Chemical mapping of silica prepared via Sol-Gel reaction in rubber nanocomposites

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INTRODUCTION

Commercially available silica, whether precipitated, fumed, or prepared otherwise, is an amorphous material that consists of silicon and oxygen atoms in a tetrahedral arrangement interconnected to form a nonregular three-dimensional network of Si–O–Si bonds with silanol groups (Si–OH) present inside the structure and on the surface. The chemical composition and the physical characteristics, such as pore size, surface area, density, and silanol content, are largely responsible for the utility of this material in various applications, including catalysis, separation operations, and composite materials. The surface silanol groups are considered responsible for the high polarity of silica.\(^1\) Several studies have been conducted using one-dimensional (1D) and two-dimensional (2D) solid-state NMR and FTIR techniques for complete surface mapping of highly dispersible (HD) silica\(^4,5\) and related systems\(^6\) in addition to clarifying specific surface interactions.\(^7\) A similar structure is expected for amorphous silica obtained via the sol–gel process, where, by means of hydrolysis of alkoxysilanes and subsequent condensation of silanol groups, silica particles are produced.\(^8,9\) One must realize that during the hydrolysis reaction, alkoxysilanes are not always fully hydrolyzed, suggesting that the non-hydrolyzed alkoxyl groups can be trapped during the sol–gel reaction inside the silica structure and/or remain on the surface. The presence of residual alkoxyl groups for sol–gel synthesized silica has in fact been reported earlier,\(^10,11\) indicating a possibly different chemical structure of silica obtained via the sol–gel reaction.

Traditionally, silica is mixed with rubber compounds via several mixing steps in the presence of a coupling agent; i.e., silica is often referred to as a filler material. The coupling agent aims at improving the silica dispersion and, with that, the rubber reinforcement.\(^12\) The in situ rubber–silica nano-
composites, reinforced with silica formed via the sol–gel reaction, can show better mechanical properties as compared to conventional nanocomposites obtained via mixing.20–22 These in situ rubber–silica nanocomposites typically consist of uniformly distributed, single silica particles, where no silica particle–particle network formation is observed. Recent studies have focused on the structural characterization of rubber–silica reinforced nanocomposites, whereas others have devoted their efforts to understand the changes in mechanical properties using a number of different experimental techniques.23–27 On the basis of these studies, it has been proposed that rubber–silica reinforced nanocomposites may form so-called “trapped entanglements” as a result of temporary bonding of rubber chains to the silica surface. The extent of such rubber–silica interactions strongly depends on the polarity of the rubber, consequently a larger extent of rubber chains to the silica surface. This will most likely result in a larger number of trapped entanglements and, consequently, a larger effect on the overall entanglement density of the rubber matrix. Thus, rubber–silica interactions are considered to be the key parameter for achieving optimal reinforcement in rubber nanocomposites, where several studies28,29 have emphasized their importance for a better understanding of rubber reinforcement.

From a structural point of view, the improved rubber–silica interactions assumed to be present in the in situ rubber–silica nanocomposites have been ascribed to the possibility of fewer silanol groups on the sol–gel synthesized silica surface. One must also note the importance of the amount of ethanol released during the silanization reaction in the compounding process of the conventional nanocomposites and the sol–gel reaction during preparation of the in situ rubber–silica nanocomposites. In general, the removal of ethanol is difficult because it has tendency to form hydrogen bonds with the silanol groups present on the silica surface.30 Therefore, in this paper, an effort is made to detect the ethanol released after the sol–gel reaction in the rubber matrix and to give an estimate for its amount as being crucial for the industrial scale of manufacturing of in situ rubber–silica nanocomposites. Moreover, there is a general lack of information regarding the chemical structure of the obtained silica produced via the sol–gel reaction and consequently little or no understanding of the interface nature or the interactions between the formed silica particle and the rubber matrix. The bound rubber content is commonly used as an indirect measure for the magnitude of these rubber–silica interactions, simply because this parameter is directly related to the surface chemistry of the filler.31 The importance of the specific silica surface was also illustrated by various modification studies32–34 conducted to provide a silica structure with a more hydrophobic surface, resulting in enhanced rubber–silica interactions, and thereby improved compatibility with the rubber matrix. Thus, in this paper we attempt to provide a better understanding of the silica structure as prepared via sol–gel reaction inside a rubber matrix, including the amount of ethanol released and the silica surface and bulk interactions with the rubber matrix.

**EXPERIMENTAL SECTION**

**Materials.** Tetraethyl orthosilicate (TEOS) and n-hexylamine (as catalyst) were purchased from Aldrich and used as received. The highly dispersible (HD) silica is used for conventional mixing under the name Zeosil 1165 MP produced by Rhodia, commonly used in the rubber industry for the production of “green” tires. It has a BET surface area of 150 m²/g and an average particle size of 10–20 nm.

The characteristics of the two rubbers, namely natural rubber (NR; Tun Abdul Razak Research Centre) and ethylene-propylene diene rubber (EPDM; LANXESS), are summarized in Table 1. It should be mentioned that it was difficult to determine the exact molecular weight of the NR sample via gel permeation chromatography due to the presence of residual gel. The literature35 suggests an approximate value for Mₙ of NR of 800 kg/mol and Mₙ/Mₘ = 2.3.

One must note that the used NR and EPDM rubbers were not chemically cross-linked; hence, no vulcanization process has been conducted. The rubbers were used as received, containing only physical network but no chemical connections between the chains. The reason the rubber was not chemically cross-linked is that from literature36 it is known that the cross-link density can influence the final particle size of the sol–gel synthesized silica.

