**High-Resolution Structural Characterization of a Heterogeneous Biocatalyst using Solid-State NMR**

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**ABSTRACT:** Solid-state magic-angle spinning (MAS) NMR spectroscopy was employed to investigate structural detail in an enzyme, human carbonic anhydrase II (hCA II) in uniformly $^{15}$N and selectively $ (^{15}$N leucine) enriched states, covalently immobilized on epoxy-functionalized silica. The immobilized hCA II retained 71% of its specific enzymatic activity when compared to the free enzyme in solution. Based on the one- and two-dimensional $^1$H, $^{13}$C, $^{15}$N and $^{29}$Si MAS NMR spectra, chemical shift assignments could be obtained from the silica support, covalent linker and the immobilized enzyme. The successful covalent immobilization of the enzyme on epoxy-silica was confirmed by the appearance of signals from the aromatic and carbonyl groups in the immobilized enzyme in addition to signals from the modified support. Most notably, our MAS NMR results suggest that the covalent immobilization of the hCA II on epoxy-silica does not significantly affect the structural integrity of the protein.

**INTRODUCTION**

The immobilization of proteins and living cells on non-biological surfaces plays a central role in a wide range of important technological applications, including industrial biocatalysis, drug delivery, medical diagnosis, wastewater treatment, biosensing, and textile and detergent manufacture. In particular, immobilized enzymes are widely used in such diverse applications as the conversion of biomass to fuel, the synthesis of fine chemicals, pharmaceutical production, and the food and cosmetics industries.

Although there have been extensive studies of the catalytic activity of enzymes immobilized by a wide range of methods, rational design of the heterogeneous biocatalytic system still remains a considerable challenge as very little is known about the state of the enzyme, support and the linker upon immobilization. Solid-state magic-angle spinning (MAS) NMR is an ideal tool for studying such heterogeneous systems, as the system under consideration does not require any long-range order. Furthermore, solid-state NMR has been successfully employed for studying complex biological systems that are not accessible by X-ray crystallography or solution-state NMR. MAS NMR has been used previously to study a variety of immobilized enzymes and supports. However, most of these research studies were focussed mainly on studying the support systems and/or the biochemical characterization of the biocatalytic system. There have also been quite extensive high-resolution MAS NMR studies of proteins and peptides involved in biomineralisation, adsorbed on the surfaces of the inorganic materials whose growth they control. But there have so far only been a few studies of signals from enzyme molecules immobilized on supports as biocatalysts. Fragai et al showed high-resolution two-dimensional spectra of $^{13}$C and $^{15}$N labelled enzymes entrapped in peptide-templated silica gel particles. This report indicated that the three-dimensional structures of the enzymes were hardly affected by immobilization. Subsequent work by the same group demonstrated enhanced sensitivity using dynamic nuclear polarisation and explored the feasibility of $^1$H-detected measurements. Natural abundance silica-entrapped lysozyme $^{13}$C signals were also detectable using DNP-enhancement. Furthermore, Ramirez-Wong et al. used $^1$H signals to determine the ratio of lysozyme to chitosan in a deposited layered conjugate.
Very recently, MAS NMR has been successfully employed to characterize a covalently immobilized enzymatic system consisting of a model enzyme α-chymotrypsin, mesoporous silica as the matrix, and (3-glycidylxypropyl)trimethoxysilane (GLYMO) as the covalent linker. However, this natural abundance MAS NMR study of covalently immobilized chymotrypsin was hampered by poor NMR sensitivity, making it impossible to obtain detailed structural and dynamic information from the immobilized enzyme.

In this research work, we report finer structural details of a model enzyme, human carbonic anhydrase II (hCA II) in the isotopically labelled (15N) state before and after covalent immobilization on epoxy-functionalized silica. Human carbonic anhydrase II is a relatively low molecular weight (~29 kDa, 260 residues) enzyme suitable for catalyzing the reversible hydration of carbon dioxide. The structure of the enzyme has been solved by X-ray crystallography and extensively studied by NMR and is therefore an ideal candidate for covalent immobilization studies using MAS NMR. Since the size of the protein (~29 kDa) still poses challenges for NMR to provide atomic-level structural information, hCA II samples were prepared in natural abundance and isotopically enriched (15N) states (uniformly labelled samples termed hereafter as [U-15N]/hCA II). Furthermore, the samples were selectively 15N labelled for the most abundant amino acid residue leucine (termed hereafter as [15N Leu]/hCA II) to reduce problems arising from spectral degeneracy in multi-dimensional MAS NMR experiments. Our results show that MAS NMR can be successfully employed to characterize the support, linker and the enzyme before and after immobilization and a simple schematic model of the complete heterogeneous biocatalytic system can be proposed.

