Hydrolyzed tetraethylorthosilicate (TEOS) formed SiO2 gel films after being spin-coated onto silicon substrates and heated to temperatures ranging between 200 and 800 °C to effect different degrees of transformation from gel to glass. The index of refraction of the films increased from 1.42 at 25 °C to 1.45 after thermal treatment at 800 °C; the thickness of the films decreased by 29 ± 6%. These thermally pretreated gels were coated with a second film of sol–gel-derived SiO2 doped with cesium chloride. Diffusion of the cesium ion through the undoped SiO2 gels was studied by measurement of the depth profile of cesium by Rutherford backscattering spectroscopy. Increasing the temperature to which the undoped layer had been exposed decreased the extent of its infiltration or diffusion by cesium at room temperature. Diffusion of cesium at high temperature (≥750 °C) was, however, independent of the thermal history of the undoped layer, indicating that the rate at which the silica film transforms from gel to glass is faster than that at which cesium diffuses.

Introduction

The mobility of dopants in SiO2 films made from sol–gel precursors—suspensions of silicate particles produced by polymerization involving hydrolysis and condensation reactions of silicon alkoxides—is important in applications of these films. Studies of diffusion in sol–gel systems have focused on diffusion of liquids into porous gels.
Diffusion of Cesium Ion in SiO₂ Films

The objective of this work was to study the diffusion of cesium ion—chosen as a representative monovalent cation for its availability and ease of analysis by Rutherford backscattering spectrometry (RBS)—in SiO₂ films coated on a silicon substrate. The system examined comprised two superimposed layers. The layer adjacent to the silicon substrate originally contained no cesium. It was prepared by coating a sol–gel solution obtained by hydrolysis of tetraethylorthosilicate (TEOS) onto the substrate and heating the resulting film at temperatures between 200 and 800 °C to effect different degrees of transformation from gel to glass. A second film of gel, obtained by addition of cesium chloride to previously hydrolyzed TEOS, was coated onto the first. The two-layer system—one originally free of cesium and the second containing cesium—was heated. The depth profile of the cesium was subsequently determined by RBS. The focus of the investigation was the rate of diffusion of the cesium ion from the doped into the undoped layer and the influence of the thermal history and physical state of the undoped layer on the rate of this diffusion. We wished to define the mildest heating cycle necessary to transform the undoped gel to a coating that acts as a barrier to diffusion of cesium. Previous work has illustrated the usefulness of RBS for studies of high-temperature diffusion in glass and ceramic films: TiC in WC, Fe in SiO₂, and Cu in carbon.

Several factors affect the rate of diffusion of cations in conventional SiO₂ glasses: the charge of the cation; the amount of water present in the glass; the mobility of the interdiffusing ion or counter ion; the composition and free volume of the glass; the presence and nature of other ions.

Experimental Section

Materials. We purchased ethyl ether from Mallinckrodt, hexane from Fisher Scientific, cesium chloride and tetraethylorthosilicate (TEOS) from Alfa, silicon wafers from Monsanto, and absolute ethanol from USI Chemicals Co. All reagents were used without further purification unless noted.

Preparation of Samples and Conditions for Studying Diffusion. Figure 1 outlines our experimental protocol schematically. We coated a silicon (100) wafer (2-in. diameter) by pipetting enough (undoped) TEOS solution to cover the wafer (3 mL), spinning the substrate at 1500 rpm for 2 min on a...
spin-coater, pipetting more solution onto the coated substrate, and spinning again. The samples were then heated in a programmable furnace at 1 °C/min. held for 2 h at either 200, 400, 600, or 800 °C, and cooled at 1 °C/min.12

In parallel with these experiments, using the same undoped sol-gel suspension of TEOS, we coated a KBr plate and collected IR spectra at each stage of thermal treatment. The melting point of KBr is 774 °C.

Thermogravimetric analysis (TGA) of the unheated coating was carried out on a powder obtained by scraping a film from the silicon substrate with a razor blade.

For studies of diffusion, we recoated the undoped, thermally pretreated SiO2 film with a sol-gel suspension prepared by addition of CsCl to previously hydrolyzed TEOS. RBS established the depth profile of cesium. After initial heating of the sample to 300 °C, the signal due to chloride in the RBS profile decreased to 40 mol % relative to cesium; by 700 °C, it was less than 10%. This observation indicated that chloride ion had volatilized (probably as HCl), and the anionic counterion of the cesium cation had become either OH- or O2- (from water) or Si-O- (from the silicate network).

