Diffusion of hydrogen atoms in silicon layers deposited from molecular beams on dielectric substrates

Kirill V. Chizh\textsuperscript{a,*}, Larisa V. Arapkina\textsuperscript{a}, Dmitry B. Stavrovsky\textsuperscript{a}, Peter I. Gaiduk\textsuperscript{b}, Vladimir A. Yuryev\textsuperscript{a,*}

\textsuperscript{a}Prokhorov General Physics Institute of the Russian Academy of Sciences, 38 Vavilov Street, 119991 Moscow, Russia
\textsuperscript{b}Department of Physical Electronics and Nanotechnology, Belarusian State University, 4 Prospekt Nezavisimosti, 220030 Minsk, Belarus

Abstract

In the paper, the processes occurring during low-temperature growth of non-hydrogenated amorphous Si and polycrystalline Si films on multilayer Si\textsubscript{3}N\textsubscript{4}/SiO\textsubscript{2}/c-Si substrates from molecular beams under conditions of ultrahigh vacuum are studied in detail. Diffusion of hydrogen atoms from a dielectric layer into the growing film is shown to accompany the growth of a silicon film on a Si\textsubscript{3}N\textsubscript{4} layer deposited by CVD or on a SiO\textsubscript{2} layer obtained by thermal oxidation of a silicon wafer. The process of hydrogen migration from the dielectric substrates into the silicon film is studied using FTIR spectroscopy. The reduction of IR absorption at the bands related to the N–H bonds vibrations and the increase of IR absorption at the bands relating to the Si–N bonds vibrations in IR spectra demonstrate that hydrogen atoms leave Si\textsubscript{3}N\textsubscript{4} layer during Si deposition from a molecular beam. The absorption band assigned to the valence vibrations of the Si–H bond at $\sim$ 2100 cm\textsuperscript{-1} emerging in IR spectra obtained at samples deposited both on Si\textsubscript{3}N\textsubscript{4} and SiO\textsubscript{2} layers indicates the accumulation of hydrogen atoms in silicon films. The difference in chemical potentials of hydrogen atoms in the dielectric layer and the silicon film explains the transfer of hydrogen atoms from the Si\textsubscript{3}N\textsubscript{4}.

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\textsuperscript{Corresponding author}

Email addresses: chizh@kapella.gpi.ru (Kirill V. Chizh), arapkina@kapella.gpi.ru (Larisa V. Arapkina), stavr@nsc.gpi.ru (Dmitry B. Stavrovsky), gaiduk@bsu.by (Peter I. Gaiduk), vyuryev@kapella.gpi.ru (Vladimir A. Yuryev)

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or SiO$_2$ layer into the growing silicon film.

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1. Introduction

Presently, MEMS technology is a rapidly developing branch of science, one of the promising areas of which is the production of IR sensors based on amorphous (α-Si) or polycrystalline (poly-Si) silicon films formed on Si3N4/SiO2 dielectric layers. The properties of α-Si and poly-Si films are determined both by the technological modes of deposition and by the properties of the substrate, primarily, the upper layer of silicon nitride of a complex Si3N4/SiO2/c-Si substrate. The most commonly used methods of depositing Si3N4 are LPCVD with thermal and PECVD with plasma-chemical decomposition of nitrogen-containing components. Application of any of these methods leads to the production of films saturated with hydrogen atoms. The concentration of hydrogen atoms in PECVD films can reach 25%, and in LPCVD films 8% [4, 5, 6, 7]. Such hydrogen concentrations can significantly affect the properties of α-Si and poly-Si films grown on the surface of a complex Si3N4/SiO2/c-Si substrate due to desorption of hydrogen from the Si3N4 film during and after the deposition of silicon layers. The protective layer of Si3N4 is known to facilitate a decrease in the recombination of non-equilibrium charge carriers and can reduce the concentration of the majority carriers in the near-surface region of the Si layer due to the formation of a built-in positive charge [8]. Hydrogen atoms are able to diffuse from the Si3N4 layer and passivate the broken bonds at the interface with the Si layer, thereby improving the electrophysical properties of the material [9, 10, 11, 12, 13]. At present, protective coatings of Si3N4 deposited by low-temperature PECVD are widely used as a source of hydrogen atoms to prevent degradation of electrophysical properties of solar cells based on single crystalline (c-Si), multicrystalline and amorphous silicon [14]. However, the study of hydrogen diffusion in such structures is complicated by the fact that the Si3N4 and α-Si (poly-Si) layers are hydrogenated and the hydrogen atoms can freely move between the layers.