**Preparation of in Situ EPDM and NR Nanocomposites.** In the in situ rubber–silica nanocomposites were prepared as described by Ikeda et al.37 Nonvulcanized compression molded rubber sheets with dimensions of 10 × 10 × 1.5 mm³ were swollen in TEOS and then immersed in an aqueous hexylamine solution with a hexylamine concentration of 0.096 mol/L, using a constant weight ratio of H₂O:TEOS (5:1 w/w). After swelling the rubber matrix in TEOS, the sol–gel reaction proceeded at 80 °C. The hexylamine catalyst gradually hydrolyzes the TEOS leading to formation of silica nanoparticles in the rubber matrix that resulted in in situ NR and EPDM nanocomposites with ~10 wt % of silica, respectively. The amount of silica in wt % can be controlled by varying the degree of swelling with the TEOS and the reaction time, as summarized in Table 2. The resulting in situ rubber–silica nanocomposites were dried for 72 h at 50 °C under vacuum.

**Table 1. Characteristics of Rubber Materials Used**

<table>
<thead>
<tr>
<th>rubber type</th>
<th>NR SMR L</th>
<th>EPDM Keltan S12</th>
</tr>
</thead>
<tbody>
<tr>
<td>composition [wt %]</td>
<td>isoprene (1,4-cis)</td>
<td>ethylene/propylene/ENB</td>
</tr>
<tr>
<td>impurity [wt %]</td>
<td>0.5</td>
<td>55/41/4.3</td>
</tr>
<tr>
<td>Mₙ [kg/mol]</td>
<td>~800</td>
<td>260</td>
</tr>
<tr>
<td>Mooney viscosity ML (1 + 4)</td>
<td>50–60</td>
<td>46 (125 °C)</td>
</tr>
<tr>
<td>Tg [°C]</td>
<td>~65</td>
<td>~54</td>
</tr>
</tbody>
</table>

**Table 2. Preparation of in Situ NR and EPDM Nanocomposites with 10 wt % Silica at 80 °C**

<table>
<thead>
<tr>
<th>type</th>
<th>TEOS amount [wt %]</th>
<th>reaction time [min]</th>
</tr>
</thead>
<tbody>
<tr>
<td>in situ NR nanocomposite</td>
<td>50</td>
<td>60</td>
</tr>
<tr>
<td>in situ EPDM nanocomposite</td>
<td>33</td>
<td>600</td>
</tr>
</tbody>
</table>

**Preparation of Sol–Gel Synthesized ex Situ Silica Powder.** A sample of pure silica powder was prepared via the sol–gel reaction to allow study of the surface chemistry via solid-state NMR, XPS, and FTIR-TGA-MS. The sol–gel synthesized ex situ silica powder was prepared in the same way as the in situ rubber–silica nanocomposites, retaining the 5:1 weight ratio of H₂O:TEOS. In absence of rubber, 50 mL of TEOS was mixed with 250 mL of aqueous solution of hexylamine (0.096 mol/L), and the sol–gel reaction was performed under intense magnet stirring for 2 h at 40 °C. At the beginning of the reaction the mixture was transparent and thereafter turned milky. After 2 h reaction time the mixture was allowed to evaporate overnight in a fume hood and subsequently dried in an oven at 50 °C under vacuum for 72 h. The dried silica powder was additionally heated for 48 h at 110 °C under vacuum and nitrogen flow to remove any residual water or ethanol. A fine fluffy, clear white powder was obtained with BET measured surface area of 336 m²/g and average particle size in a range...
of 40–100 nm. The exchange of protons with deuterons for the HD silica and sol–gel synthesized samples was obtained by stirring the compounds in D2O overnight followed by drying under vacuum at 120 °C.

Nuclear Magnetic Resonance (NMR). To study the chemical structure of the rubber–silica nanocomposites, single-pulse 1H magic-angle spinning (MAS) NMR spectra were recorded on a Bruker AVANCE I 700 spectrometer operating at a Larmor frequency of 700.25 MHz for 1H. These experiments employed a 2.5 mm double-resonance 1H-X MAS probe (Bruker) and a spinning frequency of 5.0 kHz. Both single-pulse 29Si and 29Si{1H} cross-polarization/magic-angle spinning (CP/MAS) experiments were carried out on a Bruker AVANCE I 700 spectrometer with a Larmor frequency of 139.12 MHz for 29Si. These experiments used a 4.0 mm double-resonance 1H-X MAS probe with a MAS frequency of 10.0 kHz and proton decoupling (TPPM) during acquisition.40 These experiments used a radio frequency of 50.0 kHz corresponding to a 1.0 μs 90° pulse. For the single-pulse 29Si MAS NMR spectra a flip angle of ~30° was employed together with a relaxation delay of 30 s. The 29Si{1H} CP/MAS NMR measurements used a relaxation delay of 1.5 s and a CP contact time of 4.0 ms. In the 2D 29Si{1H} heteronuclear correlation (HETCOR) experiments, the indirect dimension was sampled in 128 steps using a t1 dwell time of 30.0 μs. The 2D 1H{13C} HETCOR spectrum was recorded on the Bruker AVANCE I 700 spectrometer—the same NMR spectrometer with a resonance frequency of 176.08 MHz for 1C in a 4.0 mm probe. A MAS frequency of 5.0 kHz and 1H TPPM decoupling during acquisition were applied. A relaxation delay of 3 s was used, and the indirect dimension had 64 steps for a t1 dwell time of 50.0 μs and a CP contact time of 3.0 ms. The 15N{1H} CP/MAS NMR experiment was performed on a Bruker Avance I 300 spectrometer operating at a 7.014 MHz Larmor frequency of 30.43 and 300.23 MHz for 1H. A relaxation time of 3 s, 5.0 kHz MAS, 1H TPPM decoupling, and a CP contact time of 5.0 ms were used. The 2D 13C{1H} wide-line separation (WISE)43 NMR experiment was also carried out on the on a Bruker Avance I 300 spectrometer with a Larmor frequency of 75.49 MHz for 13C. The spectrum was recorded at 40.0 kHz MAS using a 7.0 mm double-resonance 1H-X probe with a CP contact time of 3.0 ms. To prevent spinning side bands (ssbs) in the 13C dimension, the total suppression of spinning sidebands (TOSS) technique was applied.43 Chemical shifts for 1H and 13C are reported in ppm relative to TMS using solid adamantane (δadam = 1.85 ppm for 1H, δadam = 38.484 for 13C) as an external reference.44 Chemical shifts for 15N were referenced to the ammonium site of NH4NO3 at −358.4 ppm.45 The deconvolution of 29Si MAS NMR spectra was performed using MATLAB by taking into account the resonance positions and their Gaussian/Lorentzian line widths as described by a pseudo-Voigt function. The final integration used the spectral edges as limits.

