DNP-supported solid-state NMR studies of $^{13}$C,$^{15}$N,$^{29}$Si-enriched biosilica of *Cyclotella cryptica* and *Thalassiosira pseudonana*

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**Abstract**

Solid-state NMR spectroscopy represents a powerful method for the investigation of diatom biosilica but detailed studies regarding its chemical composition and structural organization can be prohibited by insufficient spectroscopic sensitivity. Here, we used two-dimensional (2D) Dynamic Nuclear Polarization (DNP)-supported solid-state NMR experiments to obtain information about the molecular composition and supramolecular organization of proteins and carbohydrates in $^{13}$C,$^{15}$N,$^{29}$Si-labeled biosilica of *C. cryptica*. As a reference, we conducted DNP experiments on isotope-labeled biosilica of *T. pseudonana*. DNP-enhancement factors for different NMR signals, and thus, for different organic compounds, provide information about the supramolecular architecture of the biosilica. In addition, DNP-supported heteronuclear nitrogen-carbon correlation experiments allowed us to prove the presence of different structural elements of long chain polyamines (LCPAs) and revealed the occurrence of amine-nitrogen moieties exhibiting a correlation with carbonyl carbons that may indicate cross-linking of LCPAs to proteins as previously seen in studies on proteins extracted from other diatoms.

**Keywords** Solid-state NMR spectroscopy · DNP · Diatoms · *Cyclotella cryptica* · *Thalassiosira pseudonana* · Chitin · LCPAs

**Abbreviations**

2D Two-dimensional

AMUPol 15-[(7-Oxyl-3,11-dioxa-7-azadispiro[5.1.5.3]hexadec-15-yl)carbamoyl][2-(2,5,8,11-tetraoxatridecan-13-ylamino)][3,11-dioxa-7- azadispiro[5.1.5.3]hexadec-7-yl]oxidanyl

ASW Artificial seawater

*C. cryptica* *Cyclotella cryptica*

CP Cross polarization

DNP Dynamic nuclear polarization

EDTA Ethylenediaminetetraacetic acid

LCPA Long-chain polyamine

MAS Magic angle spinning

NC $^{15}$N–$^{13}$C correlation spectrum

NCC $^{15}$N–$^{13}$C–$^{13}$C correlation spectrum

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1 Introduction

Solid-state NMR spectroscopy is a powerful method for the investigation of biomaterials [1–7] including diatom biosilica [8]. Diatoms are unicellular algae. They are frequently studied model organisms for biominalization research and also of significant interest in the context of designing bioinspired materials [3, 8–11]. Their ability to build up micro- and nano-structured cell walls out of amorphous silica makes them an interesting model organism. Since various biomolecules show proximity to the silica phase, this biosilica may be understood as composite material. The silica-attached biomolecules include different proteins, e.g., sillafins, carbohydrates and long-chain polyamines (LCPAs). Especially different proteins and LCPAs are able to precipitate silica and are assumed to be involved in cell wall synthesis [12–16]. A mixture of different LCPAs which differ in chain-length and degree of methylation is usually found in diatom biosilica sample. Structure and amount of LCPAs are also species-specific [17, 18]. Note that LCPAs could even be found in sedimented diatomite samples from the ocean floor [19]. Previous studies show that especially LCPAs are embedded within the silica phase while proteins and carbohydrates can be mainly found on the surface of silica [3, 10]. The silica phase of the cell walls can be characterized using 29Si solid-state NMR spectroscopy and the organic compounds by 13C and 15N solid-state NMR spectroscopy [3, 8–11]. The sensitivity of such experiments is strongly enhanced by isotope-labeling which, as shown by Brunner et al. [8] can be elegantly achieved by diatom growth in isotope-enriched culture medium. In recent years, high-frequency Dynamic Nuclear Polarization [20] (DNP), in which polarization is transferred from free electrons to atomic nuclei has become a powerful method to enhance solid-state NMR signal intensities by up to two orders of magnitude. These advancements have greatly expanded the use of ssNMR to study complex biomolecular systems including membrane [21–26] and amyloid [27, 28] proteins as well as cellular preparations [29–33] and biomaterials [2–4, 34]. Such experiments not only enhance the prospects to study local molecular structure but also offer a spectroscopic means to probe local as well as supramolecular arrangements [27] as we have previously shown for diatom biosilica from Stephanopyxis turris [3].