**MATERIALS AND METHODS**

**Protein Expression, Purification and Characterization**

The hCA II plasmid (pACA) used for the production of hCA II mutants was a generous gift from Carol A. Fierke (University of Michigan, USA). hCA II double mutants were prepared by site-directed mutagenesis. The wild-type hCA II plasmid was used as the initial template for the first mutation (C206S) while plasmids already containing the C206S mutation were used for the final double mutants. The site-directed mutagenesis steps were carried out according to the procedure described by Zheng et al. All constructs were overexpressed in [U-15N]/hCA II and in [15N Leu]/hCA II form; the S50C-C206S construct – all expressions were carried out in E.coliBL21(DE3)pLysS cells by standard methods. Selective 15N Leu labelling was achieved by using a minimal medium and a mixture of 18 unlabelled amino acids (all standard amino acids with the exception of Leu and Asn) as the main culture medium. Simultaneously with IPTG induction, 60 mg/L 15N-labelled Leu was added. The harvested cells were lysed by three freeze thaw cycles and resuspended in 25 mL of lyso buffer (50 mM Tris-SO4, pH 8.0, 50 mM NaCl, 0.5 mM ZnSO4, 1 mM DTT and 10 µg/mL PMSF). The suspension was shaken vigorously (300 rpm) at room temperature (RT) for 30 min then deoxyribonuclease I (DNaseI, 1 µg/L) was added and the mixture was shaken for another 30 min. The lysate was centrifuged at 16880 g for 30 min at 4°C. The supernatant was recovered and the pellet was resuspended twice further in 25 mL lysis buffer. The total 75 mL solution from the extraction was filtered through a 0.45 µm filter and used directly for affinity chromatography. Affinity chromatography was performed using 25 mL of 4-(2-aminoethyl)benzenesulfonamid agarose resin packed into a XK16 column (GE Healthcare, Glattbrugg, Switzerland). The column was equilibrated with 5 column volume (CV) activity buffer (50 mM Tris-SO4, pH 8.0, 0.5 mM ZnSO4, 1 mM DTT) and the protein in 75 mL of lysis buffer was loaded onto the column at a slow flow rate (1 mL/min). Then the column was washed with 5 CV of wash buffer (50 mM NaSO4, 50 mM NaClO4, 25 mM Tris-SO4, pH 8.8, 1 mM DTT) and the protein was eluted with 10 CV of elution buffer (200 mM NaClO4, 100 mM NaAc, pH 5.6, 1 mM DTT). 10 mL fractions were collected and those containing the protein (detected by UV absorption) were pooled and dialyzed at 4°C against activity buffer for 12 h, followed by deionized H2O for 24 h, and finally against ultrapure water for another 24 h. Dialysis buffer contained 100 µM DTT and was exchanged at least three times a day. The resulting solution was frozen in liquid nitrogen and lyophilized. The resulting protein was stored at 4°C. The proteins were all characterized by SDS-PAGE (12% acrylamide, 200 V, 1.5 h) and by ESI-MS on a Bruker Daltonics microTOF instrument. Additionally, solution NMR spectroscopy was carried out on a Bruker Avance III HD spectrometer operating at 600 MHz proton frequency, equipped with a cryogenic QCI probe 1H/13C/15N/2H with z-axis pulsed field gradient.