X-ray Photoelectron Spectroscopy (XPS). XPS is a surface analysis method that provides information for the chemical composition of the material surface based on the obtained binding energies. The measurements were carried out with a Thermo Scientific K-Alpha, equipped with a monochromatic small-spot X-ray source and a 180° double focusing hemispherical analyzer with a 128-channel detector. Spectra were obtained using an aluminum anode (Al Kα = 1486.6 eV) operating at 72 W and a spot size of 400 μm. Survey scans were measured at a constant pass energy of 200 eV. The background pressure was 2 × 10−9 mbar, and during measurement 4 × 10−8 mbar of argon was used for the charge compensation as dual beam source. The low-energy ions generated cause no sample damage but compensate for charging, preventing a shift in peak values that possibly could result in incorrect assignment of the peaks.

Transmission Electron Microscopy (TEM). The morphology of the nanocomposites was studied using TEM. Ultrathin sections (70–100 nm) were obtained using a Reichert-Jung Ultracut E microtome equipped with a diamond diamond knife for dry trimming (45° angle) and sectioning (35° angle) at a temperature of −140 °C. The sections were placed on a 200 mesh copper grid with a carbon supportive layer and examined with a Sphera electron microscope, FEI Tecnai 20 operated at 200 kV.

Bound Rubber Determination. The bound rubber content was estimated by immersing the rubber/silica nanocomposites with various silica amounts in THF. The ratio between the rubber/silica nanocomposites and the solvent was kept constant (2% w/v) and the samples were immersed in the solvent for 48 h at room temperature while shaking. The residues were dried for 72 h at 50 °C under vacuum. The bound rubber content was calculated using

$$\text{wt % bound rubber content} = \frac{W_{\text{residue}} - W_{\text{silica}}}{W_{\text{original}} - W_{\text{silica}}} \times 100$$

where $W_{\text{residue}}$ (g) is the weight of the sample after solvent treatment, $W_{\text{original}}$ (g) is the weight of the sample before solvent treatment, and $W_{\text{silica}}$ (g) is the weight of silica in the nanocomposite.

Thermogravimetric Analysis–Infrared Spectroscopy–Mass Spectrometry (TGA-FTIR-MS). TGA-FTIR-MS measurements were performed at the Materia Nova Institute in Mons, Belgium. The purpose of the measurements was to determine the organic components, including the ethanol released from the sol–gel synthesized silica at a particular temperature. The thermogravimetric analyses were performed on a TGA Q5000 from TA Instruments using nitrogen as purge gas. The analysis of the evolved gases during the thermal degradation was carried out with a Bruker Tensor 27 FTIR spectrometer. Spectra were recorded (4000 to 600 cm−1) using a gas cell heated at 225 °C and a mercury cadmium telluride (MCT) detector. The evolved gases were also analyzed by mass spectrometry on a Thermoset quadrupolar apparatus from Balzers Instruments covering a mass range from 0 to 300 amu. Evolved gases were ionized by electron impact (70 eV). The simultaneous analysis of the gases by FTIR and MS spectrometers was made possible by a double coupling device. It consists of a Teflon-coated stainless steel T” connection which was heated at 225 °C, thus avoiding any condensation of evolved gases. To this connection the mass spectrometer is connected via a 1 m long quartz capillary (i.d. 150 μm) heated at 200 °C, while the gas cell of the FTIR spectrometer is connected through a 1 m long stainless steel tube (i.d. 2 mm) heated at 225 °C (N2 flow of 25 mL/min, heating rate 20 °C/min).

RESULTS AND DISCUSSION

Silica Structure via Solid-State NMR. The structure of the silica particles formed within the in situ rubber–silica nanocomposites is of crucial importance from several points of view. First, it is indicative for the completion of the sol–gel reaction (full hydrolysis of TEOS with no residual ethoxy groups), and, second, it determines the rubber–silica interaction and thereby the degree of rubber reinforcement. This triggered us to perform a detailed analysis of the chemical structure of sol–gel synthesized silica in comparison with commercially available, highly dispersible (HD) silica using solid state NMR. A D2O treatment was applied to both silicas to distinguish between isolated and exchangeable silanol groups (SiOH) and H2O present on the surface or/and inside the silica particles. Such differences may be indicative for possible changes in hydrophobicity.