In this paper, the processes occurring during low-temperature growth from molecular beams under conditions of ultrahigh vacuum (UHV) of non-hydrogenated α-Si and poly-Si films on a multilayer Si3N4/SiO2/c-Si substrate are studied in detail. The Si3N4 layer is the source of hydrogen atoms in our experiment.
Previously, the diffusion of hydrogen atoms into α-Si and poly-Si layers from an external source was studied, e.g., in the works [15, 16, 17, 18, 19]. At low process temperatures, diffusion of hydrogen atoms with an activation energy from $\sim 0.2$ to $\sim 0.4$ eV was observed in those works. Hydrogen atoms first passivated the broken silicon bonds, and when the temperature was raised above 300°C, a rupture in the weak Si–Si bonds was observed. The study of the saturation of silicon layers with hydrogen atoms during deposition of Si$_3$N$_4$ layers using PECVD showed that hydrogen atoms penetrated into c-Si to a depth from 10 to 20 nm at temperatures from 500 to 900°C, and pre-treatment of the c-Si surface in NH$_3$ plasma increased the concentration of hydrogen atoms passing into c-Si [18, 19].

2. Sample preparation, experimental methods and equipment

2.1. Samples

Experimental samples were prepared by depositing Si layers on different substrates. There were three types of substrates used in the experiments: Type N, Type O and Type S.

Type N substrates, Si$_3$N$_4$/SiO$_2$/Si(100), were produced as follows. First, finished on both sides 450 µm thick (100)-oriented Czochralski grown (CZ) boron doped silicon wafers ($p$-type, $\rho = 12$ Ω cm) were cleaned using the RCA process [20, 21, 22]. Then, 530 nm thick SiO$_2$ layers were formed on both sides of silicon wafers using thermal oxidation. This process was performed in three consecutive steps: at first, the wafers were treated in dry oxygen for 30 min, then in wet oxygen for 60 min, and finally, in dry oxygen for 30 min. A 175 nm thick Si$_3$N$_4$ layer was deposited on both sides of the oxidized wafers by pyrolysis of a monosilane-ammonia mixture at the temperature of 750 °C for 60 min [1, 2, 23].

On Type O substrates, SiO$_2$/Si(111), only 700 nm thick SiO$_2$ film was formed by thermal oxidation of CZ Si(111):P ($n$-type, $\rho = 100$ Ω cm) wafers. Type S substrates were $p$-type CZ Si(100):B wafers, $\rho = 12$ Ω cm.

Silicon films were deposited on the substrates from molecular beams (MB) in the ultra-high vacuum EVA 32 (Riber) molecular-beam epitaxy (MBE) chamber using a
solid source with Si evaporation by an electron beam from a high resistivity float zone ingot.

The MBE chamber was evacuated down to about $10^{-11}$ Torr before the processes. The pressure did not exceed $5 \times 10^{-9}$ Torr during Si deposition. The deposition rate and coverage were measured using the Inficon XTC751-001-G1 (Leybold-Heraeus) film thickness monitor equipped with the graduated in advance quartz sensors installed in the MBE chamber. Si deposition rate was $\sim 0.3 \text{ Å/c}$; the film growth temperature was varied from 30 to 650°C in different processes for samples produced on Type N and Type O substrates. The thickness of the Si layer was 200 nm.

Before depositing of silicon nitride or silicon, the substrates of all types were cleaned using an identical process that we routinely apply for obtaining clean Si surfaces [24, 25, 26]. At first, they were washed in the ammonia-peroxide solution ($\text{NH}_4\text{OH} (27\%): \text{H}_2\text{O}_2 (30\%): \text{H}_2\text{O} [1:1:3]$, boiling for 10 min), then rinsed in deionized water, boiled in high-purity isopropyl alcohol ([C$_3$H$_7$OH] $> 99.8$ wt%, $T \approx 70°C$), and dried in the isopropyl alcohol vapor (for 10 min) and the clean air. Additionally, before moving into the MBE chamber, the substrates were annealed at 600°C at the residual gas pressure of less than $5 \times 10^{-9}$ Torr in the preliminary annealing chamber for 6 hours [1, 2, 23, 27].