**Synthesis of Epoxy-Functionalized Silica**

The epoxy-activated silica (epoxy-silica) was prepared according to previously established protocols. The SP-100-15-P Daiso silica gel (pore size of 100 Å, 5 g) was calcined at 200 °C under vacuum for 24 h. The freshly activated dry silica was then suspended in dry toluene (80 ml) and degassed by sonication under vacuum for 30 minutes. A ten-fold theoretical excess of GLYMO was added and the mixture was heated to reflux for 4 h. The amount of GLYMO added to the reaction mixture was calculated based on the specific surface area of silica (452 m²/g) and the wetting area of GLYMO (330 m²/g). After the completion of the reaction, the reaction mixture was cooled to room temperature. The resulting solid was washed with 250 ml of dry toluene, 125 ml of THF, and 250 ml of methanol, and finally dried under vacuum at 150 °C for 24 h. The epoxide content grafted onto the silica surface (epoxy equivalent weight, EEW) was determined by nonaqueous titration of the oxirane groups according to the modified Dubertret method.

**Covalent Immobilization of hCA II and Enzymatic Assays**

Covalent immobilization of hCA II on epoxy-silica was performed according to the previously established methods. Human carbonic anhydrase II was dissolved in 1.88 M ammonium sulfate, 0.010 M potassium phosphate (pH 8.0) resulting in 10 mg/mL protein solution. A 1 mL volume of the above solution was mixed with 100 mg of epoxy-silica in a 5 mL capped vial. The mixture was gently mixed for 40 h at room temperature using an orbital rotator. The modified support was recovered by centrifugation and washed twice successively with 1 mL of 0.010 M potassium phosphate (pH 7.0) and 2 mL of 0.010 M sodium acetate (pH 4.5) containing 0.3 M sodium chloride. Finally, the immobilized enzymatic system was equilibrated with 2 mL of 0.10 M potassium phosphate (pH 6.0). The amount of protein covalently immobilized on the epoxy-silica was calculated indirectly from the difference between the amount

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* Based on the information provided by the supplier of bare silica (Daiso Chemical Co. Ltd, Japan).
of protein added initially for immobilization and the amount of the enzyme in the supernatant after immobilization (including washings). Quantification of the protein was carried out using a spectrophotometer based on the colorimetric method of Bradford.\textsuperscript{21} The esterase activity of the hCA II was determined spectrophotometrically using p-nitrophenyl acetate (p-NPA) as a substrate, as described elsewhere.\textsuperscript{21} Briefly, the assay system comprised 0.2 mL of hCA II solution (0.1 mg/mL), or appropriate amounts of immobilized hCA II, in a 1 cm spectrophotometric cell containing 1.8 mL of potassium phosphate buffer (0.1 M, pH 7.0) and 1 mL of 3 mM p-NPA. For the free enzyme, the change in absorbance at 410 nm at 25 °C was recorded over the first 20 min. In the case of immobilized hCA II, the reaction mixture was stirred using a magnetic stirrer, the solution was centrifuged and the absorbance of the supernatant at 410 nm at 25 °C was measured every 3 minutes for the first 30 minutes. The product concentrations were corrected for the autohydrolysis of p-NPA by conducting blank experiments. The specific activity (μmol min⁻¹ mg⁻¹) of the hCA II in solution and in the immobilized state were calculated from these corrected product concentrations. Specific activity is defined as the micromoles of substrate transformed per minute per milligram of the total protein sample.

**Solid-State NMR Experiments**

Solid-state NMR experiments were performed on a Bruker Avance III 400 MHz spectrometer equipped with a widebore 9.4 T magnet, a Bruker Avance III 700 MHz spectrometer with a widebore 16.4 T magnet, and a Bruker Avance III 850 MHz spectrometer with a widebore 20 T magnet at room temperature. The powdered samples were packed into 1.0 and 3.2 mm ZrO₂ rotors and were spun at frequencies of 75 and 22 kHz in 1.0 and 3.2 mm MAS probes, respectively. Chemical shifts were referenced externally relative to TMS for ¹H (adamantane: 1.87 ppm), ¹³C (adamantane left peak: 38.4 ppm), ¹⁵N (glycine: 32.4 ppm) and ²⁹Si [octakis(trimethylsilyloxy)silsesquioxane left peak: 12.1 ppm]. For insensitive nuclei (¹⁰C, ¹⁵N and ²⁹Si), pulse sequences were employed with linearly ramped cross-polarization (CP)\textsuperscript{45} and with small phase incremental alternation (SPINAL)\textsuperscript{46} ¹H decoupling. All the NMR data were processed using TopSpin software. Further experimental and processing details can be found in the corresponding text and figure captions.

