand fraction, $1/r = f(1/[L_f])$) and Hill (the independent variable is the logarithm of ligand free fraction concentration, $\log[r/(1-r)] = f(\log[L_f])$) equations, the association constants $K_a$, which determine the stability of formed drug-albumin complex, were determined. The number of acetohexamide molecules forming a complex with one molecule of both (af)gHSA and (af)gHSA phys at equilibrium state ($n$) for a specific class of binding sites was also obtained. Moreover, Hill interaction factors ($n_H$) were determined to investigate the possible cooperation of acetohexamide binding to the macromolecule. The data are presented in Table 2.

**Table 2.** Association constants $K_a$ (mol$^{-1}$·L), mean number of AH moles bound with one mole of (af)gHSA and (af)gHSA phys ($n$), the Hill coefficient ($n_H$) in AH-(af)gHSA and AH-(af)gHSA phys system; $\lambda_{ex} = 275$ nm, $\lambda_{ex} = 295$ nm.

|                      | Klotz Method |                      |                      |                      |
|----------------------|--------------|----------------------|----------------------|----------------------|
|                      | $\lambda_{ex} = 275$ nm | $\lambda_{ex} = 295$ nm |                      |                      |
|                      | $K_a \pm \text{RSD}$) ($10^4$ (mol$^{-1}$·L) | $n \pm \text{RSD}$) | $K_a \pm \text{RSD}$) ($10^4$ (mol$^{-1}$·L) | $n_H \pm \text{RSD}$) |
| AH-(af)gHSA         | 5.42 ± 0.06  | 0.93 ± 0.02          | 4.64 ± 0.01          | 0.98 ± 0.02          |
| AH-(af)gHSA phys    | 0.50 ± 0.17  | 1.00 ± 0.43          | 0.43 ± 0.02          | 0.95 ± 0.02          |
| AH-(af)gHSA         | 3.87 ± 0.07  | 0.93 ± 0.03          | 3.45 ± 0.03          | 0.98 ± 0.02          |
| AH-(af)gHSA phys    | 0.74 ± 0.20  | 0.96 ± 0.32          | 1.52 ± 0.02          | 1.27 ± 0.01          |

* Relative standard deviation.
From the straight line of Klotz equation for AH-(af)gHSA, AH-(af)gHSA_{phys} systems and Hill equation for AH-(af)gHSA, AH-(af)gHSA_{phys} systems, the binding of AH to macromolecules within one class of binding sites has been obtained. For AH-(af)gHSA complex, the association constants $K_a$ are more than 10-fold (for $\lambda_{ex} = 275$ nm) and more than 5-fold (for $\lambda_{ex} = 295$ nm) higher than the constants $K_a$ values obtained for AH-(af)gHSA_{phys} complex (Table 2), which indicates that AH has higher affinity for (af)gHSA than for (af)gHSA_{phys} albumin binding sites. This phenomenon is probably caused by the reduction of AH-albumin complex stability in the presence of fatty acids, when both tryptophanyl residue (Trp-214) and tyrosyl residues (Tyrs) have been excited ($\lambda_{ex} = 275$ nm) and $\lambda_{ex} = 295$ nm. The same results have been received in our other study (data not published). We have concluded that tolbutamide (TB), a hypoglycaemic drug, binds more easily to defatted albumin than in the presence of fatty acid. In the treatment of TB high fat diet increased the therapeutic effect of the drug. The number of binding sites $n$ close to unity indicates the existence of one specific AH binding site in the (af)gHSA molecule. Based on $n_H$ value ($n_H \approx 1$) lack of cooperativity in binding of AH to (af)gHSA has been observed ($\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm) (Table 2). For AH-(af)gHSA_{phys} complex at $\lambda_{ex} = 295$ nm, a Hill coefficient value higher than 1 ($n_H > 1$) points to a positive co-operability phenomenon, which probably means that binding of AH in one place increases the affinity of the drug/drugs for other macromolecule binding sites. The occurrence of cooperative binding results from the fact that under physiological conditions, fatty acid molecules (FA4 and FA5) bind to IIIA and IIIB subdomains [31], thus affecting the albumin conformation, i.e., changes the affinity of a macromolecule for acetoxyhexamide. These observations complement the results discussed earlier that the presence of fatty acids induces changes in the structure of the macromolecule. Under physiological conditions, human serum albumin binds a maximum of two fatty acid moles (FA). The formation of an albumin complex with two FA moles is possible when their molar FA:albumin ratio is within 1:1 and 1:8 [31]. Therefore, in order to assess the binding of AH to glycated human serum albumin in various clinical states, proceeding with increased concentration of saturated (FA_S) and unsaturated (FA_US) fatty acids, in the next part of the study albumin with four and eight times more amount of saturated ((af)gHSA_{4S}, (af)gHSA_{8S}) and unsaturated ((af)gHSA_{4US}, (af)gHSA_{8US}) fatty acids compared to physiological values have been analyzed.