Type S substrates were additionally deoxidized in the MBE chamber at the temperature of 800°C at a flux of Si atoms. The Si deposition rate during the deoxidation process was $< 0.1 \text{ Å/s}$; the measured Si coverage was about 30 Å [24]. The pressure in the MBE chamber increased to nearly $2 \times 10^{-9}$ Torr at most during this process. RH20 reflection high-energy electron diffraction (RHEED) tool (Staib Instruments) installed in the MBE chamber was used for deoxidation monitoring.

The growth of Si films on Type S substrates was conducted in two stages. First, a 100 nm thick Si layer was deposited at 650°C on the wafer surface purified of silicon oxide; then, after sample cooling to the room temperature, a 200 nm thick Si layer was deposited at this temperature.

Samples were heated from the rear side using tantalum radiators in both preliminary annealing and MBE chambers. The temperature was monitored with chromel-alumel and tungsten-rhenium thermocouples of the heaters in the preliminary annealing and
MBE chambers, respectively. The thermocouples were mounted in vacuum near the rear side of the samples. They were in situ graduated beforehand against the IMPAC IS12-Si infrared pyrometer (LumaSense Technologies) that measured the sample temperature through MBE chamber windows (for \( T > 300^\circ C \); the graduation curves were extrapolated to the room temperature for \( T < 300^\circ C \)).

The composition of residual atmosphere in the MBE chamber was monitored using the RGA-200 residual gas analyzer (Stanford Research Systems) before and during the deposition process.

2.2. FTIR analysis

Fourier transform IR (FTIR) transmission and reflection spectra of the samples were explored using a vacuum IFS-66v/S spectrometer (Bruker). The spectral resolution was 10 cm\(^{-1}\) that enabled both an adequate recording of all spectral features and filtering away of the high-frequency components related to light interference in silicon wafers. An opaque golden mirror was used as reference sample during recording of reflectance spectra. The instrument was evacuated to the residual air pressure of 2 mbar during spectra recording that enabled a considerable reduction of spectral interferences associated with carbon dioxide and water vapor. Direct analysis of the obtained spectra was hampered by interference of probe radiation in thin layers of materials on silicon wafers. In analyzing the spectra, the transmission spectrum and the reflection spectrum of each sample were summed. Such procedure of spectra processing has enabled the reduction of amplitude of baseline variations arisen due to the interference. As a result, it has become possible to observe and analyze relatively weak absorption bands.

Additionally, the peak wavenumbers and relative intensities of spectral components were analyzed using the deconvolution procedure. We used Gaussian functions for fitting. Spectra to be compared were normalized to the intensity of the strongest band assigned to the vibrations of the Si–O bond.

3. Results and their interpretation

Both in-situ structural investigations of the surface of the growing Si layer using RHEED and ex-situ examinations of the deposited Si layer using high resolution trans-
mission electron microscopy have shown that an amorphous silicon layer forms on Type N substrates at the temperature range from 30 to 420°C [23, 28]. Transition from the amorphous structure of the growing film to the polycrystalline one after the film thickness reaches a critical value—i.e. the formation of a layered poly-Si/amorphous Si film—is observed at the temperature interval from 420 to 500°C. Finally, only polycrystalline Si layers form at the higher temperatures (for details, see Ref. [23]).