**RESULTS AND DISCUSSIONS**

**Protein Expression, Purification and Enzymatic Assays**

We successfully expressed four different double mutants of hCA II (SS0C-C206S, S166C-C206S, S217C-C206S, S220C-C206S) in [U-¹⁵N]/hCA II as well as [¹⁵N Leu]/hCA II forms in good to excellent yields (60 to 200 mg/L culture). All protein constructs could be obtained in pure, homogeneous form as indicated by SDS-PAGE (Figure S1, Supporting Information). Only two bands were observed, one around 30 kDa corresponding to hCA II and a second one at about 60 kDa which proves that the cysteine residues are solvent-accessible and can form disulfide-linked hCA II dimers. A similar observation was made in the ESI-MS (Figure S2, Supporting Information). For freshly DDT reduced [U-¹⁵N]/hCA II samples, only m/z = 29,449.7 was obtained, in excellent agreement with the mass calculated (m/z = 29,448.7) for C₁₁₆H₁₉₂O₇N₁₅S₁₅, corresponding to the loss of the N-terminal methionine and a ¹⁵N enrichment of 98%. On prolonged standing of the protein sample before MS analysis, m/z = 58,897.4 was also found, corresponding again to a loss of two hydrogen atoms and the formation of a disulfide bond.

The enzymatic activities of the free and immobilized hCA II were assayed by measuring the hydrolysis of p-NPA as substrate. A plot of the rate of product formation at increasing concentration of hCA II is shown in Figure S3. The measured specific activity of the immobilized enzyme 0.166 μmol min⁻¹ (mg immobilized preparation)⁻¹ was converted to a value in terms of the protein present, 1.68 μmol min⁻¹ (mg protein)⁻¹ using the measured protein loading of 9.9 mg protein per 100 mg support. The converted value was then compared with the measured value for the free enzyme 2.3 μmol min⁻¹ (mg protein)⁻¹ to show 71% retention of activity.

**Solid-State NMR Experiments**

Figure 1 shows the single 90° pulse ¹H MAS NMR spectra of bare silica (Figure 1a), epoxy-silica (Figure 1b), [¹⁵N Leu]/hCA II (Figure 1c), [U-¹⁵N]/hCA II (Figure 1d), [¹⁵N Leu]/hCA II immobilized on epoxy-silica (Figure 1e) and [U-¹⁵N]/hCA II immobilized on epoxy-silica (Figure 1f), all recorded at 75 kHz MAS on a B₀ = 20 T magnet.

![Figure 1](image-url)
groups (between 1 to 2 ppm), hydrogen-bonded silanol groups (4 ppm) and the broad peak (5 to 9 ppm) from the water molecules physically adsorbed onto the surface silanol groups. The spectrum of epoxy-silica (Figure 1b) is characterized by peaks appearing from H1 (0.8 ppm), H2 (1.5 ppm), H3, H4, H5, SiOCH3 (3.3 ppm) and H6ep (2.5 ppm) groups and is in good agreement with similar previous reports.

The numbering scheme for the protons and the carbons from the epoxy-silica (green labels) and from the immobilized hCA II (red labels) is shown in Figure 3c. As in the spectrum of bare silica, the spectrum of epoxy-silica (Figure 1b) is also characterized by a broad peak spanning from 5.0 to 9.0 ppm arising from the hydrogen-bonded water on the surface of the epoxy-silica. The width of the water peak can be ascribed to a lack of dynamics and the inhomogeneous broadening associated with a wide range of chemical shifts contributed by the different modes of hydrogen bonding with the surface silanol groups. The 1H NMR spectra of the [15N Leu]hCA II (Figure 1c) and [U-15N]/hCA II (Figure 1d) are almost identical and reveal mainly the signals from the aliphatic (0 to 3 ppm), water (4.7 ppm), and the aromatic and amino groups (6 to 10 ppm). On the other hand, the spectra of the immobilized [15N Leu]/hCA II (Figure 1e) and [U-15N]/hCA II (Figure 1f) are characterized by the signals appearing from the epoxy-silica (0 to 4.0 ppm), water (4.7 ppm) in addition to the signals (6.0 to 10 ppm) from the aromatic and amino groups from the immobilized enzyme. When compared to the spectrum of the epoxy-silica (Figure 1b), the spectra of the immobilized hCA II are characterized by the absence of the signals from the C6ep groups (2.2 ppm), possibly indicating the epoxy-ring opening reaction during the process of immobilization under aqueous conditions.