2.2. The Interaction of Acetoxyhexamide with Human Serum Albumin at the Increased Concentration of Saturated ((af)gHSA_{4S}, (af)gHSA_{8S}) and Unsaturated ((af)gHSA_{4US}, (af)gHSA_{8US}) Fatty Acids

In the next part of the study, emission fluorescence spectra of glycated albumin at $5 \times 10^{-6}$ mol L$^{-1}$ concentration containing four ((af)gHSA_{4S}) (4.1 $\times$ 10$^{-5}$ mol L$^{-1}$) and eight times ((af)gHSA_{8S}) (6.9 $\times$ 10$^{-5}$ mol L$^{-1}$) more saturated fatty acids (FA_S) and four ((af)gHSA_{4US}) (5.9 $\times$ 10$^{-5}$ mol L$^{-1}$) and eight times ((af)gHSA_{8US}) (1.11 $\times$ 10$^{-4}$ mol L$^{-1}$) more unsaturated (FA_US) fatty acids in relation to the physiological value (2 $\times$ 10$^{-5}$ mol L$^{-1}$) have been obtained. Fluorescence of macromolecules excited at $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm decreases with the increase of acetoxyhexamide (AH) concentration (data not shown). Lack of shift in the fluorescence emission band of albumin relative to the albumin spectrum in the presence of AH ($\Delta \lambda_{max}$) may indicate the invariability of the hydrophobic/hydrophilic properties of AH binding site (near the Trp-214 residue in subdomain IIA and Tyrs residues in subdomain IB, IIIB, IIA, and IIIA of albumins).

As it was previously mentioned, after excitation of albumin at excitation wavelength $\lambda_{ex} = 295$ nm, the observed emission of fluorescence comes almost exclusively from the tryptophan residue, while for $\lambda_{ex} = 275$ nm from both the tryptophan residue and tyrosyl residues. Differences in quenching of albumin fluorescence at both excitation wavelengths $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm, in AH-(af)gHSA_{4S} (Figure 6c, in the main view), AH-(af)gHSA_{8S} (Figure 6c, in the insert) and AH-(af)gHSA_{4US} (Figure 6d, in the insert) systems, indicate the simultaneously participation of Trp-214 and Tyrs residues in the interaction of AH with albumin at appropriate binding site. The lack of differences in albumin fluorescence quenching curves in AH-(af)gHSA_{4US} system (Figure 6d, in the main view) after excitation at both wavelengths indicates the participation mainly Trp-214 residue.
For albumin with two times more saturated ((af)gHSA4S) and unsaturated ((af)gHSA4US) fatty acids, a stronger fluorescence quenching with increasing AH concentration has been recorded than for albumin (af)gHSA4S and (af)gHSA4US both after excitation λex = 275 nm (Figure 6a,b, in the main view) and λex = 295 nm (Figure 6a,b, in the insert). This demonstrates the greater ability of AH to absorb energy from excited fluorophores, as a result of conformational changes caused by a higher content of fatty acids in the structure of albumin.

In addition, for each concentration of AH in the drug-albumin system, the degree of albumin fluorescence quenching is greater after excitation at λex = 275 nm than at λex = 295 nm, which would indicate a significant participation of tyrosyl residues in the interaction of AH with (af)gHSA4S, (af)gHSA4US, (af)gHSA4US and (af)gHSA4US. Tyrosyl residues in position 401 (Tyr-401) and 411 (Tyr-411) located in IIIA subdomain of macromolecule play a major role in drugs binding [20,32]. The observed significant contribution of Tyrs residues in the interaction of AH with albumin indicates the IIIA subdomain as the likely acetohexamide binding site.

The analysis of Stern–Volmer plots (F/F0 = f[CAH]) showed a straight line for AH-(af), AH-(af)gHSA4US, and AH-(af)gHSA4US systems, indicating a collision or static mechanism of albumin quenching fluorescence by AH. However, for AH-(af)gHSA4S system, F/F0 = f[CAH] relationship showed a non-linear course (slight deviation in the direction of OY), which would indicate the simultaneous occurrence of dynamic and static fluorescence quenching mechanism, a similar effect that we observed in our previous work [33] concerning the theophylline (Th)–human albumin system.

