Favourable Influence of Hydrophobic Surfaces on Protein Structure in Porous Organically-Modified Silica Glasses

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Favourable influence of hydrophobic surfaces on protein structure in porous organically-modified silica glasses

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\textbf{A B S T R A C T}

Organically-modified siloxanes were used as host materials to examine the influence of surface chemistry on protein conformation in a crowded environment. The sol–gel materials were prepared from tetramethoxysilane and a series of monosubstituted alkoxysilanes, RSi(OR)\textsubscript{3}, featuring alkyld groups of increasing chain length in the R-position. Using circular dichroism spectroscopy in the far-UV region, apomyoglobin was found to transit from an unfolded state to a native-like helical state as the content of the hydrophobic precursor increased from 0 to 15%. At a fixed molar content of 5\% RSi(OR)\textsubscript{3}, the helical structure of apomyoglobin increased with the chain length of the R-group, i.e. methyl < ethyl < n-propyl < n-buty < n-hexyl. This trend also was observed for the tertiary structure of ribonuclease A, suggesting that protein folding and biological activity are sensitive to the hydrophilic/hydrophobic balance of neighboring surfaces. The observed changes in protein structure did not correlate with total surface area or the average pore size of the modified glasses, but scanning electron microscopy images revealed an interesting relationship between surface morphology and alkyl chain length. The unexpected benefit of incorporating a low content of hydrophobic groups into a hydrophilic surface may lead to materials with improved biocompatibility for use in biosensors and implanted devices.

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1. Introduction

In the past few years, numerous silica- and/or porous siloxane-based organic–inorganic materials have been employed to produce protein-doped hybrid materials as biosensors and as heterogeneous catalysts for industrial applications [1–6]. In particular, the design of organically-modified hydrophobic glasses for encapsulating lipases, a class of enzymes that mediate lipid reactions and function naturally at hydrophobic interfaces, has received much attention due to the potential application of these enzymes in the conversion of fats and oils to other products of value [7–18].

It is understood that protein function is dependent on proper folding of the polypeptide chain into a specific, three-dimensional structure and that protein structure, in turn, may be affected by its immediate surroundings [19–21]. Thus, in the crowded environment of a silica matrix, the physical and chemical properties of the glass are expected to influence protein structure and activity. This hypothesis may be tested by encapsulating proteins into chemically-modified silica networks and by monitoring slight alterations in protein conformation by spectroscopic techniques. Even though many reports indicate that sol–gel glass encapsulation retains the activity of a wide variety of enzymes, as reviewed in Refs. [1–6], few of these reports attempt to quantify the fraction of properly-folded protein, and the specific enzyme activities (normalized per mass of protein) are often omitted. Also, in the case of encapsulated enzymes, activity measurements may be influenced by multiple factors; diffusion of substrates into the pores, accessibility of enzyme molecules, and the adsorption of substrates to the silica surface, in addition to changes in protein structure, may reduce enzyme activity relative to the activity measured in bulk solution. Thus, the prevalence of unfolded or partially-unfolded proteins in sol–gel materials, as well as the contribution of unfolded proteins to lower specific enzyme activity, is hard to discern from the current literature.

We have reported previously that apomyoglobin, a protein of moderate stability in solution, becomes largely unfolded following glass encapsulation using a sol–gel protocol adopted by many investigators [22–24]. A change in protein conformation following glass entrapment also has been detected by other researchers [25,26]. Consequently, surface effects on protein structure are a genuine concern for encapsulation studies and for the field of biomaterials, in general. A careful examination of the relationship between protein structure and the physical properties of the host material may be critical for improving current applications and for the...
development of new biocompatible materials [6,27]. At the same time, such research may enrich our understanding of the biophysical forces that govern the protein folding–unfolding process, an important topic in the biomedical sciences because several human diseases are associated with misfolding of specific proteins [28].

With these issues in mind, organic–inorganic hybrid porous silica-based materials may constitute an ideal system for delineating the effects of surface chemistry on protein structure in crowded environments, including the crowded cytoplasm of a living cell [29].