Figures 1a and 1b display the single-pulse 1H MAS NMR spectra of sol–gel synthesized and HD silica and (black lines), respectively, and their deuterated analogues (red lines). We note that the chemical exchange of protons with deuterons in both silica samples is a uniform process, where protons that are accessible at the surface and in the bulk may be exchanged. From a comparison of the two 1H MAS NMR spectra (black and red) in Figure 1a it is evident that the protons at 1.9 ppm from OH groups at the surface can be exchanged by deuterons, showing a lower intensity peak. Bulk OH (hydrogen-bonded silanol groups) and silanol groups (OH and (OH)2) trapped inside the silica structure, resonating at the positions indicated in Figure 1a, do not show any changes in peak intensity after

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exchange with the D₂O. Moreover, both the bulk OH and water signals do not occur as sharp, single resonances but are spread out over a ¹H chemical shift range of up to 6 ppm. For isolated OH groups at the surface, a ¹H chemical shift range between 1.8 and 2.0 ppm has been reported, which is in good agreement with data presented in Figure 1a.⁴⁷ We assign the additional proton signal at 1.1 ppm to trapped, isolated OH groups inside the silica particles.⁴⁸

The ¹H MAS NMR spectrum of sol–gel synthesized silica (Figure 1b) does not include well-resolved proton resonances from trapped OH and (OH)₂ groups, possibly due to the more intense hexylamine resonances that dominate the spectrum in that region. The signals assigned to the aliphatic tail of hexylamine show a significant intensity decrease after the D₂O treatment, suggesting that hexylamine molecules adsorbed on the silica surface to some extent can be removed by washing with D₂O. Another interesting difference between the sol–gel synthesized silica and HD silica is the difference in the ratio between H₂O and bulk OH, resonating between 3.5–6.0 ppm and 6.0–9.0 ppm, respectively (dashed lines in Figure 1a,b). The lower relative intensity of the H₂O peak observed for the sol–gel synthesized silica possibly originates from having fewer OH groups (i.e., a more hydrophobic surface) and therefore fewer physisorbed H₂O at the surface.

To further characterize the differences in chemical structure for sol–gel synthesized and HD silica, we have used 2D ²⁹Si{¹H} heteronuclear correlation (HETCOR) NMR spectroscopy as summarized in Figure 2. Figure 2a shows a strong, intense correlation between Q³ silicon atoms and OH groups, which is also observed for HD silica in Figure 2b. This correlation provides clear evidence for the existence of OH groups at the silica surface (see Figure 2c). Interestingly, the remaining spectral features observed in the 2D ²⁹Si{¹H}
HETCOR NMR spectra for sol–gel synthesized and HD silica are very different. In Figure 2a, the N–CH₂ (hexylamine) resonates at proton chemical shift of 2.5 ppm and the CH₃ (methyl) at 0.8 ppm. The ethoxy groups show correlations to both Q² and Q⁴ silicon atoms, indicating that their presence is not only at the surface but also in the bulk. The 2D correlation signals between the N–CH₂ group of hexylamine and the Q⁴ sites further indicate that the polar amine headgroup is only located at the surface of the silica nanoparticles. Moreover, bulk OH and hydrogen bonded OH groups, represented by a broad proton resonance at 3–8 ppm for HD silica, are not observed for sol–gel synthesized silica (cf. Figures 2a and 2b). A possible reason for this difference between the two silicas is that the ethoxy groups and hexylamine present at the silica surface shield the OH groups, preventing them from forming hydrogen bonds. To confirm the presence of hexylamine and ethoxy groups for sol–gel synthesized silica, we performed 2D ¹³C{¹H} HETCOR measurements, the results of which are shown in Figure 3a. This spectrum primarily shows the intramolecular correlations of the hexylamine (carbons 1–6) with their directly bonded alkyl protons. The correlation peaks originating from the ethoxy groups (denoted by A and B) are very weak, reflecting a low abundance of these groups. The presence of hexylamine in both sol–gel synthesized silica and the in situ synthesized NR nanocomposite is verified by the ¹⁵N{¹H} CP/MAS NMR spectra displayed in Figures 3b and 3c, which both include a single ¹⁵N resonance at −347 ppm.

We note that for both of these samples ¹⁵N-labeled hexylamine was used during the sol–gel reaction, since nitrogen has a very low abundance (0.37%) and consequently a low NMR sensitivity. Thus, using ¹⁵N-labeled hexylamine, we are able to confirm the presence of hexylamine in both sol–gel synthesized materials (Figure 3b,c). This result further demonstrates that the sol–gel synthesis and the inherent modification of the silica surface can be realized without additional chemical reactions and use of modifiers, highlighting its potential for industrial manufacturing processes and applications.

Additional evidence for the presence of hexylamine at the silica surface is provided via the 2D ¹³C{¹H} wideline separation (WISE) NMR spectrum shown in Figure 4, which gives an estimation of the molecular flexibility. A rigid hexylamine (trapped inside the silica structure) would appear as a broad proton signal with a spectral width of up to 100 kHz (vertical dimension), while a mobile hexylamine (free, nonbonded) would give rise to quite narrow proton lines. On the basis of the observed vertical width of ~20–50 kHz (Figure 4), we conclude that the hexylamine molecules are not trapped inside the silica nanoparticles, but more likely in the vicinity of the silica surface in an intermediate state, i.e., neither fully rigid nor fully mobile. It is expected the presence of hexylamine near the silica surface to contribute toward a more hydrophobic surface nature of the silica nanoparticles and improve its compatibility with the rubber, leading to improved mechanical properties of a rubber–silica nanocomposite.