Figure 6. Fluorescence quenching of (a) AH-(af)gHSA4S and AH-(af)gHSA4US, (b) AH-(af)gHSA4US and AH-(af)gHSA4US system for λex = 275 nm (in the main view) and λex = 295 nm (in the insert); (c) AH-(af)gHSA4S (in the main view), AH-(af)gHSA4US (in the insert) and (d) AH-(af)gHSA4US (in the main view), AH-(af)gHSA4US (in the insert) system for λex = 275 nm and λex = 295 nm; the error bars are smaller than the symbols.
study. Quenching parameters ($K_{SV}, k_q = K_{SV}/\tau_0$) and the fraction of maximum available fluorescence of albumin $f_a$ for AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4US}$ and AH-(af)gHSA$_{8US}$ were determined from $F_0/F = f[C_{AH}]$ relationship, while for AH-(af)gHSA$_{4S}$ from $F_0/\Delta F = f(1/[C_{AH}]$ relationship (Table 3).

$$\lambda_0 = \lambda_{ex} = 275\text{ nm}$$

| $\lambda_{ex} = 275\text{ nm}$ | $K_{SV} \pm \text{RSD}^\dagger \times 10^4$ (mol$^{-1}$·L$^{-1}$) | $a_k \pm \text{RSD}^\dagger \times 10^{11}$ (mol$^{-1}$·L$^{-1}$·s$^{-1}$) | $f_a \pm \text{RSD}^\dagger$ |
|--------------------------------|-------------------------------------------------|-------------------------------------------------|------------------|
| AH-(af)gHSA$_{4S}$            | 0.83 ± 0.19                                     | 13.83 ± 3.17                                    | 0.13 ± 0.04      |
| AH-(af)gHSA$_{4US}$           | 0.49 ± 0.01                                     | 8.17 ± 0.17                                     | 1.00 ± 0.01      |
| AH-(af)gHSA$_{8US}$           | 0.08 ± 0.01                                     | 1.33 ± 0.01                                     | 1.00 ± 0.01      |
| AH-(af)gHSA$_{8US}$           | 0.14 ± 0.01                                     | 2.33 ± 0.01                                     | 1.00 ± 0.01      |

| $\lambda_{ex} = 295\text{ nm}$ | $K_{SV} \pm \text{RSD}^\dagger \times 10^4$ (mol$^{-1}$·L$^{-1}$) | $a_k \pm \text{RSD}^\dagger \times 10^{11}$ (mol$^{-1}$·L$^{-1}$·s$^{-1}$) | $f_a \pm \text{RSD}^\dagger$ |
|--------------------------------|-------------------------------------------------|-------------------------------------------------|------------------|
| AH-(af)gHSA$_{4S}$            | 0.77 ± 0.06                                     | 12.83 ± 1.00                                    | 0.08 ± 0.01      |
| AH-(af)gHSA$_{4US}$           | 0.25 ± 0.01                                     | 4.17 ± 0.17                                     | 1.00 ± 0.01      |
| AH-(af)gHSA$_{8US}$           | 0.09 ± 0.01                                     | 1.50 ± 0.01                                     | 1.00 ± 0.01      |
| AH-(af)gHSA$_{8US}$           | 0.11 ± 0.01                                     | 1.83 ± 0.01                                     | 1.00 ± 0.01      |

$\dagger$ relative standard deviation; $^*$ calculated using: $k_q = \frac{K_{SV}}{\tau_0}$, where: $\tau_0 = 6.0 \times 10^{-9}$ s $^{[30]}$—the average fluorescence lifetime of albumin without quencher.

An order of quenching constant $k_q$ equals to $10^{11}$ clearly indicates the presence of static fluorescence quenching mechanism of albumin (af)gHSA$_{4S}$, (af)gHSA$_{4US}$ and (af)gHSA$_{8US}$ by AH. A 2-fold increase in saturated (AH-(af)gHSA$_{4S}$ system) and unsaturated (AH-(af)gHSA$_{4US}$ system) fatty acids in albumin (af)gHSA$_{4S}$ and (af)gHSA$_{4US}$, respectively, caused about two-fold ($\lambda_{ex} = 275\text{ nm}$) and three-fold ($\lambda_{ex} = 295\text{ nm}$) decrease and about two-fold ($\lambda_{ex} = 275\text{ nm}$) and one-fold ($\lambda_{ex} = 295\text{ nm}$) increase in the quenching parameters ($K_{SV}$ and $k_q$). This may suggest that in the presence of an increased amount of saturated fatty acids, acetohexamide binds at a considerable distance from the albumin fluorophores, which hinders the transfer of energy between them. However, a greater amount of unsaturated fatty acids facilitates energy transfer in the AH-albumin system (Table 3). In addition, it can be seen that the availability of albumin (af)gHSA$_{4S}$ fluorophores (Trp-214 residues, Tyrs residues) for individual AH binding sites is significantly hindered.