Figure 1 demonstrates IR absorbance spectra in the range from 650 to 1350 cm$^{-1}$ for the samples grown from MB at different temperatures of Type N substrates. The spectra obtained from a Type N substrate and from samples grown at three different temperatures are shown in Figure 1a (the spectra are normalized to the maximum of the highest peak at 1090 cm$^{-1}$). The spectra are characterized by the presence of two wide bands peaked at 825 and 1090 cm$^{-1}$, and the latter, the strongest of them assigned to the Si–O bond vibrations, exhibits a wide shoulder on the side of large wavenumbers spreading to nearly 1300 cm$^{-1}$ that is associated to the vibrations of Si–O and N–H bonds [29, 30, 31]. The deconvolution of these bands using Gaussian functions and analysis of vibrational frequencies presented in the panel (b) demonstrates that they are composed of eight spectral components peaked at about 735, 805, 875, 965, 1075, 1100, 1150 and 1190 cm$^{-1}$. The peaks at 1150 and 1190 cm$^{-1}$ are assigned to the vibrations of the N–H and Si–O bonds, respectively [29, 30, 31]. Two strong peaks around 1090 cm$^{-1}$ are assigned to the stretching vibrations of the Si–O bond and a peak around 800 cm$^{-1}$ is attributed to the deformation vibrations of the Si–O bond [32, 33, 34]. Absorption peak at 965 cm$^{-1}$ likely relates to the fundamental antisymmetric stretching vibrations of the $\text{Si}_2\text{O}_6^{4-}$ silicate units [35]. A weak absorption peak at 735 cm$^{-1}$ is also assigned to the vibrations of the Si–O bond, whereas the band around 875 cm$^{-1}$ relates to the absorption of radiation by the Si–N bond [31, 36].

Comparison of spectral peaks plotted in Figure 1c shows that decrease in the intensity of the absorption band relating to the vibrations of the N–H bond and simultaneous increase in the intensity of the absorption band attributed to the vibrations of the Si–N bond occur due to the silicon layer growth.

The peaks associated with the N–H bond are seen to virtually match for the samples produced at the temperatures of 300, 450 and 650°C; for the samples produced at the
temperatures of 30 and 160°C these peaks also coincide but go some higher than those from the higher temperature group. Only the peak recorded at Type N substrate goes separately being the most intense. On the contrary, the peak related to the vibrations of the Si–N bond is much less intense in the spectrum obtained at Type N substrates than those recorded at samples with deposited Si films regardless their processing temperature; and the latter are identical within the standard errors of their parameters. This observation shows that the additional Si–N bonds begin emerging in the silicon nitride layer as soon as the Si film starts forming, even at the temperature close to the room temperature. This process may be caused both by hydrogen diffusion from the interface of Si nitride and Si into the growing Si film and by Si accumulation at the interface. Breaking rate of the N–H bonds initially low at low temperatures likely becomes higher at elevated temperatures.

Figure 2 presents the sums of IR transmittance and reflectance spectra obtained in the wavenumber range from 1900 to 3500 cm\(^{-1}\) for the samples grown from MB on Type N substrates at different temperatures of silicon deposition. Radiation absorption band at about 3300 cm\(^{-1}\) is known to be related to the valence vibrations of the N–H bonds [31, 36]. The intensity of this peak decreases as the silicon deposition temperature increases. For the samples grown at low temperatures, 30 and 160°C, two absorption lines emerge in the wavenumber interval from 2000 to 2300 cm\(^{-1}\): the line peaked at about 2100 cm\(^{-1}\) is connected with the Si–H bond vibration; that peaked at ca. 2250 cm\(^{-1}\) presumably may also be assigned to the Si–H bond vibration.

Figure 3 depicts the sums of IR transmittance and reflectance spectra recorded in the wavenumber interval from 1900 to 3500 cm\(^{-1}\) for the samples with silicon layers deposited at 30°C from MB on different types of substrates. It is seen that the bands peaked at \(~2100\) and \(~2250\) cm\(^{-1}\) are present in the spectra obtained at the samples grown on Type O substrates, but they are absent in the spectra obtained at the samples with Type S substrates.

Figure 4a demonstrates IR absorbance spectra recorded in the range 400 to 670 cm\(^{-1}\). The absorbance curves are normalized to the maximum of the peak at \(~450\) cm\(^{-1}\) corresponding to the vibrations of the Si–O bond. Two characteristic bands are observed
in this spectral region: a band peaked at $\sim 460 \text{ cm}^{-1}$ and that peaked at $\sim 615 \text{ cm}^{-1}$.