Due to the large variety of commercially available, organically-substituted silane reagents, wet-aged-based silica glasses can be used as host materials for examining the structure and stability of a protein as a function of many variable glass properties (porosity, surface chemistry, surface area, etc.). Typically, these host materials are synthesized via the sol–gel technique, a method involving hydrolysis and polycondensation reactions with alkoxysilane precursors such as tetramethoxysilane (TMOS) or mixtures of TMOS and monosubstituted reagents, RSi(OR)₃ [30]. This approach is compatible with protein encapsulation because the process can be performed under mild conditions of temperature and pH, avoiding irreversible denaturation of the protein. Moreover, the resulting, optically transparent glass materials facilitate the use of spectroscopic techniques for examining protein structure. In this regard, circular dichroism (CD) spectroscopy has been demonstrated to be a particularly useful tool for detecting modest changes in conformation of glass-encapsulated proteins relative to their conformation in dilute solutions [22–24,31,32]. Although CD measurements may be considered less sensitive than fluorescence techniques and may require more protein per experiment, the average global structure of the protein may be monitored by CD; in the far-UV region, the entire protein serves as the chromophore due to the differential absorbance of left-handed and right-handed circularly polarized light by the peptide units that define the backbone [33,34].

Here we report on the conformation and stability of two model proteins, apomyoglobin and ribonuclease A, as a function of glass hydrophobicity. The secondary structure of apomyoglobin (apoMb) has been shown previously to be highly sensitive to changes in glass composition by far-UV CD [24], whereas ribonuclease A (RNase A) provides a means of checking the tertiary structure of a protein by CD analysis in the near-UV region. The structural behaviour of each protein is examined in TMOS-derived silica glasses and in organically-modified glasses made from a series of alkyltrimethoxysilanes (methyl, ethyl, n-propyl, n-butyl, and n-hexyl) at different contents of the modifying precursor (0.5, 10, 15%, the physical properties of the host silica matrices are characterized by porosimetry, N₂ adsorption, and microscopy techniques in order to search for a relationship between glass hydrophobicity and the observed changes in protein structure.

2. Materials and methods

2.1. Protein preparation

Apomyoglobin was prepared by 2-butanol extraction of myoglobin from horse heart (Sigma–Aldrich, lot M-1882) with subsequent dialysis against 5.0 M potassium acetate buffer, pH 5, containing 1.0 mM dithiothreitol and 0.50 mM ethylenediaminetetraacetic acid (EDTA from Aldrich). The dialyzed solution was filtered using a 0.22 µm syringe filter (Millex, Millipore) to remove any potential aggregates prior to encapsulation. The concentration of the apomyoglobin stock was determined by CD using the following formula [34]:

\[ C = \frac{\theta \times 100 \text{mM}}{[\theta]_{\text{ DBNull}}} \]

where \([\theta]_{\text{DBNull}}\) is the molar ellipticity for apoMb (~19,000 deg cm² dmol⁻¹ at 222 nm), \(\theta\) is the path length of the cuvette (0.2 cm), MW is the protein molecular weight (16,951 g mol⁻¹ for apoMb), \(N_a\) is the number of amino acid residues per protein (Nₐ = 153 for apoMb), the factor 100 originates from the conversion of the molar concentration to the dmol/cm³ concentration unit, and C is the protein concentration (mg/ml). After prolonged storage at 5°C, the structural integrity of apomyoglobin was confirmed by reappearance of the Soret band at 408 nm upon 24-hr denaturation (Hewlett-Packard 8452A Diode Array Spectrophotometer). Pure ribonuclease A Type XII–A from bovine pancreas was purchased from Sigma–Aldrich (R5000). RNase A (MW = 13,674 g mol⁻¹, Nₐ = 124) was dissolved in potassium phosphate buffer (10 mm) to obtain an approximate concentration of 30 mg/ml. The final concentration of the RNase A stock solution was determined from the absorption peak at 276 nm using \(A_{276} = 0.19\) for a 1-cm path length [35]. MilliQ-purified water was utilized for all solutions (Millipore).