The quality of the sol–gel synthesized and HD silicas can be determined from the ratio between Q²:Q³:Q⁴ silicon atoms, and Figure 5 shows the quantitative single-pulse ²⁹Si MAS NMR spectra for HD silica, sol–gel synthesized silica, and the in situ NR nanocomposite. In general, all ²⁹Si MAS NMR spectra of Figure 5 include three different resonances at −91.7, −101.4, and −110.9 ppm, corresponding to Q², Q³, and Q⁴ silicon tetrahedra, respectively. We note that it was difficult to distinguish the different types of silanol groups for the sol–gel synthesized silica from FTIR (data not shown). Instead, solid-state ²⁹Si MAS NMR provides a clear distinction and visualization of silicon tetrahedra bearing OH groups in addition to the silicon tetrahedra with residual (or non-hydrolyzed) ethoxy groups remnant from the TEOS precursor.

On the basis of the ²⁹Si MAS NMR spectra in Figure 5, we have performed a spectral deconvolution to determine the
intensities for Q^2, Q^3, and Q^4, as illustrated in Figure 5 by the dashed lines. These show that the quality of the sol–gel synthesized silica as described by the Q^2:Q^3:Q^4 ratio is 1.4(0.2):23.9(1.1):74.7(1.3), while this ratio for HD silica is 1.4(0.2):23.2(1.1):75.4(1.3). A slightly different ratio of 1.8(0.3):30.0(1.2):68.2(1.4) is obtained for the silica particles in the in situ NR nanocomposite. In this regard, the sol–gel synthesized silica has a structure that appears to be of very similar quality to that of HD silica. This can be ascribed to the confinement effect of the rubber on the growing silica particles resulting evidently in a somewhat lower degree of particle condensation. One must note that the silica particles formed in the in situ NR nanocomposite show a somewhat lower quality when compared to the sol–gel synthesized and the HD silica. This can be ascribed to the diffusion limitations of the reactants (TEOS and hexylamine) inside the rubber matrix, resulting evidently in a somewhat lower degree of particle condensation. Interestingly, since the Q^2:Q^3 ratios for sol–gel synthesized and HD silica are almost the same, and the only difference is that the sol–gel synthesized silica contains not only Si–OH but also ethoxy (Si–OCH$_2$CH$_3$), one may assume that fewer OH groups are present in the sol–gel synthesized silica. This observation finally resolves a piece of the puzzle for the chemical structure of the sol–gel synthesized silica. Here, one may indeed conclude that not only the sol–gel synthesized silica nanoparticles do contain fewer Si–OH groups, but in addition also residual ethoxy groups and hexylamine on or near the silica surface. These structural features contribute to the hydrophobic nature of the silica nanoparticles, regardless whether these are prepared ex situ or inside a rubber matrix. Though the above results from solid-state NMR enable a detailed structure analysis of the different types of silica, it cannot provide easy access to information regarding the type of interactions existing between the silica nanoparticles and hexylamine and ethoxy residuals on or near the surface. Such phenomena become an issue when the stability of silica structure is addressed, and for this purpose we used X-ray photoelectron spectroscopy (XPS) to gain a better insight.

Mapping the Sol–Gel Synthesized Silica Surface Species by XPS. XPS is used for mapping the elemental composition of sol–gel synthesized silica in powder form and HD silica. The reason we did not use the silica particles formed directly in the rubber matrix is twofold. First, XPS only works for fine powders, and second, burning off the rubber at higher temperatures or treating it with an organic solvent is not desirable as it might affect the silica surface and lead to wrong conclusions. The chemical species from the analysis of the sol–gel synthesized and HD silica are summarized in Table 2. These data confirm the difference in the surface chemistry between the sol–gel synthesized and HD silica. The first difference originates from the total carbon content, which for sol–gel synthesized silica is 7.2% and 2.0% for HD silica. The higher carbon content for sol–gel synthesized silica can be ascribed to the presence of remnant ethoxy groups, hexylamine, and adsorbed carbon dioxide. The carbon content observed for HD silica can be only assigned to adsorbed carbon dioxide, as proven by the absence of a carbon XPS signal after removing a 10–15 nm layer from the surface of the HD silica with argon sputtering (data not shown).

Three types of carbon signals are included in Table 3 and assigned to C−C/C−H, C=O, and C=O bonds appearing at 284.4, 285.8, and 287.3 eV, respectively. Particularly, the higher concentration of the C−O bond (1.42%) for the sol–gel synthesized silica is consistent with the presence of ethoxy groups on the surface, although part of this signal can also originate from ethanol that is a byproduct of the sol–gel reaction.51 Moreover, the amino group of the hexylamine resulted in two characteristic signals: one at 401.4 eV assigned to a hydrogen-bonded amine and another at 401.6 eV. Earlier studies suggested that the origin of the signal appearing at 399.16 eV is due to either free amine52 or chemisorbed amine forming a Si–N bond.53,54 The TGA-FTIR-MS measurements performed in this work (see below) showed release of hexylamine at about 500 °C, suggesting that it is more likely that chemisorbed amine is present rather than free amine at those temperature conditions. Moreover, since the sol–gel synthesized silica proceeds via an ionic mechanism, where hexylamine becomes protonated by accepting a proton from water, one expects to observe evidence for the ionic interactions between the positively charged hexylamine and possibly some remnant negatively charged silica entities situated at the silica surface. However, such ionic interactions should result in an additional amine signal located at 402.8 eV,52 which in our case is not observed. The O(1s) binding energy for both the sol–gel synthesized and HD silica appears at 532.6 eV and corresponds to a hydrogen bonded O(1s) C=O bond (1.42%).