The course of binding isotherms, which determine the binding specificity of ligand to albumin, for AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4US}$ and AH-(af)gHSA$_{8US}$ systems is linear. The straight-line relationship of $r = f([L]_t)$ Taira and Terada $^{[23]}$ explained by the non-specific interaction of ligand with the hydrophobic fragments of macromolecule surfaces. However, the association constants $K_a$ determined from the Klotz relationship for the complexes AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4US}$, AH-(af)gHSA$_{8US}$ systems also prove the specific nature of acetohexamide binding within the molecule (Table 4). On the other hand, the non-linear course of Hill dependence obtained for the AH-(af)gHSA$_{4S}$ complex, confirms the interaction of AH mainly with albumin surface fragments. Straight line Klotz plots for AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4US}$, AH-(af)gHSA$_{8US}$ and Hill plots for AH-(af)gHSA$_{4S}$, AH-(af)gHSA$_{4US}$ and AH-(af)gHSA$_{8US}$ also indicate the presence of one independent class of AH binding sites in the albumin structure (or one binding site). Binding parameters ($K_a$, $n$) and Hill $n_H$ coefficient (interaction factor) has been summarized in Table 4.
Table 4. Association constants $K_a$ (mol$^{-1}$·L) and Hill coefficient ($n_H$) of AH-(af)gHSA$_{BS}$, AH-(af)gHSA$_{US}$, AH-(af)gHSA$_{4US}$ and AH-(af)gHSA$_{BSUS}$ systems; $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm

| $\lambda_{ex}$ (nm) | $K_a \pm$ RSD ($^{a}$) $10^4$ (mol$^{-1}$·L) | $n \pm$ RSD ($^{a}$) | $K_a \pm$ RSD ($^{a}$) $10^4$ (mol$^{-1}$·L) | $n_H \pm$ RSD ($^{a}$) |
|---------------------|--------------------------------------------|----------------------|--------------------------------------------|----------------------|
| 275 nm              | AH-(af)gHSA$_{BS}$                           | 0.92 ± 0.20           | 0.93 ± 0.25                                | 0.93 ± 0.25           |
|                     | AH-(af)gHSA$_{BS}$                           | 0.39 ± 0.07           | 0.98 ± 0.19                                | 0.98 ± 0.19           |
|                     | AH-(af)gHSA$_{4US}$                          | 0.70 ± 0.09           | 0.97 ± 0.07                                | 0.97 ± 0.07           |
|                     | AH-(af)gHSA$_{BSUS}$                         | 0.23 ± 0.05           | 0.99 ± 0.05                                | 0.99 ± 0.05           |
| 295 nm              | AH-(af)gHSA$_{BS}$                           | 0.85 ± 0.06           | 0.94 ± 0.08                                | 0.94 ± 0.08           |
|                     | AH-(af)gHSA$_{BS}$                           | 0.52 ± 0.04           | 0.98 ± 0.09                                | 0.98 ± 0.09           |
|                     | AH-(af)gHSA$_{4US}$                          | 0.87 ± 0.08           | 0.95 ± 0.05                                | 0.95 ± 0.05           |
|                     | AH-(af)gHSA$_{BSUS}$                         | 0.67 ± 0.06           | 0.96 ± 0.08                                | 0.96 ± 0.08           |

* relative standard deviation; $^{a}$—unable to calculate.