2.2. Protein encapsulation in sol–gel glasses

Proteins were encapsulated in sol–gel glasses made from TMOS (Acros) and a series of alkyltrimethoxysilane reagents, RSi(OME)₃, where R = —ethyl, ethyl, n-butyl, or n-hexyl, (purity ~ 99%) from Gelest, Inc. (Morrisville, PA, USA). n-Propyltrimethoxysilane was supplied by Aldrich. Molar ratios of 5, 10, and 15% of the organically-modified silane relative to total silane were calculated on the basis of 6.78 mmol of total Si content (2.0% MW, in the case of pure TMOS). The sol was obtained by co-hydrolysis of TMOS with the alkoxysilane precursors. For a typical experiment, the necessary volume of silane precursors for a given glass was mixed in the same tube, divided into two equal aliquots, and then 214 µl of water and 30 µl of 0.040 N HCl were added to each aliquot. Hydrolysis of the sol was performed by sonication in an ice-water bath for 30 min (Branson 1510 ultrasonic cleaner). After verifying by eye that a homogenous solution was achieved, the two aliquots were combined, and a 2.0 ml volume of the sol was added to 3.0 ml of the aqueous sol–gel containing the desired protein in 5 mm potassium phosphate buffer, pH 7. The final concentration of protein per total volume of sol–gel mixture was 0.49 mg/ml for apoMb and 7.0 mg/ml for RNase A; the higher concentration of RNase A is required for far-UV CD analysis in the near-UV region. The sol–gel mixture was homogenously mixed and immediately transferred to a plastic cassette of 1-mm thickness (Invitrogen #NC2010). The cassette was covered with parafilm and stored upright at 5°C for 2–3 weeks before use. After the wet-aging period, the glass sheets were removed by breaking open the cassette. Rectangular pieces of glass were cut with a razor blade and immersed in 3 ml of phosphate buffer solutions (pH 7) at two different concentrations for equilibration (10 mm and 100 µM). The glass samples were left immersed overnight at room temperature prior to CD analysis. No leaching of the protein from the glasses was observed at ambient temperatures.

2.3. Circular dichroism spectroscopy

Changes in protein structure were monitored by CD spectra as recorded on an Aviv Model 215 circular dichroism spectrometer equipped with a Peltier-type thermodenamic cell holder. Glass samples were placed in a 2 mm path quartz cuvette filled with the corresponding equilibration buffer during CD analysis. Spectra were recorded at 25°C in the far-UV region for apoMb and in the near-UV region for RNase A to monitor the secondary and tertiary structure of each protein, respectively. Potassium phosphate buffer was used as the sample blank, and the background signal was subtracted from all spectra before conversion to units of molar ellipticity. The thermal stability curves were recorded in five-degree steps from 25 to 90°C with a 30-s delay at each temperature before recording the ellipticity at 222 nm. Longer delays for temperature equilibration had no significant effect on the thermal denaturation profiles. The largest source of uncertainty in all CD measurements is the protein concentration (~5%). For a given figure, the same volume of stock solution of protein was used for every glass sample such that a small uniform error in calculation of protein concentration and molar ellipticity will have no effect on the interpretation of results.

2.4. Aerogel preparation

Prior to the physical characterizations described in Sections 2.5 and 2.6, the gels were dried at ambient temperature in a manner that minimizes shrinkage due to collapse of the pores [36,37]. Multi-step solvent exchange was undertaken with non-polar solvents to avoid the capillary action associated with drying and shrinkage [38,39]. First the gels were washed with acetone for 3 days in a capped vial, exchanging the acetone once per day. Subsequently, the process was repeated with pentane, a solvent of low surface tension. Next, the vials were opened and covered with aluminum foil through which a 1 mm diameter pinhole was made to allow slow evaporation of the pentane. The resulting solid materials were dried for 24 h at 37°C to form aerogels suitable for physical analysis.

2.5. Porosity and Brunauer–Emmett–Teller (BET) surface area measurements

The surface area and porosity determinations of aerogel samples were carried out using a Gemini apparatus from Micromeritics. The samples (ca. 40 mg) were cleaned by flowing nitrogen (ca. 30 ml/min) for 6 h at 50°C and for 12 h at 120°C in a Flow-Pep 060 oven, also from Micromeritics. The gels, helium for calibration (99.999%) and nitrogen for adsorption–desorption (99.999%), were obtained from L’Air Liquide (Spain). Full adsorption–desorption isotherms were recorded, and the
multipoint BET method was used for analysis. Specific surface area was determined from the adsorption branch, whereas pore size distribution was determined from the desorption branch using the Wheeler method.