Table 3. XPS Results for Sol–Gel Synthesized and HD silica

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<tr>
<th>atomic orbitals</th>
<th>bond type</th>
<th>binding energy [eV]</th>
<th>atomic conc [%]</th>
<th>binding energy [eV]</th>
<th>atomic conc [%]</th>
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<tr>
<td>C(1s)</td>
<td>C−C/C−H</td>
<td>284.4</td>
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<td>C(1s)</td>
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<td>1.42</td>
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<tr>
<td>C(1s)</td>
<td>C=O</td>
<td>287.3</td>
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<tr>
<td>N(1s)</td>
<td>NH$_2$ (free)</td>
<td>399.2</td>
<td>0.42</td>
<td>405.2</td>
<td>0</td>
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<tr>
<td>N(1s)</td>
<td>NH$_2$ (HB)$^{a}$</td>
<td>401.4</td>
<td>0.28</td>
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<td>Na(1s)</td>
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<td>O−(SiH$_2$C)</td>
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<td>68.5</td>
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<td>Si(2p)</td>
<td>SiO</td>
<td>103.3</td>
<td>27.8</td>
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</table>

$^{a}$HB = hydrogen bonded.
to all bonded oxygen atoms (O−Si, O−H, and O−C), having very close binding energies and for this reason are difficult to distinguish. The Si(2p) binding energy is 103.3 eV and corresponds to Si−O−Si bonds. Thus, the XPS study shows that hexylamine is attached to the silica surface only via hydrogen bonding and chemisorption.

**Time-Resolved Thermal Treatment of Sol−Gel Synthesized Silica via FTIR-TGA-MS.** The physical and chemical changes of HD silica during a thermal treatment are reported in the literature while similar reports for sol−gel synthesized silica are still very limited. The purpose of using a combinatory TGA-FTIR-MS in this work is to be able to estimate whether ethanol is present inside the silica particles and to assess the way it is bonded to the silica surface (physical or chemical interactions) using the evaporation temperature as an indicator. It should be noted that a precise determination of the exact temperature at which one compound begins to evolve is difficult because there is always a certain delay between the evaporation of the compounds in the TGA instrument and their detection by the FTIR and the MS instruments. For this reason, when assigning a temperature, one must consider a narrow temperature region instead of a particular temperature. Spectra of the thermal treatment of sol−gel synthesized and HD silica using TGA-FTIR-MS are presented in Figure 6.

Figure 6 shows the evolution of chemical compounds at particular temperature regions during heating of HD and sol−gel synthesized silica. Both 3D representations show water peaks, mainly observed in the ranges of 1300−1800 and 3200−3800 cm$^{-1}$, and the carbon dioxide (CO$_2$) peak at 2350 cm$^{-1}$, all indicated with black dashed lines to guide the reader’s eye. The extensive carbon dioxide evolution around 750 °C is assigned to the intensive decomposition of the organics evolving from the sol−gel synthesized silica at these high temperatures. The large peak of CO$_2$ for sol−gel synthesized silica in the same temperature region is also observed in the HD spectra. The presence of adsorbed CO$_2$ on the HD silica surface was earlier confirmed by XPS (see Table 3) expected to be removed already at lower temperatures, so the large CO$_2$ peak appearing around 750 °C in HD silica can be ascribed to presence of degradation products. Moreover, the spectrum of the sol−gel synthesized silica in Figure 6b contains other, smaller peaks appearing at various wavelengths, suggesting the evolution of other organics, besides water and CO$_2$.

A more quantitative view of the chemical changes during the thermal treatment of the sol−gel synthesized and HD silica is given by the weight loss shown in Figure 7. The total weight loss of the sol−gel synthesized silica (15.3%), as calculated from the TGA data, is higher than for the HD silica (9.5%), indicating a higher concentration of organic compounds present.

In region 1 (25−150 °C) of Figure 7, weight losses are caused only by the removal of physical water from both materials. For sol−gel synthesized silica, FTIR and MS analysis confirmed that only water (no other organics) was removed in this temperature interval. Even though the two silica materials were dried under the same conditions, the sol−gel synthesized silica released less adsorbed physical water (1%) than the HD silica (2%), despite the significantly larger BET surface area, 336 m$^2$/g. We would expect silica with larger surface area to have higher absorbing capacity for moisture, hence more physically bounded water. However, as shown in the study so far, the sol−gel synthesized silica has a unique hydrophobic surface due to the presence of hexylamine and ethoxy groups that prevents additional absorption of physically bounded water despite the larger surface area. Therefore, the lower absorbed physical water (1%) by the sol−gel synthesized silica can be clearly ascribed to the hydrophobic surface. One must consider that determining the “specific” surface area, hence considering the density of the silica, is very questionable. The sol−gel synthesized silica is expected to have lower density than 2.1 g/
cm$^3$ (HD), not fully completed condensation step during the sol–gel reaction, as it was shown from several density measurements (using a picnometer) that density ranges between 1.7 and 1.95 g/cm$^3$.

This points toward a more hydrophobic nature of the sol–gel synthesized silica due to the ethoxy groups and the hexylamine residing at or near the surface. In region 2 ($150\text{–}330 ^\circ \text{C}$) and region 3 ($330\text{–}490 ^\circ \text{C}$), more intensive chemical changes occur in the sol–gel synthesized and HD silica, resulting in the release of chemically bonded water, rapid condensation of the silanol groups in both materials and, for the sol–gel synthesized silica, the additional release of organic components as indicated by the FTIR spectra shown in Figure 8.