Further confirmation of the chemical shift assignments from the epoxy-silica before and after immobilization were made by recording 13C CPMAS experiments. Figure 2 shows the 13C CPMAS spectra of epoxy-silica (Figure 2a) and [15N Leu]/hCA II immobilized on epoxy-silica (Figure 2b). The spectrum of epoxy-silica (Figure 2a) is characterized by the peaks appearing from C1 (8.5 ppm), C2 (23.1 ppm), C3 (74.2 ppm), C4 (72.2 ppm), CSop (51.1 ppm), C6op (44.3 ppm) and -SiOCH3 (49.4 ppm) groups and is in good agreement with previous reports on a similar epoxy-silica. The spectrum of [15N Leu]/hCA II immobilized on epoxy-silica (Figure 2b) is characterized by the 13C signals from the aromatic (110 to 150 ppm), arginine side chain (zeta-carbon of the guanidine group; 160 ppm) and the carbonyl groups (175 ppm) from the protein, in addition to the signals from the modified epoxy-silica. Two distinct low 13C frequency shoulder peaks from the Ile side chains (Cε: between 15 to 20 ppm) could be also identified, in addition to the signals from the aliphatic region (between 20 to 40 ppm) from the immobilized hCA II. Furthermore, it is worth noting the relative decrease in the intensities of the peaks from the CSop (51.1 ppm) and C6op (44.3 ppm) groups and the appearance of new C6op (63.4 ppm) and C5 (40.4 ppm) peaks compared to the peaks associated with the remainder of the carbon atoms (C1, C2, C3, C4).

To further confirm the chemical shift assignments from the epoxy-silica and the immobilized hCA II on epoxy-silica, two-dimensional 1H-13C HETCOR (HETero-nuclear CORrelation) spectra were recorded. Cross-polarization relies on dipolar couplings and, as a result, the cross-peaks in the two-dimensional 1H-13C FSLG-HETCOR spectra appear from strongly dipolar-coupled nuclear spins and so can provide information about the spatial proximity of the atoms involved. The spectrum of the epoxy-silica is well resolved (Figure 3a) and is characterized by 1H-13C cross-peaks appearing from H1-C1 (0.5-6.4/8.4 ppm), H2-C2 (1.5-23.0 ppm), H3-C3 (2.8-74.1 ppm), H4-C4 (2.8-72.2 ppm), H5-C5 (2.7-51.1 ppm), H6-C6 (2.2-44.3 ppm) and SiOCH3 groups (2.8-49.6 ppm) in agreement with the single 90° pulse 1H and 13C CPMAS spectra. The presence of two C1 peaks in the spectrum of epoxy-silica can perhaps be attributed to the different bonding environments of the GLYMO with the surface silicon species, with the first C1 (6.4 ppm) peak assigned to the carbon atoms from the T1 silica species and the second C1 (8.4 ppm) peak assigned to the carbon atoms from the T2 species. The spectrum is also characterized by cross-peaks appearing from weakly dipolar-coupled 1H-13C spin pairs including H1-C2 (0.5-23.0 ppm), H3-C2 (2.8-23.0 ppm), H2-C3 (1.5-74.1 ppm) and H2-C4 (1.5-72.2 ppm). Furthermore, cross-peaks between the bulk 1H water signals (4.3 ppm) with C2, C3,
Figure 3. Two-dimensional $^1$H-$^{13}$C FSLG-HETCOR spectra of (a) epoxy-silica, (b) $[^{15}$N Leu]/hCA II immobilized on epoxy-silica in natural abundance and (c) simplified schematic representation of epoxy-silica surface before (top row) and after (bottom row) covalent immobilization with hCA II. Both spectra were acquired using homonuclear FSLG decoupling and a $^1$H-$^{13}$C CP contact time of 2 ms on a $B_0 = 16.4$ T spectrometer at a MAS frequency of 22 kHz. Spectrum (a) was acquired using 1024 transients for each of the 128 increments and spectrum (b) using 4096 transients for each of the 28 increments. High-power homonuclear and heteronuclear proton decoupling (~90 kHz) was applied during the FSLG $\tau$ intervals and also during $^{13}$C acquisition. Spectrum (a) was processed without window functions and spectrum (b) using 50 Hz of line broadening in the direct ($F_2$) dimension, with no line broadening in the indirect ($F_1$) dimension. Chemical shift assignments from epoxy-silica are shown by green labels, bulk water in blue labels, and from the fingerprint regions from the covalently immobilized hCA II using red labels.