2.6. Scanning electron microscopy (SEM) and atomic force microscopy (AFM)

The surface morphology and microstructure of the modified and unmodified silica-based aerogels were characterized by SEM using a Hitachi S-4300 Scanning Electron Microscope. Samples were attached to aluminum stubs with double-coated carbon conductive tabs and were gold sputtered (4.5–5.2 nm) using Auto Sputter Coater by Pelco (Model SC-7). The edges of the coated samples were brushed with colloidal graphite for improved conductivity of the sample surface, dried overnight, and analyzed. The images were acquired at 5 kV. AFM measurements were performed using a Veeco Dimension 5000 instrument. AFM tips from Veeco, model TESP, made of antimony (n) doped Si of radius of 8–12 nm were used. The vibrating frequency was in the range of 320 kHz, and the tapping mode was employed at a scanning rate of 1 Hz. Average domain sizes (i.e. agglomerated SiO₂ particles) were determined from the SEM images according to the given scale.

3. Results

3.1. Protein structure in alkyl-modified silica glasses

The average structures of proteins encapsulated in the pores of wet-aged-based silica glasses were determined by CD spectroscopy in TMOS-based glasses [Si(OME)₄] that were modified by addition of alkyltrimethoxysilane precursors [RSi(OEt)₃]. The CD signals originating from the protein in glass samples may be compared directly to the protein signal obtained in dilute solution because the data are normalized to units of molar ellipticity. At a fixed value of 5% RSi(OEt)₃ for each glass, a significant increase in the secondary structure of apomyoglobin is observed in the following order: unmodified < methyl < ethyl < n-propyl < n-butyl < n-hexyl (Fig. 1). The molar ellipticity of apoMb in the far-UV region approaches the spectrum of the native protein in dilute solution as the alkyl chain length increases. At a wavelength of 222 nm, where molar ellipticity is an approximate measure of helical content, the value for apoMb in solution is 19,000 deg cm²/dmol, whereas the molar ellipticity for the 5% hexyl-modified glass is 19,750 deg cm²/dmol, indicating slightly more helical structure in the glass than observed in solution. By contrast, the 100% TMOS-derived glass yields the weakest CD signal indicating that apoMb is largely unfolded in the control matrix, as reported previously [22–24].

The relationship between protein structure and glass hydrophobicity is confirmed and strengthened by monitoring apoMb structure in a set of glasses of increasing molar content for each of the alkylsilane reagents employed in this study. When the molar content of RSi(OEt)₃ is varied from 0 to 15%, a corresponding increase in helicity of apoMb is observed for each set of glasses (Fig. 2). The hexyl-modified glass could not be analyzed above a value of 5% incorporation due to a loss of optical transparency; the other four glasses were observed to lose transparency above 15%. We also attempted to make a glass with n-octyltrimethoxysilane, but this reagent led to a phase separation such that formation of a homogeneous transparent glass was not possible, even at a low molar content of 5%.

The near-UV CD spectra of another protein, RNase A, were obtained following encapsulation in the same glasses as apoMb (Fig. 3). The near-UV spectra arise from asymmetry in the packing around aromatic residues and, thus, report on changes in the tertiary structure of RNase A. The changes in molar ellipticity indicate that RNase A structure also is enhanced as the hydrophobicity of the glass increases (Fig. 3). Unfortunately, changes in the tertiary structure of apoMb cannot be monitored by this technique because the signal originating from its two tryptophan residues offset each other, resulting in a featureless spectrum [40].

In previous studies, molar salt solutions were shown to have a significant effect on the structure of apoMb in unmodified glasses [23]. As shown in Fig. 4, an increase in phosphate concentration from 10 mM to 1.0 M yields an increase in the molar ellipticity of apoMb in the unmodified glass from 4300 to 15,000 deg cm²/dmol at 222 nm, as expected. When the alkyl-modified glasses were equilibrated in the 1 M potassium phosphate solution, the salt effect on apoMb structure was more subtle and diminished with increasing chain length of the alkyl group. In the 10% ethyl-modified glass, for example, phosphate addition did not induce the same helical endpoint as the control glass made from 100% TMOS (Fig. 4). Thus, the favourable effects of molar phosphate concentration and glass hydrophobicity on protein structure are not additive. This result strongly suggests that the effects of altered glass hydrophobicity and molar salt addition are mediated by different mechanisms. Not surprisingly, the addition of 1 M phosphate had no effect on the

![Fig. 1. The helicity of encapsulated apoMb increases with increasing chain length of the organic modifier. CD spectra in the far-UV region for apoMb in unmodified and modified glasses (solid curves) and in dilute solution (dashed curve). All spectra were measured in the presence of potassium phosphate (10 mM, pH 7).](image1)

![Fig. 2. ApoMb structure as a function of molar content and chain length of the alkylsilane precursor. The molar ellipticity at 222 nm is proportional to the helical content of the protein, and the ellipticity value for the native protein in dilute solution is 19 × 10⁴ deg cm²/dmol.](image2)
native-like structure of apoMb in the 10% butyl-modified glass (Fig. 4).