For AH-albumin complex with two-times the amount of saturated ((af)gHSA$_{BS}$) and unsaturated ((af)gHSA$_{4US}$) fatty acids, the $K_a$ constants are smaller than those obtained for the AH-AH-(af)gHSA$_{BS}$ and AH-(af)gHSA$_{4US}$ systems at $\lambda_{ex} = 275$ nm and 295 nm (Table 4). This means that a higher concentration of both saturated and unsaturated fatty acids in glycated albumin reduces the stability of the complex formed with acetohexamide. For AH-(af)gHSA$_{BS}$, AH-(af)gHSA$_{BS}$ and AH-(af)gHSA$_{4US}$, AH-(af)gHSA$_{BSUS}$ complexes, on average one ligand molecule binds to one albumin molecule ($n = 1$). In contrast, Hill interaction coefficient $n_H$ equals to unity ($n_H = 1$) and indicates a lack of cooperativity in the binding of AH to albumin (af)gHSA$_{BS}$, (af)gHSA$_{4US}$ and (af)gHSA$_{BSUS}$ in the vicinity of Trp residues. The same value of $n_H$ we received in our previous work [33] when we determined the Hill interaction coefficient for theophylline (Th)-albumin complex with 8-fold higher saturated fatty acid amount compared to physiological value (Th-dHSA-FA$_{BS}$). $n_H$ value greater than unity ($n_H > 1$) obtained for the complexes AH-(af)gHSA$_{4US}$ and AH-(af)gHSA$_{BSUS}$ excited at $\lambda_{ex} = 295$ nm indicates the phenomenon of positive cooperativity, which probably means that the binding of AH in one place (near Trp-214 residue) increases the affinity of the drug/drugs for the rest macromolecule binding sites. The presence of both saturated and unsaturated fatty acids affects the quenching of glycated, defatted albumin ((af)gHSA) fluorescence by acetohexamide (AH). For albumin (af)gHSA, fluorescence is quenched more strongly with the increase of drug concentration ($5 \times 10^{-6}$ mol·L$^{-1}$ - $5 \times 10^{-5}$ mol·L$^{-1}$) than for albumin containing four times ((af)gHSA$_{BS}$) ($4.1 \times 10^{-5}$ mol·L$^{-1}$) and eight times ((af)gHSA$_{BS}$) ($6.9 \times 10^{-5}$ mol·L$^{-1}$) more saturated fatty acids (FA$_{S}$) and four times ((af)gHSA$_{4US}$) ($5.9 \times 10^{-5}$ mol·L$^{-1}$) and eight times ((af)gHSA$_{BSUS}$) ($1.11 \times 10^{-4}$ mol·L$^{-1}$) higher unsaturated (FA$_{US}$) fatty acids in relation to the physiological value ($2 \times 10^{-5}$ mol·L$^{-1}$) (Figure 7). The largest differences in quenching of internal fluorescence relative to the control system (AH-(af)gHSA) were noted for the AH-(af)gHSA$_{4US}$ system at $\lambda_{ex} = 275$ nm (by 26.08%) and $\lambda_{ex} = 295$ nm (by 27.44%), while the smallest differences for AH-(af)gHSA$_{BS}$ at $\lambda_{ex} = 275$ nm (by 10.15%) and at $\lambda_{ex} = 295$ nm (by 20.29%) have been recorded. Stronger fluorescence quenching in AH-(af)gHSA$_{phys}$, AH-(af)gHSA$_{BS}$, AH-(af)gHSA$_{BS}$, and AH-(af)gHSA$_{4US}$ systems at excitation wavelength $\lambda_{ex} = 275$ nm than at $\lambda_{ex} = 295$ nm, probably indicates much easier drug access to tyrosyl residues (Tyr) than to the tryptophanyl residue (Trp-214). Only for AH-(af)gHSA and AH-(af)gHSA$_{4US}$ systems determined percentage differences in quenching fluorescence of albumin excited at $\lambda_{ex} = 275$ nm and $\lambda_{ex} = 295$ nm are within error limits of measurement, which indicates a greater participation of Trp-214 than Tyr residues in the interaction. For the molar ratio of AH-albumin 10:1, a two-fold increase in the amount of saturated and unsaturated fatty acids compared to the
physiological value of the acids causes (50.65% and 33.49% at $\lambda_{ex} = 275$ nm and 55.40% and 16.70% at $\lambda_{ex} = 295$ nm) a stronger quenching of the macromolecule fluorescence by AH.

Stern–Volmer $K_{SV}$, biomolecular quenching rate $k_q$ and association $K_a$ constants determined from the Stern–Volmer, Klotz, and Hill methods, respectively, for AH complex with glycated, defatted albumin ((af)gHSA) are characterized by higher values than for AH complex glycated in the presence of fatty acids albumin (AH-(af)gHSAphys, AH-(af)gHSA4S, AH-(af)gHSA8S, AH-(af)gHSA4US, AH-(af)gHSA8US), as previously discussed. The decrease in the association constant $K_a$ value demonstrates the effect of fatty acids on the reduction of AH affinity towards human serum albumin. Similar conclusions were reached by Shum and Jusko [34], who showed a weakness of theophylline–albumin interaction with an increase in the concentration of fatty acids under the physiological concentration in the macromolecule. Based on the in vitro results regardin the effect of fatty acids on the binding of acetohexamide to glycated human serum albumin, it can be assumed that under conditions of abnormal fat content in the body, the pharmacokinetics of the drug may be affected. A four-fold increase in saturated and unsaturated fatty acids amount relative to the physiological value causes $(50.65\% \text{ and } 33.49\% \text{ at } \lambda_{ex} = 275 \text{ nm and } 55.40\% \text{ and } 16.70\% \text{ at } \lambda_{ex} = 295 \text{ nm})$ a stronger quenching of the macromolecule fluorescence by AH.