In the present work, we have used DNP-ssNMR to examine isotope-labeled diatoms of Cyclotella cryptica biosilica [35], a centric diatom which has a massive insoluble organic matrix. This diatom species forms external β-chitin fibrils as well as silica-attached chitin. In particular, the use of DNP allowed us to study the presence of silica-attached chitin. In addition, we could spectroscopically probe LCPAs and we examined the organic–inorganic interface in C. cryptica. We compare our findings to experimental results obtained on Thalassiosira pseudonana, a centric diatom, which is one of the most studied diatoms and our previously published results on S. turris biosilica [3]. Interestingly, the amount of silica-attached biomolecules in S. turris is remarkably smaller compared to those of T. pseudonana and especially C. cryptica. The higher amount of biomolecules in the samples studied here in combination with the strong sensitivity gain provided by DNP allowed us to investigate the silica-attached biomolecules in additional detail.

2 Methods

2.1 DNP sample preparation of 13C, 15N, 29Si-enriched and 13C-enriched C. cryptica & T. pseudonana biosilica

Diatom species of both C. cryptica and T. pseudonana were cultivated as described previously [35]. Likewise, (13C, 15N, 29Si) isotope labeling, cell pellet cleaning and preparation of the biosilica samples using lysis buffer solution [0.1 M EDTA (ethylenediaminetetraacetic acid), 2 wt-% SDS (sodium dodecyl sulfate)] followed earlier procedures [10, 35, 36].
2.2 DNP-supported NMR experiments

2.2.1 NMR sample preparation using the incipient wetness method

3.2 mm sapphire rotors were filled with isotope-labeled C. cryptica or T. pseudonana biosilica which were wetted directly before measurements. The wetting of the biosilica was performed using the method of incipient wetness impregnation with the radical solution at room temperature [21]. The radical stock solution with a concentration of 15 mM was prepared by dissolving AMUPol in D$_2$O/H$_2$O 9:1. The solution was stored at -20 °C. 10.2 mg of dry biosilica were wetted with 50 µL AMUPol solution and immediately packed into the MAS rotor. Directly afterwards, the filled rotors were transferred to the NMR probe which was cooled to 92 K.

2.2.2 DNP-supported NMR experiments

DNP experiments were acquired using a $^1$H–$^{13}$C–$^{15}$N triple resonance probe at a static magnetic field of 9.4T corresponding to proton/electron resonance frequencies of 400 MHz/263 GHz (Bruker BioSpin). The temperature was set to 100 K and an MAS spinning speed of 9 kHz was used. The recycle delay was set to 2 s for all experiments (see, e.g. Ref. [37]). For the 1D cross-polarization (CP) spectra, 90 degree $^1$H, $^{13}$C and $^{15}$N pulses were applied using field strengths of 77 kHz, 43 kHz and 40 kHz, respectively. H–C CP matching was achieved by using a 62 kHz, 70–100% ramped 1H pulse and a 45 kHz r.f. 13C pulse for a contact time of 700 µsec. HN CP conditions were optimized using a 70–100% ramped pulse on protons with a 41 kHz field, a matching 15N pulse of 33 kHz and a transfer time of 1.4 ms. High power proton decoupling at 85 kHz was achieved using SPINAL-64 [38] during evolution and acquisition in all experiments. For 2D proton-driven spin diffusion (PDSD) experiments, Carbon–carbon mixing was established with a mixing time of 50 ms. For the isotope-enriched C. cryptica biosilica sample, 224 scans were acquired with acquisition times of 15 ms and 6 ms for the direct and indirect dimensions, respectively. For the isotope-enriched T. pseudonana biosilica sample, 64 scans were acquired with acquisition times of 15 ms and 7 ms for the direct and indirect dimensions, respectively. In both cases two spectra were recorded, with and without DNP, and 1D slices were used to calculate the DNP enhancement. The $^{15}$N–$^{13}$C correlation experiment was performed using a H–N CP step of 1 ms and a SPECIFIC-CP [39] transfer step of 2.5 ms. Acquisition times were 8 ms and 7.5 ms for the direct and indirect dimensions, respectively. 256 scans were acquired and the spectrum was processed using a 0.5 π shifted sine squared window function in both dimensions The $^{15}$N-edited PDSD experiment [40] was recorded using a SPECIFIC-CP transfer step of 2.5 ms which is typical for one-bond NC correlations experiments [39] and a carbon–carbon mixing time of 50 ms. 1728 scans were acquired with acquisition times of 10 ms and 4 ms for the direct and indirect dimensions, respectively. The center frequencies were set to 50 ppm for $^{15}$N and $^{13}$C dimensions. The spectrum was processed using a 0.33 π shifted sine squared window function in both dimensions.