3.2. Thermal stability of the proteins in the glasses

One major goal of glass encapsulation is to enhance protein stability. The data in Fig. 5 indicate that apoMb is extremely stable against thermal denaturation in both the modified and unmodified glasses when heated from 25 to 90 °C. For example, the molar ellipticity of apoMb in the 10% butyl-modified glass only decreases from 18,000 to 17,300 deg cm$^2$ dmol$^{-1}$ at a wavelength of 222 nm. At the other extreme, apoMb in solution proceeds to unfold through a broad transition from 19,000 to 8500 deg cm$^2$ dmol$^{-1}$ (Fig. 5(a)). When a glass sample is held at 90 °C for extended times (30 and 60 min), apoMb structure remains stable. The CD spectra of the 10% butyl-modified glass show the features of a folded protein even after a long period of heat treatment; the CD signal at 222 nm decreases only slightly from 18,000 to 17,100 deg cm$^2$ dmol$^{-1}$ after holding the sample at 90 °C for 60 min (Fig. 5(b)). This modest change in signal may originate from a minor leaching problem at the elevated temperature. In contrast, the ellipticity of the protein in dilute solution decreases drastically after extended times at 90 °C due to aggregation and precipitation of the unfolded protein (Fig. 5(b)).

3.3. Specific surface area and porosity of the silica-based aerogels

The CD results suggest that the alkyl chain of the organically-modified glasses may induce a length-dependent change in one or more physical properties of the host matrix, contributing to an apparent increase in the fraction of properly-folded protein. To address this possibility, the specific surface areas and porosities of both the modified and unmodified glasses were determined after conversion to aerogels. The results for TMOS and for the modified host matrices of 10% molar composition (R = methyl, ethyl, propyl, butyl) or 5% composition (R = hexyl) are summarized in Table 1. The measurements indicate that pore volume and pore size generally decrease upon alkyl modification of the silica surface, both in the presence and absence of protein. The average pore size decreases...
from 10.8 nm to about 4 nm in the absence of apoMb, and the average pore size decreases from 8.9 nm to about 4 nm when apoMb is present. Thus, all of the modified glasses, from methyl to hexyl, exhibit the same approximate pore size (~4 nm) with or without protein encapsulation. The reduced pore sizes are accompanied by a 25–75% increase in the specific surface area of the modified silica matrices relative to the unmodified glass (Table 1).

In Fig. 6, two physisorption isotherms are shown that are typical of those obtained for the modified and unmodified silica glasses. The isotherms were found to be independent of protein addition; glasses of the same composition yield the same profile, with and without apoMb encapsulation. Although both isotherms have the characteristics of a mesoporous silica material, it can be seen that the unmodified glass yields a different type of isotherm compared to the organically-modified host matrix (Fig. 6). The hysteresis profile for the 100% TMOS-derived glass, with or without protein encapsulation, corresponds to type B in de Boer’s classification, now referred to as type H3 by IUPAC [41,42]. This classification is associated with adsorption into slit-shaped pores or into the space between parallel platelets. In contrast, all of the modified silica samples exhibit type E hysteresis in de Boer’s classification (type H2 by IUPAC). This profile is associated with adsorption into bottleneck-shaped pores, that is, pores for which the inner compartment is wider than the entrance.