The strong wide band with a maximum at $460 \text{ cm}^{-1}$ is formed by the superposition of several absorption bands related to the vibrations of the Si–O bond with the maximums at about 400, 425, 440, 460 and 510 cm$^{-1}$ (Figure 4b); the peak with the maximum around 500 cm$^{-1}$ is assigned to the vibrations of the Si–N bond. As the silicon deposition temperature is increased, the intensity of the absorption line at $\sim 500 \text{ cm}^{-1}$ decreases (Figure 4b). Such behavior of this absorption band is associated in the literature \[37\] with the decrease of H atoms content in the Si$_3$N$_4$ layer and hence with the reduction of the number density of the Si–H bonds in it. This, in turn, increases an order in the Si$_3$N$_4$ layer and decreases the absorption at $\sim 500 \text{ cm}^{-1}$ since, the more disordered silicon nitride, the higher the absorption at this line. It should be noticed, however, that the peak at ca. 500 cm$^{-1}$ demonstrates significant narrowing after poly-Si film depositing at the temperatures higher than 450°C, yet keeping the general trend to decrease in intensity with the growing Si deposition temperature.

The absorption band at $\sim 500 \text{ cm}^{-1}$ was not observed in the samples grown at 30°C on Type O and Type S substrates.

4. Discussion

As it is seen from the above data of our investigation of silicon layers grown on different substrates at different deposition temperatures using different deposition methods, the main changes are observed at the bands assigned to the vibrations of the N–H, Si–H and Si–N bonds. All the Si layer deposition processes used in this work were carried out at the temperatures lower than the deposition temperatures of the Si$_3$N$_4$ and SO$_2$ layers. The preliminary UHV thermal treatment at 600°C did not change the absorption spectra of the original substrates. Previously, in CVD-deposited Si$_3$N$_4$ films, the reduction of the hydrogen atoms content was observed only after annealing at the temperatures that exceeded their deposition temperature \[38\]. Thus, we can state that the observed changes in the IR absorption spectra are due to Si deposition.

We have found that the reduction of IR absorption at the lines related to the valence vibrations of the N–H bond at $\sim 3300 \text{ cm}^{-1}$ and the increase in the intensity of the Si–N
bond valence vibrations at ~ 875 cm\(^{-1}\) is observed regardless of the method of depositing the Si layer on a Type N substrate at the growth temperature in the range from 30 to 650°C. Additionally, the absorption band intensity of the Si–N bond vibration at ~ 500 cm\(^{-1}\) decreases. These changes in the absorption spectra are highly likely result from the reduction of the hydrogen content in the \(\text{Si}_3\text{N}_4\) layer: hydrogen atoms diffuse into the growing amorphous or polycrystalline Si layer. They either remain in the growing Si film or desorb from it depending on the deposition temperature. If the latter is 30 or 160°C that is below the hydrogen desorption temperature \[39\], the absorption band connected with the valence vibrations of the Si–H bond at ~ 2100 cm\(^{-1}\) appears in the spectra. Emergence of the band at ~ 2250 cm\(^{-1}\) is usually connected in the literature with H–Si(O\(_3\)) or Si–O–Si–H structures forming in SiO\(_x\):H layers \[40, 41, 42, 43\]. In addition to the band of valence vibrations at ~ 2250 cm\(^{-1}\), a band of deformation vibrations of those structures at ~ 880 cm\(^{-1}\) is present in the samples studied in the cited articles. In the current study, the absorption band assigned to the valence vibrations of the Si–N bond is registered in the wavenumber range around 875 cm\(^{-1}\). The increase in the absorption peak intensity at this spectral range does not correspond with the change in the intensity of the band at ~ 2250 cm\(^{-1}\). The accumulation of hydrogen atoms in the amorphous Si film growing at 30°C on a Type O substrate also manifests itself in the appearance of the absorption band at ~ 2100 cm\(^{-1}\); the band at ~ 2250 cm\(^{-1}\) also appears in this case, but no increase in the absorption intensity at ~ 875 cm\(^{-1}\) is observed. Thus, the origin of the absorption band at ~ 2250 cm\(^{-1}\) is presumably also related to the vibrations of the Si–H bond since the conditions for its emergence are the same as for the band peaked at ~ 2100 cm\(^{-1}\). During Si deposition on a Type O substrate, the SiO\(_2\) layer is a source of hydrogen atoms, which are known to be present in pretty high concentration in SiO\(_2\) layers obtained using thermal oxidation. These bands were absent in the IR absorption spectra obtained at the reference sample grown at 30°C on a Type S substrate.