2.3 Scanning electron microscopy

SEM samples were prepared on an aluminum sample holder with an aluminum tape on top. A small amount of the biosilica sample was suspended in water and 50 µL of the suspension were transferred to the sample holder. After drying overnight at 40 °C, the sample was sputtered with 5 nm gold using a Q150R ES sputter coater from Quorum Technologies. SEM images were acquired on a Hitachi SU 8020. An acceleration voltage of 5 kV was used for acquisition. Secondary electrons were used for detection of surface topography. Images with a magnification of 20,000 and 90,000 were acquired.

3 Results and discussion

In our previous study [35] of C. cryptica using conventional 1D and 2D ssNMR spectroscopy, we observed that a high amount of proteins is associated to the biosilica and we detected two different chitin conformations (Fig. 1). In the following, we used DNP-supported NMR experiments to further enhance ssNMR signal intensities. As mentioned above, DNP relies on spin polarization transfer from electron spins to nuclei. In our experiments, samples were impregnated with a “DNP juice” (see methods) containing the biradical AMUPol [41]. First, we conducted one-dimensional $^{13}$C and $^{15}$N detected CP experiments (Fig. 2) under DNP conditions and, as in our previous study [3], defined the DNP enhancement factor by the ratio of signal intensities with and without microwave irradiation. In $^{13}$C CP experiments, we observed enhancement factors that (except for carbohydrates and possibly some methyl signals, see also Additional file 1:
Figures 1–4) ranged between 18 and 26 (Fig. 2, left). In the 15N CP measurement, we observed an enhancement of 26 for both amide and amine 15N signals (Fig. 2, right).

As we have discussed elsewhere [35], *C. cryptica* contains a massive organic matrix and produces extracellular chitin fibrils of ca. 50 nm thickness [42, 43] (see also Fig. 1). To further investigate molecule-specific DNP enhancements, we...
performed a two-dimensional DNP-enhanced PDSD (Proton-Driven Spin Diffusion) experiment (Fig. 3). We observed similar $^{13}\text{C}^{13}\text{C}$ correlations as in the standard PDSD experiment [35], however with considerably higher signal intensities. This improvement enabled the detection of new signals which are not observable in the conventional experiment. In particular, two spin systems were observable (Fig. 3, blue box). One spin system shows correlations from 92 to 81 ppm, 75 ppm, and 72 ppm. The other system exhibits connectivities between 95 and 81 ppm, 75 ppm, 73 ppm, as well as 70 ppm. These correlations are characteristic for carbohydrates, where the C1 is not connected. These unconnected anomeric carbon atoms characteristically give rise to peaks between 90 and 100 ppm [44]. Both spin systems also contain a shift higher than 80 which indicates a substitution on the carbocycle [45] by e.g. another sugar. This points towards the presence of reducing ends of polysaccharides [46] or would be consistent with sugar–protein linkages which often yield lower chemical shifts for the anomic carbon than a connection to another sugar (see, e.g. Ref. [47]). The different line widths of the resonance peaks of the sugar spin systems compared to the much narrower chitin peaks can be explained by two independent effects. Firstly, the relative position and exposure of the molecules/sugars towards the radical increased T$_2$ relaxation rates due to paramagnetic relaxation enhancement (PRE), resulting in broader NMR lines. Secondly, it could be an indication of different morphologies of the sugar networks, where there is crystalline chitin present with narrow line widths due to the increased isotropy in crystals as well as amorphous parts of chitin and potentially other sugars, which then exhibit broader line widths due to their anisotropy especially under freezing conditions. A monosaccharide analysis which provides information about the total carbohydrate composition revealed glucose and mannose as major part of the monosaccharides fraction. Furthermore, xylose, ribose, fucose, galactose and glucosamine are present in significant concentrations [35]. However, a differentiation between monosaccharides and monosaccharide units from polysaccharides is not possible due to the performed hydrolysis step.