### 3. Materials and Methods

#### 3.1. Reagents

Crystallized and lyophilized human serum albumin fatty acid free (fraction V) ((af)HSA, Lot No. 6312A) was purchased from MP Biomedicals SARL (Illkirch Cedex, France). Myristic acid (MYR, Lot No. R28576), oleic acid (OA, Lot No. 1912J), linoleic acid (LA, Lot No. 2353J), palmitic acid (PA, Lot No. 6798H) and stearic acid (SA, Lot No. 7729H) were provided by MP Biomedicals LLC (Irvine, CA, USA), Sodium azide (NaN₃, Lot No. BCBD6941V), acetohexamide (AH, Lot No. SLBF5611V) were

![Figure 7](https://via.placeholder.com/150)
obtained by Sigma-Aldrich Chemical Co. (Darmstadt, Germany) and Sigma-Aldrich Chemical Co. (Shanghai, China), respectively. D(+)-glucose (GLC, Lot No. A0299881) was supplied by POCH S.A. (Gliwice, Poland). All chemicals were of the highest analytical quality. The stock solution of AH, MYR, OA, LA, PA, and SA were prepared by dissolving appropriate amounts in methanol from Merck KGaA (Darmstadt, Germany, Lot No. 32373611/18).

3.2. Procedure of Preparation Fatty Acids Solutions for Fluorescence and Absorbance Studies

Defatted human serum albumin (af)HSA in the presence of glucose (GLC) at 5 × 10−6 mol·L−1 and 0.05 mol·L−1 concentrations, respectively, were prepared in phosphate buffer solution (pH 7.4, 0.05 mol·L−1) in double-distilled water and in the presence of sodium azide (NaN₃) (0.015 mol·L−1). Solution of protein was passed through a sterile Millex-GP syringe filters with 0.2 µm pores, and then it was incubated in sterile closed tubes for a period of 21 days at constant temperature of 37°C. After the incubation period, to remove the excess of unbound glucose, the solutions of glycated human serum albumin (af)gHSA was dialyzed extensively against 0.05 mol·L−1 phosphate buffer at pH 7.4 for 24 h.

To investigate the influence of fatty acids on glycated human serum albumin binding with acetohexamide (AH), the mixture of fatty acids (FA) corresponding to the physiological FA composition in human blood (mix 1) and FA four mixtures at different concentrations (mix 2–5) were prepared. Six glycated albumin samples were prepared without (af)gHSA and in the presence of a suitable fatty acid mixture (af)gHSAphys, (af)gHSA4S, (af)gHSA8S, (af)gHSA4US, (af)gHSA8US. Fatty acids physiological mixture at 2 × 10−5 mol·L−1 concentration (mix 1) contains FA mixtures at 8 × 10−6 mol·L−1 (oleic acid, OA), 6 × 10−6 mol·L−1 (palmitic acid, PA), 0.5 × 10−6 mol·L−1 (stearic acid, SA), 0.5 × 10−6 mol·L−1 (myristic acid, MYR) and 5 × 10−6 mol·L−1 (linoleic acid, LA) concentrations. The next two mixtures contain four (mix 2) and eight times (mix 3) more saturated fatty acids (PA, MYR, SA) in relation to the physiological value. The total concentration of these solutions was 4.1 × 10−5 mol·L−1 and 6.9 × 10−5 mol·L−1. The last two solutions contain four (mix 4) and 8 times (mix 5) more unsaturated fatty acids (OA and LA) in relation to the physiological value. Total concentration of these solutions was 5.9 × 10−5 mol·L−1 and 1.11 × 10−4 mol·L−1. The stock solution of MYR, OA, LA, PA, and SA at 1 × 10−3 mol·L−1 concentration was prepared by dissolving the right amount of acid in methanol, while mixtures of FA were prepared in phosphate buffer. A stock solution of acetohexamide (AH) at 5 × 10−3 mol·L−1 concentration was prepared in methanol. The content of methanol in the samples did not exceed 1% of tested protein solution total volume.

3.3. Instruments and Measurements Conditions

The fluorescence and absorbance measurements of the samples were recorded at 37°C using spectrofluorimeter JASCO FP-6500 (Hachioji, Tokyo, Japan) equipped with Peltier thermostat (ΔT ± 0.2°C) (error apparatus ± 1.5 nm) (Peltier thermostat is a part of JASCO FP-6500) and spectrophotometer JASCO V-760 (correcting error of apparatus for wavelength and photometric is equal to ± 0.3 nm and ± 0.002 Abs. at 0.5 Abs) (Hachioji, Tokyo, Japan), respectively. For the measurement standards quartz cuvettes 1 cm × 1 cm × 4 cm were used. The fluorescence spectra presented in the paper were corrected for the solvent dispersion (phosphate buffer) using the Spectra Manager program, and then analyzed using OriginPro version 8.5 SR1 software (Northampton, MA, USA). Finally, light scattering caused by buffer was subtracted from fluorescence of samples in each spectrum using software supplied by JASCO (Spectra Manager). The results of the study were expressed as a mean ± relative standard deviation (RSD) from three independent experiments. Linear regression by fitting experimental data to the appropriate equations were analyzed using OriginPro version 8.5 SR1 software (Northampton, MA, USA).