Ellipsometry of Samples. Measurements were made using a thin-film ellipsometer equipped with a He-Ne laser. Thicknesses and indexes of refraction were obtained by using "A FORTRAN Program for Analysis of Ellipsometric Measurement".25

RBS Analysis. The depth profiles of cesium were obtained with a 2-MeV He+ beam. Depth profiles of cesium and oxygen were calculated by using SPECMAN ANALYSIS.26 The depth profiles presented in Figure 5 were smoothed, and representative profiles from unsmoothed data are presented in Figure 2 for comparison.

Hydrolysis of TEOS. A solution containing TEOS (61 mL, 273 mmol of Si), ethanol (43 mL), doubly distilled water (5 mL), and aq HCl (0.2 mL of 1 M acid) was heated at 60 °C for 1.5 h.27 After cooling the solution to room temperature, an additional 4.0 mL of doubly distilled water and 12 mL of HCl were added to a 100-mL aliquot of the TEOS solution, and the mixture was stored in a freezer (−8 °C), where it was stable (did not gel) for several months. For coating, the suspension was diluted by a factor of 3 with ethanol (0.62 mmol of Si/mL).27

Preparation of CsCl-Doped TEOS. To a suspension of previously hydrolyzed TEOS (10 mL, 6.2 mM in Si) was added a water/ethanol (1.8/5.0 v/v) solution of cesium chloride (0.5 mL, 0.265 mM). The solution was coated immediately.

Results and Discussion

Characterization of SiO2 Films. Ellipsometry of the Supported SiO2 Films as a Function of Their Thermal History. Analysis of the undoped SiO2 coatings by ellipsometry, after heating, gave both their thicknesses and indexes of refraction (Figure 3). Data in Figure 3 represent the average values of six measurements; the error bars, 1 standard deviation.

The index of refraction for the SiO2 gel prepared at 25 °C was 1.42 ± 0.01 (98 ± 1% dense) and 1.45 ± 0.01 (99 ± 1% dense) after heating to 800 °C (the index of refraction for dense SiO2 glass is 1.46).38 The room-temperature values of the index of refraction on our spin-coated samples are consistent with measurements on dense dip-coated samples made from a solution of TEOS prepared by the same procedure that we followed.27 These latter samples had less than 5% porosity,29 a mean pore volume of less than 4 A (inaccessible to nitrogen),29 and a surface area of 0.93 cm2/cm3 of sample.30 Shear stress increases the density of coatings for formulations designed

(26) SPECMAN ANALYSIS, a FORTRAN program based on algorithms from ref 35, was written by Patrick M. Smith, Division of Applied Sciences, Harvard University.

Figure 2. Depth profiles of oxygen (O) and cesium (●) from unsmoothed data (a, best case; b, worst case). The data in a are for the SiO2 undercoat previously heated to 600 °C and in b are for the SiO2 undercoat previously heated to 800 °C. Both samples had no thermal treatment after the cesium chloride doped TEOS was applied.

Figure 3. Plot of the index of refraction (n, O) and thickness (d, ▲) obtained from ellipsometry of the SiO2 films, as a function of the maximum temperature to which they had been heated in air. The temperature profile in the heating is summarized in Figure 1. The lines are included only to guide the eye.

Figure 4. Loss of weight from TEOS-derived SiO2 as a function of temperature under nitrogen. The temperature of the sample increased at 10 °C/min.
to give porous coatings, and we expected the shear stress due to spin-coating would contribute to the formation of dense coatings for α-Si samples as well. The index of refraction did not change significantly until the sample was heated to 800 °C.

The films decreased in thickness by 29 ± 6% on heating to the highest temperature studied (800 °C). This decrease in thickness represents a loss of volatiles (water, organics, and possibly low molecular weight silicon containing oligomers). The loss of volatile organosilicon oligomers would allow for the loss of material without a large change in the index of refraction and would be consistent with the unexpectedly small change in the index of refraction over this temperature range.

Thermogravimetric Analysis (TGA) of the SiO₂ Coating. Figure 4 shows the TGA of a gel, heated under nitrogen at 10 °C/min. The sample had lost 6.2 ± 1.5% of its weight by 200 °C, and an additional 8.5 ± 1.0% by 800 °C. The rate of loss of weight is faster between 25 and 200 °C than between 200 and 800 °C. These observations correlate with processes important in the gel-to-glass transition: at 400 °C and below, surface and physisorbed water and organic groups are lost; up to 700 °C, a gradual loss of water results from further condensation of the Si-OH groups. The loss of volatile organosilicon oligomers could occur throughout the thermal treatment until condensation reactions incorporated them into the gel network.