C4, C5, C6 and SiOCH$_3$ groups are also visible in the spectrum, in addition to the cross-peaks between the $^1$H signals from the physisorbed water molecules (5.5 ppm) and the C3, C4, C5 and SiOCH$_3$ groups from the epoxy-silica. On the other hand, the spectrum of the $[^{15}$N Leu]/hCA II immobilized on epoxy-silica (Figure 3b) is characterized by $^1$H-$^{13}$C cross-peaks appearing from the fingerprint regions of the immobilized enzyme (red labels) in addition to the signals from the epoxy-silica (green labels). Successful immobilization of the hCA II on epoxy-silica is evident from the $^1$H-$^{13}$C cross-peaks appearing from the carbonyl and the aromatic groups in the protein in addition to the signals appearing from the epoxy-silica. The systematic absence of the C6$_{ep}$-H6$_{ep}$ cross-peak (44.3-2.2 ppm) from the epoxy ring and the appearance of a new C6$_{op}$-H6$_{op}$ cross-peak (63.4-3.7 ppm) indicates the hydrolysis of the epoxy ring during the enzyme immobilization under aqueous conditions. In similar previous work on chymotrypsin immobilized on epoxy-silica, it was possible to observe and assign $^{13}$C signals from the epoxy-silica and modified epoxy-silica, no distinct signals could be observed or assigned from the fingerprint regions from the immobilized enzyme. This is a consequence of the fact that natural abundance $^{13}$C CP MAS NMR of covalently immobilized enzymes on epoxy-silica can be extremely challenging due to the small amount of protein grafted onto the epoxy-silica surface. Furthermore, obtaining site-specific resonance assignments in natural abundance is additionally challenging due to the intense background signals arising from the epoxy-linker. Although the protein signals in the aliphatic region from the immobilized enzyme are masked by the intense background signals from the epoxy-silica, in this work chemical shift assignments from the immobilized hCA II can be made by directly comparing the chemical shifts from the epoxy-silica before immobilization. It is worth noting the overall distribution and dispersion of chemical shifts from the fingerprint regions of the immobilized enzyme, in particular the distribution of
the low $^{13}$C frequency signals from the Ile side chains ($C_\beta$ between 15 to 20 ppm) and the high frequency signals from the Glu/Gln side chains ($C_{\gamma,\delta}$: between 177 to 185 ppm). The spectrum in Figure 3a is also characterized by the reduced intensity of the $\text{-SiOCCH}_3$ (49.5-2.8 ppm) cross-peak, indicating extensive cross-linking of the silanol groups on the silica surface. This is in agreement with similar studies, where it has been reported that the silanol groups do not condense extensively during the initial silica surface modification but require the prolonged aqueous incubation during immobilization of the enzyme. This observation is also supported by the $^{29}$Si CPMAS spectrum (Figure S4, Supporting Information) of the immobilized enzyme, as this is mainly dominated by signals from the $T_1$ (-56 ppm) and $T_2$ silicon species (-66 ppm) in comparison with the spectrum of the epoxy-silica, which is mainly dominated by signals from the $T_1$ (-48 ppm) and $T_2$ (-56 ppm) silicon species. Upon covalent binding of the enzyme, the C6 carbon atom of the epoxy ring $C_{6p}$ (44.3-2.2 ppm) is shifted to low $^{13}$C frequency, labelled as $C_{6u}$ (40.1-4.2 ppm) (Figure 3b). Based on the chemical shift assignments, a simplified schematic representation of epoxy-silica surface before and after immobilization could be proposed (Figure 3c).