3.4. Surface morphology of glasses

The microstructures of the unmodified TMOS-derived and the modified alkylsiloxane-derived aerogels were characterized by scanning electron microscopy, with and without apoMb addition. Interestingly, the presence of apoMb has a major influence on the surface morphology of the host glass materials. The surface of glasses formed in the absence of protein appears highly homogeneous with an even distribution of silica particles (Fig. 7(a,c)), whereas all glasses containing apoMb exhibit serpentine-like features, as exemplified by the images of the 100% TMOS and 10% ethyl-modified glasses (Fig. 7(b,d)). Moreover, in the case of the unmodified silica aerogel, the serpentine domains appear to form semi-regular arrays (Fig. 7(b)). Digital analysis of each image indicates a correlation between alkyl chain length and the average domain size of the serpentine substructures (Fig. 8). When a 10% ethyl-modified glass was examined by atomic force microscopy in tapping mode, the higher depth contrast of the 10 × 10 μm square image reveals a distribution of spiral chains of ~20 nm size formed by the interconnected silica particles (Fig. 9).

4. Discussion

Hydrophobic, organically-modified silica glasses have been used previously to enhance the activity of glass-encapsulated lipases [7–18]. The hydrophobic character of such glasses has been verified by several means including contact angle measurements [43], solid-state 29Si MAS NMR, and Fourier transform infrared spectroscopy [17,44]. Although most enzymes prefer an aqueous environment, it is rational to place lipases in a hydrophobic environment because these enzymes mediate reactions at lipid–water interfaces and because the substrates of these enzymes are highly non-polar molecules. However, studies involving glass-encapsulated lipases typically compare rates of product formation without providing any insight on whether or not the glass composition also influences enzyme structure. If a change in glass hydrophobicity alters both the fraction of properly-folded enzyme and the partitioning of the enzyme substrate into the matrix, then it would be difficult to discern one effect from the other based solely on kinetic measurements of enzyme activity.

In the current work, we examine the structure of glass-encapsulated apoMb, a model protein that is not expected to favour a hydrophobic environment. We demonstrate that relatively small amounts of alkyl-modifying precursors (5–15%) can have a drastic effect on the helical content of the glass-entrapped protein. We were unable to examine all of the hydrophobic glasses above 15% modification due to a phase separation phenomenon that leads to a loss of optical transparency. Phase separation can be minimized by co-hydrolysis of the organosilane reagents, as employed here, instead of separate hydrolyses followed by mixing [8,45,46]. Our glasses appear to be under the reported threshold for phase separation of co-hydrolyzed samples; for example, glasses with up to 20% by volume of methyltriethoxysilane or 5% propyltrimethoxysilane have been prepared in tetraethoxysilane (TEOS) with no sign of phase separation, as monitored by several optical techniques [8,46]. The higher threshold for propyltrimethoxysilane in the current
work (15% molar) may be due to slight differences in preparation (TMOS versus TEOS) and in the commercial source of reagents.

The helical content of apoMb, measured as molar ellipticity in the far-UV region, was found to increase as the alkyl chain length of the precursor increased from one to six carbons (Fig. 1) or as the content of the hydrophobic precursor increased (Fig. 2). Importantly, the secondary structure of encapsulated apoMb did not exceed that of the native protein in dilute solution for all but one of the glasses examined in this study; the 5% hexyl-modified glass resulted in a slightly more helical protein than observed in solution. If the CD spectrum of the glass-encapsulated protein had become significantly more negative in the far-UV region than the solution structure (e.g. below 24,000 deg cm² dmol⁻¹ at 222 nm), it might indicate that the globular, tertiary structure of the protein has been compromised. We know of only one condition where a protein may gain helical content without a simultaneous gain in tertiary structure, and that is the case where water-miscible alcohols are added to the solvent phase. Because we never add alcohol to any of the samples in this work, and because apoMb is a protein with ~60% α-helical structure in the native state, it seems likely that the gain in helical content of this protein in hydrophobic, organically-modified glasses is accompanied by a gain in proper tertiary structure. Our suggestion that apoMb acquires more tertiary structure in alkyl-modified glasses is reinforced by the fact that the tertiary structure of another protein, RNase A, was not diminished and was somewhat enhanced following encapsulation in the same glasses, as monitored by near-UV CD (Fig. 3). A hyperhelical structure relative to dilute solution has been reported for apoMb following encapsulation in trifluoropropyl-modified glasses [24]. This result was attributed to the presence of organic fluorine, and, because the helicity of the protein never exceeded that of the heme-bound holoprotein, the observed hyperhelical structure may represent the true native state that prevails in a crowded environment [21].
To better understand the relationship between glass hydrophobicity and protein structure, we characterized some of the physical properties of alkyl-modified glasses using the tools of materials science. BET measurements indicate that all five alkyl precursors have a comparable effect on the glass, independent of chain length; surface area is increased and pore volume is decreased to a similar extent for all modified glasses relative to the unmodified glass (Table 1). This result is consistent with the formation of smaller silica particles prior to gelation due to a preponderance of reaction-terminating alkylsilane groups on the surface of the growing particle; at much higher concentrations of the alkyltrimethoxysilane precursor, phase separation can lead to the formation of crystalline oligomers, resinous materials, and precipitates [47].