The loss of weight of the sample of 15 ± 2%, as measured by TGA, is not consistent with a decrease in film thickness of 29 ± 6% with an accompanying decrease in porosity (or increase in density) of 1.5 ± 1.0%. These changes in thickness and density would require a loss of weight of 28 ± 6%.

IR Analysis of SiO₂ Coatings as a Function of Thermal History. Figure 5 presents IR spectra, collected at room temperature and after each thermal cycle, of an SiO₂ coating on a KBr plate. Samples were heated at 1 °C/min, held for 2 h at 200, 400, or 600 °C, and cooled at 1 °C/min. Below 200 °C the decrease in intensity of the H₂O and Si-OH absorbances is consistent with loss of physisorbed water. When the sample is heated from 200 to 800 °C the decrease in the Si-OH absorbances is consistent with further condensation of Si-OH groups with accompanying loss of H₂O.

Measurement of the Depth Profile of Cesium in SiO₂ by RBS. To examine diffusion of cesium, we heated samples to 750–800 °C under argon. At these temperatures, two processes could, in principle, occur concurrently: transformation (or further transformation, for the undoped SiO₂ layer) of the silica layer from gel to glass, and diffusion of cesium. By studying the relative rates of migration of cesium into gels that had been carried to different stages of the gel-to-glass transformation by prior heating, we hoped to determine, at least qualitatively, whether diffusion...
To induce detectable diffusion of cesium, the samples were heated under argon by using the same profile as shown in Figure 1: heating and cooling rates of 1 °C/min, and 2 h at the holding temperature (750 or 800 °C). For calibration, one sample in each set was not heated. We note from the depth profiles of cesium in these samples that the thermal history of the undoped silica layer influences the extent to which cesium infiltrates accessible pores (<5% total porosity) and/or diffuses before heating the two-film sample. For example, at a depth of 180 nm (20 nm above the interface between the undoped silica film and the silicon substrate), the cesium content is approximately 0.1 × 10^{20} atoms/cm^3 for the sample treated initially at 200 °C for 2 h, 0.1 × 10^{20} atoms/cm^3 at 400 °C, and less than 0.05 × 10^{20} atoms/cm^3 for samples taken to both 600 and 800 °C.

Estimation of the Apparent Diffusion Coefficient of Cesium in SiO_2 Gels. We estimated the apparent diffusion coefficient, D_{app} (cm^2/s),\textsuperscript{(37)} of cesium in the SiO_2 gel using the thin-film solution.\textsuperscript{(38)} Figure 8 presents the calculated apparent diffusion coefficients. Those for the samples that were previously heated to 200 and 400 °C represent an effective diffusion coefficient, reflecting an aggregate of the room-temperature infiltration of cesium and the diffusion of cesium at elevated temperature. There is no significant difference in the apparent diffusion coefficients calculated for samples previously heated to 200, 400, 600, and 800 °C; they are all in the order of magnitude of 10^{-15} cm^2/s.

(37) For a film with a quantity of solute, a, diffusing into a material from the surface, the thin-film solution (Crank, J. The Mathematics of Diffusion; Clarendon: Oxford, 1975; pp 11-13) is

\[ c = \frac{a}{x \cdot D} \exp\left(-\frac{x^2}{4Dt}\right) \]

To extract the diffusion coefficient, D, ln c is plotted versus r^2. The slope of the line is (4Dt)^{1/2}. The thin-film solution is valid for diffusion into an undoped gel, provided that the gel can be considered infinite; no loss of material occurs through the other side of the film at the SiO_2 gel/Si interface.

(38) The thin-film solution is a reasonable assumption for the estimation of an apparent diffusion coefficient for our system, where a thin film of solute is placed on a “semifinite” and solute-free medium. The boundary condition that the flux at the surface of the thin-film solution is zero satisfies the thin-film solution. The underlying assumptions here are that the diffusion coefficient is not a function of the concentration of cesium and that the driving force to diffusion, the gradient in the chemical potential, is the same as the gradient in concentration—a measurable quantity. The SiO_2 coating can be considered an infinite medium only if it passes the “leak test.” Following Shewmon (Shewmon, P. G. Diffusion in Solids; D. Williams Book Co.: Jenks, OK, 1983; p 9): “a short bar can be considered infinite if the quantity of solute which would lie outside its length in a truly infinite bar is an insignificant portion of the total solute present.” With 0.1% as the insignificant portion of material to “leak” out, eq 1 is the ratio of the solute that has

\[ 10^{-3} = \left[ \int_0^D \int_0^x e^{-x^2/4Dt} dx \right] \left[ \int_0^D \int_0^x e^{-y^2/4Dt} dy \right] \]