The analysis of the interaction of acetohexamide (AH) with glycated human serum albumin without (af)gHSA and in the presence of (af)gHSAphys, (af)gHSA4S, (af)gHSA8S, (af)gHSA4US, (af)gHSA8US fatty acids were done using albumin fluorescence quenching method. To excite fluorescence of the proteins, λex = 275 nm (exites tyrosyls and tryptophanyl residues, λem = 285 nm − 400 nm)
and $\lambda_{ex} = 295 \text{ nm}$ (excites tryptophanyl residue, $\lambda_{em} = 305 \text{ nm} - 400 \text{ nm}$) wavelengths were used. The measurements were conducted when the spectral width of the band (for monochromator of excitation and emission radiation) was equal to 3 nm, sample scanning speed of 200 nm/min, signal sensitivity "Medium", response time 4 s. For measurements of drug–albumin and drug–albumin-FA mixtures (mix 1–5), solutions of proteins at constant 5 $\times$ 10^{-6} mol L^{-1} concentration were used, without (0) and in the presence of acetohexamide (AH) at final concentration: 5 $\times$ 10^{-6} mol L^{-1} (1), 1 $\times$ 10^{-5} mol L^{-1} (2), 1.5 $\times$ 10^{-5} mol L^{-1} (3), 2 $\times$ 10^{-5} mol L^{-1} (4), 2.5 $\times$ 10^{-5} mol L^{-1} (5), 3 $\times$ 10^{-5} mol L^{-1} (6), 3.5 $\times$ 10^{-5} mol L^{-1} (7), 4 $\times$ 10^{-5} mol L^{-1} (8), 4.5 $\times$ 10^{-5} mol L^{-1} (9), 5 $\times$ 10^{-5} mol L^{-1} (10). Samples for fluorescence measurements have been made by titration method. By the use of Hamilton syringe a suitable volume of titrant (3 µL in 10 portions) has been added to 3 mL of albums immediately before the fluorescence measurement. The final AH:(af)gHSA, AH:(af)gHSA_{phys}, AH:(af)gHSA_{4S}, AH:(af)gHSA_{AS}, AH:(af)gHSA_{4US}, AH:(af)gHSA_{US} molar ratio was 10:1. The degree of glycated albumin fluorescence quenching by the acetohexamide was determined relative to the fluorescence of the non-ligand albumin solutions. Due to the inner filter effect (IFE) caused by the presence of the drug, the recorded fluorescence was corrected using the following formula [20]:

$$F_{cor} = \frac{F_{obs}}{10^{(\frac{\Delta \lambda_1 + \Delta \lambda_2}{2})}}$$

where: $F_{cor}$ and $F_{obs}$ are corrected and observed fluorescence intensity, respectively; $\Delta \lambda_1$ and $\Delta \lambda_2$ are the absorbance at excitation ($\lambda_{ex} = 275 \text{ nm}$ or $\lambda_{ex} = 295 \text{ nm}$) and emission wavelength for (af)gHSA ($\lambda_{em} = 275 \text{ nm}$: $\lambda_{em} = 327$ – 323 nm or $\lambda_{ex} = 295 \text{ nm}$: $\lambda_{em} = 334 \text{ nm}$) and (af)gHSA_{phys}, (af)gHSA_{4S}, (af)gHSA_{AS}, (af)gHSA_{4US}, (af)gHSA_{US} ($\lambda_{ex} = 275 \text{ nm}$: $\lambda_{em} = 323 \text{ nm}$ or $\lambda_{ex} = 295 \text{ nm}$: $\lambda_{em} = 329$–332 nm), respectively.

3.4. The Analysis of Acetohexamide–Albumins Interaction

Based on the calculated fluorescence emission intensities without and in the presence of fatty acids glycated human serum albumin, the quenching curves ($F/F_0$) vs. ligand:albumin molar ratio, where: $F$ and $F_0$ is the fluorescence intensity at the maximum wavelength of albumin in the presence and absence of a quencher, respectively) of (af)gHSA, (af)gHSA_{phys}, (af)gHSA_{4S}, (af)gHSA_{AS}, (af)gHSA_{4US} and (af)gHSA_{US} in the presence of acetohexamide (AH) have been plotted.

The course of Stern–Volmer curves (Equation (2)) allowed to conclude the mechanism of quenching fluorescence of albumins fluorphores by the studied drug [27].

$$F_0 = 1 + k_q \tau_0 |L| = 1 + K_{SV} |L|$$

where: $k_q$ is the bimolecular quenching rate constant [mol^{-1}\text{-L-s}^{-1}]; $\tau_0$ is the average fluorescence lifetime of albumin without of quencher $\tau_0 = 6.0 \times 10^{-9} \text{ s}$ [30]; $K_{SV}$ is the Stern–Volmer constant [mol^{-1}\text{-L}]; $|L|$ is the ligand concentration [mol\text{-L}^{-1}]; $|L|$ is the bound and free (unbound) drug concentrations [mol\text{-L}^{-1}].