**Structural Changes of Bulk and Surface Silica Species**

The changes in the silica binding environments during different stages of enzyme immobilization were monitored by comparing $^{29}$Si CPMAS NMR spectra from the bare silica, epoxy-silica and the hCA II immobilized on epoxy-silica (Figure S4, Supporting Information). The $^{29}$Si CPMAS NMR spectrum of the bare silica (Figure S4a, Supporting Information) reveals mainly three different types of silicon species, $Q_1$ (-90 ppm), $Q_2$ (-100 ppm) and $Q_3$ (-110 ppm) in agreement with previous reports. The spectrum of epoxy-silica (Figure S4b, Supporting Information) shows new $T_1$ (-48 ppm) and $T_2$ (-56 ppm) species from the epoxy-silica in addition to the peaks from $Q_1$, $Q_2$ and $Q_3$ species and is also in good agreement with previous reports. The appearance of new $T_1$ and $T_2$ groups and the reduced intensity of the $Q_2$ and $Q_3$ peaks confirms that GLYMO is covalently grafted on the silica surface and not chemically or physically adsorbed. The spectrum of epoxy-silica is mainly dominated by $T_1$ and $T_2$ species indicating the absence of extensive cross-condensation of silanol groups on the silica surface as evident from the absence of signals from the $T_3$ species (-66 ppm). However, the $^{29}$Si CPMAS NMR spectrum of hCA II immobilized on epoxy-silica (Figure S4c, Supporting Information) shows mainly $T_1$ and $T_2$ species and is characterized by the systematic absence of $T_3$ species.

**Native Fold of the Protein before and after Immobilization**

Structural changes in the enzyme before and after immobilization were monitored by comparing $^{15}$N CPMAS NMR and two-dimensional $^1$H-$^{15}$N NMR FSLG-HETCOR spectra of [U-$^{15}$N]/hCA II and [U-$^{15}$N]/Leu/hCA II samples. The $^{15}$N NMR spectra of the lyophilized hCA II (black) and immobilized hCA II (red) are almost identical (Figure S5, Supporting information). The spectra of [U-$^{15}$N]/hCA II before and after immobilization (Figure S5a, Supporting information) consist of $^{15}$N peaks arising from the backbone amide (NH: 100 -140 ppm), lysine side chain ($N_\epsilon$: ~33 ppm), arginine side chains ($N_\epsilon$: ~73 ppm, $N_\pi$: ~85.0 ppm) and the histidine side chains ($N_\delta$: $N_\pi$: 155-185 ppm). On the other hand, the spectra of [U-$^{15}$N]/Leu/hCA II (Figure S5b, Supporting information) are characterized only by the backbone amide peaks (NH: 100-140 ppm) and the systematic absence of the side chain $^{15}$N signals from lysine, arginine and histidine residues, indicating the successful selective isotopic enrichment of leucine residues and the absence of any isotopic scrambling. It is worth noting that no significant chemical shift perturbations or line broadening could be observed from the backbone amide peaks (NH: 100-140 ppm) before and after immobilization for both [U-$^{15}$N]/hCA II and [U-$^{15}$N]/Leu/hCA II samples. Although the epoxy groups can react with different reactive amino acid residues from the protein, the preferential distribution of the exposed side chain amino groups from the lysine residues in solution (N) makes them favorable for reaction with the epoxy groups.

To obtain further structural details from the enzyme before and after immobilization, two-dimensional $^1$H-$^{15}$N FSLG-
The spectra (Figure 5a) of [U-\(^{15}\)N]/hCA II before (black) and after (red) immobilization are characterized by the protons from the histidine residues at the active site. The spectra (Figure 5b) of [\(^{15}\)N Leu]/hCA II before (black) and after (red) immobilization are similarly almost identical. Both spectra are characterized by \(^1H\)\(^{15}\)N cross-peaks between the backbone amide (NH: 100-140 ppm) and the protons from the aliphatic (2.0-4.0 ppm), bulk water (4.8 ppm), and backbone amide/aromatic groups (6-14 ppm). The overall distribution of chemical shifts from the protein before and after immobilization indicates that the structural integrity of the enzyme is not significantly altered upon covalent immobilization and is comparable to that of the lyophilized state. This result can be compared with the observation that the immobilized hCA II retained 71% of its enzymatic activity when compared to the free enzyme in solution.