Moreover, the determination of DNP enhancement factors for different signals, which are characteristic for different organic compounds, can help to probe the supramolecular architecture of the biosilica [3]. To distinguish DNP enhancements of protein and carbohydrate (including chitin) signals, we analyzed different 1D slices of the 2D spin diffusion experiment (Fig. 3, right). For the protein signals, we examined a 1D slice at 53 ppm which is typical for protein backbone C$_\alpha$ resonances and observed a strong enhancement factor of 26 in line with our 1D $^{15}\text{N}$ CP studies (Fig. 2). Thus, the proteins should be located at the solvent-accessible surface of the biosilica, where the radical solution (AMUPol) can polarize nuclei located in the biomolecules [22]. In contrast, for the carbohydrates, e.g. the chitin signals at 105 ppm, we determined significantly lower enhancement factors between 5 and 8 which is close to the $^{13}\text{C}$ enhancements seen in 1D CP MAS for carbohydrates (Fig. 2). Interestingly, when analyzing the carbohydrates with diagonal peaks at 92.4 and 95.7 (see also Additional file 1: Figure S2) we observed enhancement factors of 19 and 22–26. Their relatively high enhancement, especially when compared to the chitin signals, can —similar to the case of proteins— be explained by their

![Fig. 3](image_url)  

**Fig. 3**  DNP-supported Proton-Driven Spin Diffusion (PDSD) experiment of isotope-enriched *C. cryptica* biosilica using a mixing time of 50 ms and an MAS rate of 9 kHz. For better resolution, the enhancements were determined from 1D slices in the fingerprint region for protein C-alpha (slice A, 53 ppm, enhancement = 26) and characteristic for carbohydrates (Slice B, 105 ppm, Enhancement = 5–8)
location on the surface of the frustule where sugar moieties that are close to the radical are more strongly enhanced than the ones that are further away from it or buried/shielded like parts of the chitin.

As discussed elsewhere [3, 27, 48], the DNP enhancement in complex biomolecules depends on the internuclear distances between hydrogens responsible for the polarization transfer via proton-driven spin diffusion and the macromolecular layer size. In addition, the DNP enhancement for biomolecules close to the radical, may itself be modulated by the details of the local electron-nucleus geometry. This geometry determines the spin-diffusion barrier [49], influences the DNP magnetic field dependence of the DNP effect [50] and leads to paramagnetic line broadening [22, 51] within a sphere of about 1 nm around the DNP radical [22]. We expect this effect to be particularly strong for biosilica such as from S. turris which are characterized by small protein/organic surface layers [3]. Indeed, DNP signal enhancements for polysaccharides and polyamines amounted to about 40% to the protein signal enhancements for S. turris [3]. In contrast, the organic layer for C. cryptica is significantly larger [35] which results in almost uniform DNP enhancements in 1D 13C CP MAS data and is possibly the reason for the larger (relative) decrease in DNP enhancement from 26 to 5–8 for chitin.