Comparing columns three and six of Table 1, one observes a small but consistent decrease in average pore size of all glasses upon addition of protein. The effect of protein addition on pore size suggests a direct interaction between the silica particles and protein molecule during sol–gel processing. However, there is clearly no correlation between pore size and apoMb structure since 10% ethyl-, propyl-, and butyl-modified glasses yield the same pore size (Table 1), whereas the structure of the protein differs greatly in the same three glasses (Fig. 2).

The influence of pore size on the biological activity of enzymes in sol–gel matrices has been a subject of discussion by other investigators. Reetz and co-workers noted that lipase activity in alkyl-modified silica host materials might be expected to increase with increasing pore volume due to better accessibility of the substrate, but the opposite was found experimentally [7]. The authors concluded that the catalytic activity of lipases is not related to pore size but most likely a function of other factors related to the glass hydrophobicity, such as partitioning of the reactants or an interface-dependent conformational change in a specific loop that activates the enzyme. In a recent paper, Nourreddini and Gao came to the same conclusion after finding no correlation between lipase activity and the degree of crosslinking or the specific surface areas of their alkyl-modified glasses [18]. Our data provide direct evidence for the conformational change hypothesis; even though nature did not design apoMb to function at a hydrophobic interface, apoMb structure was greatly enhanced in alkyl-modified glasses. We propose that the fraction of properly-folded lipase also may be increased by encapsulation in hydrophobic glasses and that the increase in functional lipase is primarily responsible for the higher activity detected in modified glasses relative to unmodified glasses.

It should be noted that hydrophilic organosilane precursors also have been employed to alter the surface chemistry of sol–gel glasses and to enhance the activity of encapsulated enzymes. Some of the common hydrophilic substitutions include glasses modified with aminopropyl [48–51], carboxethyl [48,49], glycidoxypropyl [50,51], and diglycerol or gluconamidyl groups [52–54]. Of special interest with regard to hydrophilic organically-modified glasses, Ackerman and co-workers have reported outstanding enzyme activities following protein adsorption into functionalized mesoporous silica prepared using a non-ionic block copolymer surfactant as the template [48,49]. For example, the specific activity of organophosphorus hydrolase was found to increase by more than two-fold relative to dilute solution when entrapped in a functionalized glass modified with a 2% coverage of carboxethyl groups on the internal silanol wall [48]. Although the superactivity of this enzyme relative to solution measurements is promising [55], the mechanism responsible for the enhanced biocatalysis is not fully understood. Three possibilities include (a) a glass-induced enhancement in native structure following partial unfolding or aggregation of the enzyme during the reported freeze–thaw step, leading to a deceptively low activity in solution, (b) a favourable partitioning of the enzyme substrate into the silica environment such that the activities of the free and immobilized enzymes are not compared at the same effective substrate concentrations, and (c) crowding and confinement effects on the enzyme reaction due to the glass environment, allowing one to measure the true kinetics that would prevail in a living cell [19,20].

If one assumes that the average pore size of the bulk glass also reflects the average size of a protein-occupied pore, then one must conclude that glass-encapsulated apoMb inhabits an extremely confined environment; the calculated pore size is approximately 4 nm in all alkyl-modified glasses, and the largest dimension of the protein is 4.2 nm from the crystal structure [56,57]. It should be noted that the actual pore size of a wet-aged glass before drying may be much larger than 4 nm. Harrell co-workers [58] reported that the pore size obtained by drying with
supercritical CO₂ is approximately 11 nm for a 10% methyl-modified glass, whereas the same glass resulted in an average pore diameter of 3.8 nm when dried from pentane, the same method employed in the current work. Thus, a molecule of apoMb should have ample room to rotate and alter conformation within the pores of a wet-aged glass, but the environment is a crowded one which presumably leads to the striking thermal stability of the protein (Fig. 5).