It should be noted that lyophilized states of the proteins are not necessarily identical to the native structures in solution and available reports on lyophilized powders of proteins are controversial. Although some studies\(^{30,31}\) indicate significant reversible conformational change upon lyophilisation, others\(^{32,33}\) show that the lyophilized state is comparable and identical to that in solution. When compared to the crystalline state, the poor resolution from the lyophilized proteins observed by MAS NMR can be attributed to the introduction of conformational heterogeneity during the lyophilization process. Better insights about the native fold of the protein before and after immobilization might be obtained by comparing spectra (for example, \(^1H\)-detected two-dimensional \(^1H\)\(^{15}\)N HSQC experiments under fast MAS) of the micro/nano crystalline protein and of the immobilized state. Such experiments are an active area of research in our laboratory but are beyond the scope of this current manuscript.

**CONCLUSIONS**

In summary, we have shown that solid-state MAS NMR can be successfully employed to yield detailed structural information from a covalently immobilized heterogeneous biocatalytic system. Obtaining site-specific information from covalently immobilized proteins on silica using NMR at natural isotope abundance is challenging due to the limited experimental sensitivity arising from the small amount of protein grafted onto the surface of the silica support. Preparation of the hCA II samples in the [U-\(^{15}\)N]/hCA II and [\(^{15}\)N Leu]/hCA II forms has enabled us to gain additional structural insights from the enzyme before and after immobilization. \(^1H\) MAS NMR experiments at 75 kHz spinning rate at high magnetic fields (\(B_0 = 20\) T) have aided in revealing the different \(^1H\) environments from the epoxy-silica and the immobilized enzyme in the as-synthesized state. Confirmation of the chemical shift assignments from the epoxy-silica and the immobilized enzyme before and after immobilization were accomplished using one- and two-dimensional \(^13\)C CPMAS NMR experiments. A comparison of the two-dimensional \(^1H\)\(^{15}\)N FSLG-HETCOR spectra of the epoxy-silica and immobilized enzyme reveals that -SiOCH\(_3\) groups do not cross-link extensively on the surface of the silica during the synthesis of epoxy-silica; this cross-linking requires prolonged aqueous incubation during the immobilization. This was further confirmed by \(^29\)Si CPMAS NMR experiments where the spectrum of the epoxy-silica consists mainly of \(T_1\) and \(T_2\) species while the spectrum of the immobilized enzyme is mainly dominated by \(T_1\) and the \(T_2\) species. Most notably, comparison of the \(^15\)N CPMAS and two-dimensional \(^1H\)\(^{15}\)N FSLG-HETCOR spectra before and after covalent immobilization reveals that the structural integrity of the protein is not dras-
typically changed upon immobilization. This result can be compared with the observation that the immobilized enzyme system retained 71% of its effective specific activity when compared with the free hCA II in solution.

ASSOCIATED CONTENT
Supporting Information. SDS-PAGE of uniformly 15N-labelled hCA II - S50C-C206S before and after subsequent purification steps. Deconvoluted ESI-MS spectra of uniformly 15N-labelled hCA II - S50C-C206S after TCEP reduction. Plot of the rate of product formation at increasing concentration of hCA II. One-dimensional 29Si CPMAS spectra of silica, epoxy-silica and hCA II immobilized on epoxy-silica. One-dimensional 13N CPMAS NMR spectra of [U-13N]/hCA II and [3N Leu]/hCA II before and after immobilization on epoxy-silica. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interest.

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ABBREVIATIONS
hCA II, human carbonic anhydrase II; MAS, magic-angle spinning; CP, cross-polarization; DNase I, Deoxyribonuclease I; DTT, Dithiothreitol; E.coli, escherichia coli; ESI-MS, electrospray ionization mass spectrometry; HETCOR, hetero-nuclear correlation; i.e., immobilized enzyme; IPTG, isopropyl β-D-thiogalactopyranoside; FSLG, frequency-switched Lee–Goldberg; SDS, sodium dodecyl sulfate; GLYMO, (3-glycidoxypropyl)trimethoxysilane; EEW, epoxy equivalent weight; TMS, tetramethyloxysilane; PMSF, phenylmethylsulfonyl fluoride; SPINAL, small phase incremental alternation; TCEP, (tris(2-carboxyethyl)phosphine).

REFERENCES
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Epoxy functionalized silica

Covalently immobilized enzyme