Note that the increased line broadening often observed at low-temperature DNP conditions [22] prevents the differentiation between the α-like chitin- and β-chitin signals as it was possible without DNP and at ambient sample temperatures [35]. On the other hand and as mentioned earlier, C. cryptica contains a massive organic matrix and produces extracellular chitin fibrils of ca. 50 nm thickness [42, 43]. Hence, a more detailed analysis employing a classical spin-diffusion approximation to correlate relative DNP enhancements with biosilica layer thickness as performed earlier for labeled diatom biosilica of S. turris [3] was not attempted here. Instead we can draw the following general conclusions. Firstly, we would expect from previous theoretical work from our laboratory [48] only minor changes for the DNP enhancement for chitin embedded in the extracellular chitin fibrils compared to the protein layer. On the other hand, fluorescence spectroscopy [52] has shown that chitin in C. cryptica biosilica can also be directly associated to the siliceous cell walls. For this species, that is potentially surrounded by a low density of hydrogens that limit the spin-diffusion process, low DNP enhancements as found in Fig. 2 may be possible. Since ssNMR detects the entire ensembles of chitin moieties in our sample, the latter species may hence represent a prominent fraction in our C. cryptica preparations.

To further investigate the carbohydrate association to the siliceous cell walls, we conducted similar experiments on isotope-enriched T. pseudonana biosilica (Fig. 4) for which the presence of a chitin-based meshwork is already known [11]. Again, we analyzed 1D slices typical for protein and carbohydrate signals and observed a strong enhancement factor of 32 for the protein Cα signals (signal at 53 ppm). In line with our earlier observations, the proteins are thus predominately located at the solvent-accessible surface of the biosilica, where the radical solution (AMUPol) can polarize nuclei located in the biomolecules. Remarkably, for the chitin signals resonating at 105 ppm, a lower enhancement compared to the

Fig. 4 DNP-supported Proton-Driven Spin Diffusion (PDSD) experiment of isotope-enriched T. pseudonana biosilica using a mixing time of 50 ms and a MAS rate of 9 kHz. For better resolution, enhancements were again determined from 1D slices in the fingerprint region for protein C-alpha (slice A, 53 ppm, enhancement = 32) and characteristic for carbohydrates (Slice B, 105 ppm, Enhancement = 14–15)
protein signals is again observed. However, the relative enhancement of about 15 is significantly larger than in the case of *C. cryptica* biosilica (amounting to 5–8). These findings would be consistent with the notion that the organic matrix of *T. pseudonana* biosilica is smaller than in the case of *C. cryptica* [35] but larger than for *S. turris*. Such an interpretation would be consistent with previous findings [17, 18, 53].

Moreover, our DNP experiments revealed the presence of additional carbohydrate species as seen before for *C. cryptica* (Fig. 3). Interestingly, one of the observed new sugar spin systems exhibits a correlation of the anomic carbon to a signal around 165 ppm (Additional file 1: Figure S5). Sp²-configured carbon atoms of nitrogen containing functional groups including imines [54] and guanidiniums [55] as well as amide carboxyls give rise to peaks in this region. This correlation could hence indicate a cross linking between a carbohydrate C1 and an amino acid such as the guanidinium containing arginine, which has rarely been described as part of N-glycans. Another possibility would be the presence of citrulline as its carboxamide gives rise to resonances at around 164 ppm and it has been described as being linked to polysaccharides before [56]. This link could represent the interface of the sugar layers with the residual organic matrix.