The observed decrease of intensity of the absorption band peaked at ~ 500 cm\(^{-1}\) may also be related to the reduction of hydrogen atoms concentration in the \(\text{Si}_3\text{N}_4\) layer. For instance, in Ref. \[37\], the activity of the band at ~ 495 cm\(^{-1}\) has been connected with the N–Si–H vibrations in a PECVD \(\text{Si}_3\text{N}_4\) layer where hydrogen atoms...
form Si–H and N–H bonds and the ratio of concentrations of these bonds depends on the composition of the reacting mixture. According to that work, reduction of the N–Si–H bonds quantity due to breaking Si–H bonds results in absorption lowering at the band at \( \sim 495 \text{ cm}^{-1} \) and in order improvement in the arrangement of atoms. It may be supposed following that work that the observed change in the intensity of the band at \( \sim 500 \text{ cm}^{-1} \) is related to the order improvement in the Si\(_3\)N\(_4\) layer. In the case of CVD-deposited Si\(_3\)N\(_4\), hydrogen atoms form N–H bonds since they have the highest binding energy. The reduction of the concentration of hydrogen atoms and, as a consequence, of the quantity of the Si–N–H bonds leads to the enhancement of order in the arrangement of atoms and increasing of the Si–N bonds number. As a result, the IR absorption at the band around \( \sim 500 \text{ cm}^{-1} \) decreases.

Let us consider a possible mechanism for the diffusion of hydrogen atoms from the Si\(_3\)N\(_4\) or SiO\(_2\) dielectric layer into the growing Si layer. This process has an important feature: it occurs even at room temperature of Si deposition. Diffusion at such a low temperature obviously is not of an activation nature. The diffusion activation energy of hydrogen atoms, e.g., in layers of amorphous Si is about 1.5 eV [39]; the energy required for breaking the N–H bond is \( \sim 4 \text{ eV} \); the energy necessary for breaking the Si–H bond is \( \sim 3 \text{ eV} \) [44]. In the discussed case, a mechanism of hydrogen atoms diffusion may be described using a model considered in Refs. [45, 46, 47, 48, 49, 50]. The model is based on an assumption that the diffusion of hydrogen atoms in amorphous silicon is limited by capturing of hydrogen atoms by dangling bonds of Si atoms with formation of the Si–H bonds, which are deep traps. Hydrogen atoms move through interstitial states that is accompanied by the rupture of the weak Si–Si bonds. A hydrogen atom diffuses until a Si–H bond is again formed. A concept of a hydrogen atom chemical potential (\( \mu_H \)) is introduced in the model, the position of which is controlled by the concentration of hydrogen atoms in a layer. For the diffusion to start, a hydrogen atom should transit from an energy level corresponding to the Si–H bond onto a so-called transport level (to the interstitial position). The \( \mu_H \) level is located between the level of deep traps and the transport level. The greater the concentration of hydrogen atoms, the higher the \( \mu_H \) level. When the layers with different levels of hydrogen atoms are combined, hydrogen atoms will diffuse from the region with a high level of \( \mu_H \) into
the region with a low level. In our case, the content of hydrogen atoms is higher in dielectric layers of Si$_3$N$_4$ or SiO$_2$. The diffusion process of hydrogen atoms from the Si$_3$N$_4$ (or SiO$_2$) layer into the growing Si layer is possible until the chemical potentials of hydrogen atoms in them are equalized. The thickness of the Si$_3$N$_4$ layer, in which the concentration of hydrogen atoms decreases, and the depth of penetration of hydrogen atoms into the growing Si layer will expand as the deposition temperature increases. The probability of desorption of hydrogen atoms from the growth surface of the Si layer will also increase. As a result, at low growth temperatures, we observe the bands corresponding to the Si–H bonds in the absorption spectra, and as the temperature is raised, hydrogen atoms start to diffuse through the Si layer onto its surface and desorb [48]. This explains the observed reduction of IR radiation absorption at the spectral bands related to the Si–H bonds at high silicon deposition temperatures.