From the modified Stern–Volmer relationship (Equation (3)), the Stern–Volmer $K_{SV}$ constant in AH:(af)gHSA, AH:(af)gHSA_{phys}, AH:(af)gHSA_{4S}, AH:(af)gHSA_{AS}, AH:(af)gHSA_{4US}, AH:(af)gHSA_{US} systems and also the fractional maximum protein fluorescence accessible for the quencher ($f_a$) were determined [29].

$$\Delta F = \frac{F_0 - F}{F_0} = \frac{1}{|L|} \cdot \frac{1}{f_a} \cdot \frac{1}{K_{SV}} + \frac{1}{f_a}$$

where: $f_a$ is the fractional maximum protein fluorescence accessible for the quencher.

Isotherms of drug binding to without and in the presence of fatty acids glycated human serum albumin have been obtained based on the graph of the function $r = f(|L_r|)$, where: $r = \frac{|L_r|}{[af]_{gHSA}}$ is the number of ligands moles bound per mole of protein molecule; $|L_r| = \frac{\Delta F}{\Delta F_{max}} (af)gHSA_{total}$, $\Delta F$ is the difference between $F_0$ and $F$, $\Delta F_{max}$ (maximal fluorescence change with complete saturation) is
evaluated from the linear part of the $\frac{\Delta F}{[L]}$ vs. $\frac{1}{[L]}$; $[(af)gHSA]$ is glycated serum albumin concentration [mol·L$^{-1}$] [23].

From the Klotz curves (Equation (4)) [35], in addition to the values of $K_a$ association constants, the number of drug molecules bound to one protein molecule ($n$) was determined.

$$\frac{1}{r} = \frac{1}{n} + \frac{1}{n \cdot K_a \cdot [L_f]} \quad (4)$$

where: $n$ is the number of binding sites for the independent class of drug binding sites in the glycated albumin molecule.

From Hill dependence (Equation (5)) [36] association constants $K_a$ and interaction coefficients $n_H$ were calculated.

$$\log\left(\frac{r}{1-r}\right) = n_H \cdot \log[L_f] + \log K_a \quad (5)$$

where: $n_H$ is the Hill’s coefficient.

3.5. Binding Sites Visualisation. Molecular Docking Simulation

To visualize the potential binding site of acetohexamide (AH) to glucose complexed human serum albumin (GLC-HSA), molecular docking simulation has been made using the CLC Drug Discovery Workbench ver. 1.0.2. (CLC Bio, a QIAGEN Company: Aarhus, Denmark) [License number: CLC-LICENSE-51JT8-DXYBY-2A3EW-ED80P-DGW80]. The crystallographic structure of glucose complexed human serum albumin (GLC-HSA) has been obtained from the PDB protein structure database (Protein Data Bank) using the PDB ID: 4IW2 [37]. Theoretical structure of acetohexamide (AH) was minimized and optimized by the Austin Model 1 (AM1)-semi-empirical method using the Atomic and Molecular Electronic Structure System (GAMESS) [38]. The AH docking procedure for the GLC-HSA structure was conducted simulating physiological conditions, in accordance with the default program configuration.

4. Conclusions

Fluorescence spectroscopy is a technique that allows to monitor the intermolecular interactions. To determine the distribution capacity and binding properties of glycated, defatted albumin ([(af)gHSA]) and albumin containing a physiological mixture of fatty acids ([(af)gHSAphys]), albumin containing 4 times ([(af)gHSA4S]) and 8 times ([(af)gHSA8S]) higher amount of saturated fatty acids (FA$S$) and 4 times ([(af)gHSA4US]) and 8 times ([(af)gHSA8US]) higher amount of unsaturated (FA$US$) fatty acids in relation to physiological concentration, a number of fluorescent quenching measurements of albumin fluorescence by acetohexamide (AH) were made. Based on the quantitative and qualitative analysis we can conclude that the presence of both saturated and unsaturated fatty acids disturbs the binding of acetohexamide to human serum albumin. Acetohexamide binds more strongly to defatted albumin (af)gHSA than to albumin in the presence of fatty acids. Competitive binding of AH and fatty acids to albumin may influence the concentration of free drug fraction and thus its therapeutic effect. It should be taken into account that under conditions of abnormal fat content in the body, the pharmacokinetics of the drug may be affected.

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**Sample Availability:** Samples of the compounds are available from the authors.

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