Finally, we used DNP-supported ¹⁵N–¹³C correlation experiments to further investigate proteins as well as the presence of LCPA in our preparations. Long chain poly amines are based on putrescine, spermidine or 1,3-diaminopropane with a number of propylenediamine repeats ranging from 6–20. They are often positively charged and thereby able to attract negatively charged molecules, by which they are able to facilitate rapid silica precipitation. Since they are species specific their involvement in silica patterning is highly likely [57]. Again, DNP greatly facilitated such experiments because of the increased spectroscopic sensitivity, thereby reducing measurement times. To study nitrogen containing ¹³C moieties, we performed ¹⁵N filtered experiments [39, 40]. Figure 5 (red spectrum) shows a ¹⁵N–¹³C correlation spectrum. The observed correlations are characteristic for those of primary, secondary and tertiary amines to their directly adjacent carbon atom. The introduction of a spin diffusion (mixing) step between carbons allowed us to study correlations of the amines to more distant ¹³C atoms (blue spectrum). An increasing number of correlations occur, e.g., correlations to alkyl groups at lower ¹³C chemical shift between 15 and 30 ppm which would be consistent with correlations for both Lysine sidechains as well as the above mentioned structural elements of LCPAs. Due to the expected manifold of LCPAs [57] as well as the increased spectral line width under DNP conditions, we were however, not able to isolate individual spin systems or deduct exact structures for different LCPAs. Interestingly, we even observed a correlation to carbonyl-carbons at 174 ppm. This signal occurs probably due to relay transfer between close-by ¹³C labeled atoms to a carbonyl atom as usually seen in fully labeled protein samples under MAS conditions (see e.g. Ref. [58]) Such a correlation between the amine of a LCPA and a carbonyl group was not observed for other diatom biosilica, including diatom biosilica from *S. turris* [3]. However, it is well known from the model organism *T. pseudonana* [11] that LCPAs can be covalently attached to lysine residues or posttranslational modified hydroxylysine residues that may give rise to the correlation seen at 174 ppm. Additionally, the chemical shift of this carbonyl group is more typical for carbonyl groups of proteins. The above discussed

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**Fig. 5** DNP-supported NC and NCC NMR correlation experiments of isotope-enriched *C. cryptica* biosilica. In both experiments, ¹⁵N–¹³C transfer was established using a SPECIFIC-CP transfer step of 3 ms. In the NCC correlation experiment (blue) a mixing time of 50 ms was used for proton-driven spin diffusion (PDSD)
chitin should give rise to a signal at a higher chemical shift of ca. 176 ppm. Thus, proximity between amine nitrogens and protein carbonyl groups is more likely here.

4 Conclusions

Our DNP-supported solid-state NMR studies provide further insight into the composition and architecture of *C. cryptica* biosilica and allow us to draw the following conclusions:

(i) We could confirm the presence of chitin seen in ssNMR conducted at ambient temperatures [35] and we were also able to isolate two additional carbohydrate spin systems using PDSD experiments performed under DNP conditions. Their chemical shifts point towards the presence of either terminal reducing ends of polysaccharides or monomers that form a cross-connection between polysaccharides and proteins.

(ii) We also observed a reduced DNP efficiency for chitin compared to signals from proteins and other carbohydrates in preparations of *T. pseudonana* biosilica. However, this reduction was lower than for the case of *C. cryptica* biosilica. We attribute these variations to an increased size of the organic layer in the case of *C. cryptica* biosilica. Their different enhancements can be further correlated to their relative position on the diatom surface and their structural properties. Paramagnetic quenching due to the presence or binding of DNP radicals has the smallest influence on the observed protein signal under DNP conditions.

(iii) DNP-supported $^{15}$N–$^{13}$C 2D correlation experiments are consistent with the presence of different structural elements of LCPAs. Interestingly, a correlation of amine-nitrogen to a carbonyl carbon can be observed which may reflect chemical linkage to proteins as previously seen in studies on *T. Pseudonana* [59, 60].

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Authors' contributions  HLE, FK and ALP performed and analyzed the DNP solid-state NMR experiments. All authors contributed to writing the paper. All authors read and approved the final manuscript.

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Availability of data and materials  All data generated or analyzed during this study are included in this published article and its additional information files.

Declarations

Ethics approval and consent to participate  Not applicable.

Consent for publication  Not applicable.

Competing interests  The authors declare no financial interests.

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