It should be noted also that if we take into the account a mechanical stress usually present in the studied layered structures, especially in the Si layer, we should conclude that the stress decreasing the atomic bond breaking barrier in the Si–Si bonds [51] will increase the hop probability of hydrogen atom during diffusion, which, as mentioned above, is accompanied with the Si–Si bond rupture. Thus, the weakening of the Si–Si bonds will accelerate the diffusion. Even at room temperature, easier Si–Si bond breaking due to local energy fluctuations will stimulate the diffusion of hydrogen atoms from a dielectric substrate into the growing silicon layer. The same assumption may be made about the N–H and Si–H bonds in the dielectric substrates: the loaded bonds should be easier to rupture and hydrogen atoms should be easier to release and hence easier to start their diffusion into the silicon film. The internal stress, due to, e.g., difference in temperature coefficients of expansion of layers, and hence atomic bond loading is certainly extremely temperature sensitive. The local energy fluctuations may be formally considered as a sort of quasi-particles that take part in the diffusion process by breaking atomic bonds. The density and the magnitude of the local energy fluctuations, as well as their movement in solid layers, also critically depend on temperature. So, we should consider the mechanical strain in the layers and the local energy fluctuations as additional factor influencing the process of the hydrogen atoms diffusion in silicon films deposited on dielectric substrates.
5. Conclusion

Concluding the article we emphasize its main inference: the growth of a silicon film on a Si₃N₄ layer deposited using CVD or on a SiO₂ layer formed using thermal oxidation of a silicon substrate is accompanied by the diffusion of hydrogen atoms from the dielectric layer into the growing film. In a wide temperature range, the intensity of the IR absorption bands related to the vibrations of the N–H bonds reduces whereas the intensity of the IR absorption bands connected with the Si–N bonds vibrations increases that demonstrates the escape of hydrogen atoms from a Si₃N₄ layer. At Si deposition temperatures lower than the temperature of hydrogen atoms desorption from the Si surface, the absorption band assigned to the valence vibrations of the Si–H bond peaked at ~ 2100 cm⁻¹ emerges in IR spectra obtained at samples deposited both on Si₃N₄ and SiO₂ dielectric layers that demonstrates the accumulation of hydrogen atoms in silicon films.

The experimental results may be explained using a model of hydrogen atoms diffusion proposed in Ref. [43]. The migration of hydrogen atoms from the Si₃N₄ or SiO₂ layer into the growing silicon film is due to the difference in chemical potentials of hydrogen atoms in the dielectric layer and the silicon film.

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Figure 1: FTIR absorbance spectra for the samples grown from MB at different temperatures on Type N substrates: (a) spectra obtained at a Type N substrate and at samples grown at three different temperatures indicated in the plot, the spectra are normalized to the maximum of the peak at 1090 cm$^{-1}$; (b) deconvolution of the absorbance spectral band in the range from 700 to 1350 cm$^{-1}$ (the sample grown at 30°C): the band is composed of eight Gaussian bands assigned to vibrations of the Si–O, Si–N and N–H bonds; (c) evolution of the bands assigned to the vibrations of the Si–N and N–H bonds in the spectra obtained at a Type N substrate and at samples grown at three different temperatures shown in the plot.
Figure 2: Sums of FTIR transmittance and reflectance spectra for the samples grown from MB on Type N substrates at different temperatures of silicon deposition: (1) a Type N substrate, (2) 30°C, (3) 160°C, (4) 300°C, and (5) 650°C; the bands corresponding to the N–H and Si–H bonds vibrations are shown with vertical lines.
Figure 3: Sums of FTIR transmittance and reflectance spectra for the samples grown from MB at 30°C on (1) Type S, (2) Type O, and (3) Type N substrates; the bands related to the Si–H bond vibrations are shown with vertical lines.
Figure 4: (a) FTIR absorbance spectra for the samples grown from MB at different silicon deposition temperatures on Type N substrates: (1) a Type N substrate, (2) 30°C, (3) 300°C, and (4) 650°C; the spectra are normalized to the maximum of the Si–O vibration band peaked at 458 cm\(^{-1}\); the bands relating to the vibrations of the Si–O, Si–N and Si–Si bonds are shown with vertical arrows; (b) deconvolution of the absorbance spectral bands in the range from 400 to 650 cm\(^{-1}\) (the sample grown at 30°C): the bands are composed of ten Gaussian peaks associated with the vibrations of the Si–O, Si–N and Si–Si bonds; (c) bands attributed to the vibrations of the Si–N bond in a Type N substrate and samples grown from MB at different silicon deposition temperatures indicated in the plot.