In Figure 8a, the FTIR spectra of sol–gel synthesized silica suggests the presence of hexylamine ($1121 \text{ cm}^{-1}$) and ethanol ($1055 \text{ cm}^{-1}$) that begin to evolve at approximately $300 ^\circ \text{C}$. These observations indicate that both hexylamine and ethanol are bound to the silica surface via hydrogen bonds. As indicated by the XPS results (see Table 3), hexylamine is hydrogen bonded, and we expect ethanol also to be hydrogen bonded to the OH groups existing on the silica surface. As the absorption bands of a component can differ in its liquid and gaseous state, the absorption bands of hexylamine and ethanol were determined in their gas phase (FTIR-TGA), as shown in Table 4.

Another peak showing a similar trend with temperature as the two peaks from ethanol and hexylamine appears in the range 1220–1225 cm$^{-1}$. This signal is difficult to assign because it does not correspond to any of the components shown in Table 4. Since it is close to the one of ethanol (1242 cm$^{-1}$) and since MS does not suggest the presence of other relevant organics, we might assume that it originates from ethanol. The difference between sol–gel synthesized and HD silica at the same temperature conditions ($\sim 330 ^\circ \text{C}$) is clearly observed in Figure 8b, where HD silica does not reveal the presence of any organics, as indicated by the absence of peaks in the region 600–1400 cm$^{-1}$, while the sol–gel synthesized silica clearly shows the presence of organics via three distinguished peaks.

Following the thermal treatment of sol–gel synthesized silica with the mass spectrometer part (MS) of the combinatory FTIR-TGA-MS equipment until $1000 ^\circ \text{C}$ revealed an additional interesting insight into the sol–gel synthesized silica, as shown in Figure 9.

In Figure 9, hexylamine, ethanol, and water amounts as detected by MS during thermal treatment of sol–gel synthesized silica up to $1000 ^\circ \text{C}$ using a heating rate of 3 $^\circ \text{C}$/s.

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In Figure 9, hexylamine, ethanol, and water are followed by their ionization species with 30, 31, and 18 mass-to-charge ratios ($m/z$), respectively, as a function of time and temperature. These ionized species were selected because these give the highest abundance in the MS spectra of their pure components. The results suggest a relatively low content of ethanol and hexylamine when compared to water. There are several reasons for the higher amount of water released throughout the thermal treatment: (i) condensation of silanol groups, (ii) strong hydrogen-bonded water on the outer surface or inner structure, (iii) water trapped in small micropores inside the particles and trapped water molecules in the sol–gel synthesized silica structure that can be released at high temperatures. Interestingly, in this temperature region (corresponding to region 4 in Figure 7, indicated by the vertical
the challenge involved in determining the surface area of the gel synthesized silica in the rubber matrix. One must consider the possibility that ethanol molecules might also be trapped inside the silica particles and released in this temperature range. It should be noted that the ethanol amount is surprisingly low. A possible reason could be that the hydrophobic surface on the sol–gel synthesized silica limits the number of ethanol molecules that can interact via hydrogen bonds with the Si–OH groups. In addition, the presence of residual ethoxy groups inside the silica particles also creates an unfavorable environment for the ethanol to remain trapped during the sol–gel reaction. The temperature region of the observed maxima in Figure 9 corresponds well to the extensive evolution of CO$_2$ in the same temperature range (see Figure 6b), which is related to intensive decomposition of the high amount of organics released.

To our surprise, residual nonreacted TEOS was not registered in the sol–gel synthesized silicic acid powder, suggesting excellent conversion of the precursor into a solid silica particles and/or proper drying conditions.

**Physical and Chemical Structure of Sol–Gel Synthesized Silica.** The sol–gel synthesized silica is grown directly in the EPDM rubber matrix, uniformly distributed in the formed in situ EPDM nanocomposite with sizes ranging between 25 and 35 nm, as shown in Figure 10. The size, shape, and particle density (number of particles in certain volume) of the sol–gel synthesized silica strongly depend on two general factors: the type of rubber matrix in which they are grown (EPDM, NR, SBR) and the reaction parameters (e.g., amount of TEOS, temperature, and reaction rate). The morphology of the obtained in situ nanocomposites as a function of the two main factors has been extensively discussed more in detail in our previous paper.29 In this paper we also showed existence of so-called gradient in particle size along the sample thickness due to diffusion limitations, where larger silica (∼35 nm) particles were formed closer to the surface and smaller (∼16 nm) toward the sample center. This is an important factor that might significantly influence the overall surface area of the sol–gel synthesized silica in the rubber matrix. One must consider the challenge involved in determining the surface area of the sol–gel synthesized silica which can be used either prepared in ex situ conditions or isolated from the rubber matrix by burning or dissolving the rubber. In the ex situ obtained sol–gel synthesized silica one must be aware of the size (40–110 nm), shape, and particle aggregation difference from the silica grown directly in the rubber (well-separated small size particles 20–35 nm) or the consequences of burning or dissolving the rubber for isolating the sol–gel synthesized silica which can influence the surface of the silica and hence the surface area. Despite the prons and cons of both approaches, we decided to use the sol–gel synthesized silica prepared ex situ. The BET measurement showed surface area of 336 m$^2$/g, which is significantly larger when compared to the one of the HD silica (150 m$^2$/g). One possible explanation can be that the sol–gel synthesized silica particles are less aggregated due to the more hydrophobic surface, resulting in overall larger surface area unlike the strongly aggregated HD silica due to its hydrophilic surface. Regarding the chemical structure, in this study, we showed that is significantly different from the HD silica. Ethoxy groups have shown to be present inside the silica and on the silica surface, while hexylamine molecules reside mainly at the surface, in full agreement with the results of our previous study on the silica formation mechanism in rubber matrix via sol–gel reaction.49