(1)

diffused beyond the distance x to the total content of solute in the bar, where D is the diffusion coefficient (cm^2/s), t is time (s), and x is distance (cm). The solution of x is \( x = \sqrt{4Dt} \). For our system, with an apparent diffusion coefficient of 1.4 × 10^{-15} cm^2/s at an annealing time of 200 s, there is less than 0.1% diffusion beyond ~4.2 × 10^{-15} cm (400 Å). Our SiO_2 films extend to approximately 5000 Å from the surface (longer than the minimum length for an “infinite” medium for the given diffusion coefficient and annealing time), and the assumption that the thin-film solution holds is correct. Qualitatively there was little or no diffusion of cesium into silica (Figure 7).

The Si/SiO_2 interface was fixed—no detectable oxidation occurred. Samples were heated in argon to effect diffusion of the cesium at elevated temperatures. There was no significant detectable increase in the depth profiles of oxygen of the films after thermal treatment (see Figure 7, bottom profile).

Since cesium is a charged species, it must either ion exchange with another cationic species, for example, H_+ on Si-OH, or diffuse with an anion. Here we cannot examine these processes directly, because we cannot measure H_+ by RBS.
Diffusion of Cesium Ion in SiO$_2$ Films

Figure 8. Plot of $D_{Cs}$ ($\times 10^{-4}$ cm$^2$/s) at 750 and 800°C for samples previously heated to 200°C (●), 400°C (●), 600°C (●), and 800°C (▲). The points are shown displaced from the nominal temperatures to avoid overlap.

Conclusions

Increasing the temperature at which the undoped layer was treated before applying the cesium-containing gel decreased the extent of infiltration into accessible pores and/or diffusion of cesium into the undoped gel at room temperature. This observation indicates that the physical state of the gel (probably its SiOH content) was important in determining ionic mobility. Cesium diffused at room temperature into undoped films that had been heated to 200 and 400°C but not into those heated to 600 or 800°C; the extent of diffusion of cesium was greatest for the film heated only to 200°C. Thus, it seems to be necessary to have heated the undoped film to approximately 600°C to prevent infiltration and/or diffusion of cesium into it when it is overcoated with a cesium chloride containing gel layer at room temperature.

The rate of diffusion of cesium in the undoped gel correlates qualitatively with the concentration of SiOH groups in this gel, as measured by IR spectroscopy.$^{39}$ At room temperature, the gels retained chloride ion, and part of the relatively high room-temperature mobility of cesium ion into the undoped gels that had not been heated to temperatures greater than 400°C may be due to the presence of both cation (cesium) and anion (chloride) in this system in diffusible form. Since the chloride ion rapidly disappears (to 37% of the concentration of cesium by 300°C and approximately to 10% by 700°C) from the gel, we assume the counterion for most of the cesium ion at high temperature is either SiO$^-$ or O$^2-$ at temperatures above 300°C.

Cesium diffusion was negligible at $T < 750°C$. When detectable migration of cesium did begin to occur ($T \geq 750°C$), its rate was independent of the thermal history of the undoped layer over the range of temperatures used in treating these layers (200–800°C). The extent of diffusion of cesium at elevated temperatures did not depend on the temperature of thermal pretreatment of the undoped gel. Thus, when the sample reached a temperature at which detectable diffusion of cesium occurred ($T > 750°C$), the cesium “saw” silica film previously heated to any temperature between 200 and 800°C as having the same physical state. We infer from this observation that the rate at which the silica film transforms from gel to glass is fast relative to cesium migration, at the high temperatures required for this migration to be detectable by RBS.

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Registry No. Cs$^+$, 18459-37-5; vitreous silica, 60676-86-0.

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$^{39}$ The SiOH content in a glass can strongly influence its properties. Incorporation of 0.1 wt % water in SiO$_2$ reduces its viscosity by 3 orders of magnitude at 1000°C. The Si–OH groups may bind Cs$^+$ (plausibly as SiO–Cs$^+$) and increase its mobility. Doremus, R. H. Glass Science; Wiley: New York, 1973; pp 170–172.