The mobilities of glass-entrapped proteins have been characterized by fluorescence anisotropy [26,59–62], tryptophan phosphorescence [63], and photobleaching experiments [62]. In addition to apoMb, enhanced conformational stability at elevated temperatures has been observed for many glass-entrapped proteins including α-lactalbumin [22], lysozyme [22,24], human serum albumin [24,26,54], creatine kinase [31], monellin [59], oncomodulin [64], cytochrome c [65], and carbonic anhydrase [66]. The outstanding thermal stability of proteins in organically-modified glasses, as demonstrated here for apoMb, bodes well for the use of such biomaterials in industry, including the development of new nanobiodevices [6,27].

The N₂ adsorption/desorption isotherm curves indicate a change in pore shape upon addition of an alkyl-modifying precursor, whether or not apoMb is present (Fig. 6). In the unmodified glass (100% TMOS), slit-shaped pores are prevalent, whereas bottle-neck-shaped pores are observed for all of the alkyl-modified glasses. Unfortunately, the difference in pore shape fails to explain the observed changes in protein structure because, the same shape (hysteresis classification) was observed for all aerogels made from alkyl-modified glasses, independent of chain length. These experiments were necessary, however, to narrow our search for the critical parameter that links protein structure (and biological activity) to the changes in glass hydrophobicity.

The only parameter reported here that correlates with the observed changes in protein structure is the change in surface morphology of the glass, as monitored by SEM (Fig. 7). The spiral-shaped domains were only observed in the presence of apoMb, and the domain size decreased with increasing hydrophobicity (Fig. 8). This result suggests that apoMb may serve as a template or seed for agglomeration of the silica particles during the gelation process, leading to the unusual morphology. In the case of the unmodified glass, protein encapsulation resulted in the largest domain structures and appeared to form a semi-array pattern. The smaller domain size of the alkyl-modified glasses may reflect a change in the strength of the putative silica–protein interactions that lead to the serpentine microstructure. The presence of similar spiral-shaped particle domains has been observed in hydrophobic aerogel beads made from sodium silicate (water-glass) using trimethylchlorosilane and hexamethyldisilazane as surface-modifying agents [67]. The authors suggested that the observed microstructure was due to their preparation method which involved the dropwise addition of the silica sol into a stirred beaker of n-hexane. In another study, the surface morphology of a poly(vinyl alcohol)-modified glass changed significantly upon encapsulation of the protein urease [68]. The surface mean roughness increased from 0.21 nm in the absence of protein to 2.64 nm in the presence of protein, as measured by AFM. This observation is consistent with the abrupt change in surface morphology reported here upon addition of apoMb (Fig. 7).

To a biochemist, the enhanced structure of apoMb with increasing hydrophobicity of the glass is a dilemma; one would expect a hydrophobic environment to favor the unfolded conformation of a protein because, once unfolded, the hydrophobic core of the folded state should prefer to interact with its hydrophobic surroundings. The results presented in this paper suggest that silica–protein interactions do indeed occur, leading to morphological changes in the host matrix surface, but the mechanism by which glass hydrophobicity enhances the helical content of apoMb remains unresolved.

5. Conclusions

Glass hydrophobicity can have a major influence on the structure of a protein encapsulated in silica matrices. By using a series of modifying reagents that incorporate alkyl groups of increasing chain length, we were able to discern the influence of hydrophobicity on the properties of the glass and on the structure of the protein in a rational manner. The observed changes in protein structure were not linked to any specific physical property of the glass matrix, although we report an intriguing correlation between alkyl chain length and surface morphology. Our results suggest that silica-derived biomaterials can be engineered or optimized for specific applications by selecting the appropriate surface chemistry and that biocompatibility may depend on the proper balance of hydrophilic and hydrophobic groups; in our case, the most hydrophilic surface (unmodified silica) was the least compatible with protein structure. Previous studies were unable to pinpoint why lipase activity is increased in organically-modified glasses, but our work strongly suggests that the enhancement in enzyme activity is due, at least in part, to an increase in the fraction of properly-folded functional enzyme. The favourable effect of hydrophobic glasses on protein conformation may be realized by other encapsulated proteins since our model protein, apoMb, was not expected to benefit from such an environment. Further work is needed to determine the generality of encapsulation effects on the structure and function of proteins in modified glasses of low hydrophobic content.

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References