In this study using the toluene model system and several characterization techniques, we showed that the silica formation in rubber matrix proceeds via an emulsion mechanism, where the hexylamine, being the catalyst, behaves also as surfactant. We revealed that hexylamine forms inverse micelles in the rubber matrix which act as nanoreactors hosting the sol–gel reaction for silica formation. In these inverse micelles the polar groups amino groups of the hexylamine are pointing toward the enclosed water in the center of the micelle, and the nonpolar aliphatic chains are facing the rubber phase. After solidification of the silica particles, the hexylamine (if not removed via calcination at 400 °C) is expected to remain on the silica surface, resulting in silica particles with a more hydrophobic nature and consequently increased compatibility with the rubber matrix. This provides a unique chemical structure which resembles so-called “hairy” silica particles, as schematically presented in Figure 10.

These particles are called “hairy” due to the ethoxy groups and the long aliphatic chains of the hexylamine sticking out from the silica surface. One can imagine that, besides TEOS being an alkoxy precursor for silica formation in the sol–gel reaction, other precursors can also be used, such as propyl and butyl orthosilicate. These precursors have longer aliphatic tails...
that are expected to lead to even higher surface hairiness of the silica particles and consequently higher surface hydrophobicity and hence a better compatibility with the rubber matrix and potentially improved mechanical properties of in situ rubber–silica nanocomposites.

**Entrapped Rubber Chains in Sol–Gel Synthesized Silica Particles.** Considering that the silica nanoparticles are formed via a sol–gel reaction within the rubber matrix, it might be a possibility that these in situ rubber–silica nanocomposites have rubber chains trapped inside the sol–gel synthesized silica particles. To test this hypothesis, we have recorded 2D $^{29}$Si{[H]} HETCOR NMR spectra for both the in situ EPDM and the conventional nanocomposite, as shown in Figure 11.

The 2D $^{29}$Si{[H]} HETCOR NMR spectrum for the conventional composite shows the expected correlation between Q^3 species and OH groups (Figure 11a), but for the in situ EPDM nanocomposite (Figure 11b), rather surprisingly, an intense correlation peak is present at ~0.8 ppm due to the interaction between CH₃ groups and Q^4 species. While the CH₃ peak as such can be easily explained by the presence of ethoxy groups or hexylamine as observed in the 2D $^{29}$Si{[H]} HETCOR spectrum for the sol–gel synthesized silica (cf. Figures 2a and 11b), the only possible way for a CH₃ group to correlate even more strongly with Q^4 species, which is mainly present in the interior of the silica representing the Si–O–Si network, is that the CH₃ group originates from the EPDM (the propylene unit) rubber itself (Figure 11c). This conclusion relies on the fact that all 2D $^{29}$Si{[H]} HETCOR experiments (Figures 2 and 11) were recorded using identical conditions. Moreover, our choice of a long CP time of 4.0 ms will also lead to an increase in the polarization transfer from ¹H to $^{29}$Si for Q^4 compared to Q^3 with protons in their direct vicinity (see Figure 2c) due to the difference in CP kinetics for such groups.³⁰ The absence of the CH₃ peak in the EPDM conventional nanocomposite (Figure 11b) additionally contributes to our hypothesis of rubber chains entrapped inside silica particles. Obviously, entrapped chains can be, at least partially, responsible for the high bound rubber content of the in situ rubber–silica nanocomposites. The same correlation is also observed for the in situ NR nanocomposite (data not shown), which implies that this process may occur regardless of the rubber type. Thus, to the best of our knowledge, this work reports the first, and rather intriguing, evidence of entrapping polymer chains in growing sol–gel synthesized nanoparticles.

**CONCLUSIONS**

NR and EPDM rubber nanocomposites were prepared via an in situ sol–gel reaction using hexylamine as catalyst and TEOS as alkoxy precursor for the formation of silica particles directly in the rubber matrix. The chemical mapping of the silica particles was performed by solid-state NMR and XPS, revealing the presence of residual ethoxy groups and hexylamine on the silica surface. Ethoxy groups are present inside the silica structure as well as on the surface. Together with the hexylamine present at the surface, this results in an increase of hydrophobicity for the sol–gel synthesized silica as compared to silica particles in conventional rubber–silica nanocomposites. FTIR-TGA-MS confirmed the higher amount of organic material present for the sol–gel synthesized silica as well as the low amount of ethanol present. The latter is important as the ethanol amount released is crucial for the industrial production of in situ rubber–silica nanocomposites. The chemical structure of sol–gel synthesized silica inside the rubber matrix, as well as ex situ prepared sol–gel synthesized silica via an aqueous solution of hexylamine, has comparable quality to that of commercial HD silica, as judged from the Q^2:Q^3:Q^4 ratios determined using $^{29}$Si MAS NMR measurements. The excellent dispersion of the sol–gel synthesized silica particles in the rubber matrix, their hydrophobic surface, and the entrapment of the rubber chains in the growing silica particles can be used as a scaffold for creating nanocomposites with improved mechanical properties as compared to conventional nanocomposites.
AUTHOR INFORMATION

Corresponding Authors
*E-mail elena.miloskovska@sabic-ip.com (E.M.).
*E-mail mrh@chem.au.dk (M.R.H.).

Notes
The authors declare no competing financial interest.

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