Deuteration of proteins boosted by cell lysates: high-resolution amide and $H\alpha$ MAS NMR without re-protonation bottleneck

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Abstract. Amide-proton detected magic-angle spinning NMR of deuterated proteins has become a main technique in NMRbased structural biology. In standard deuteration protocols that rely on D_2O -based culture media, non-exchangeable amide sites remain deuterated, making these sites unobservable. Here we demonstrate that proteins produced with H₂O-based culture medium doped with deuterated cell lysate allow to overcome this "reprotonation bottleneck", while retaining a high level

- 5 of deuteration (ca. 80 %) and narrow line widths. We quantified coherence life times of several proteins prepared with this labelling pattern over a range of MAS frequencies (40-100 kHz). We demonstrate that under commonly used conditions (50-60 kHz MAS), amide ¹H line widths with our labelling approach are comparable to those of perdeuterated proteins and better than those of protonated samples at 100 kHz. For three proteins in the 33-50 kDa size range many previously unobserved amides become visible. We report how to prepare the deuterated cell lysate for our approach from fractions of perdeuterated cultures
- 10 which are usually discarded, and show that such media can be used identically to commercial media. The residual protonation of H α sites allows for well-resolved H α -detected spectra and H α resonance assignment, exemplified by the *de novo* assignment of 168 H α sites in a 39 kDa protein. The approach based on this H₂O/cell-lysate deuteration and MAS frequencies compatible with 1.3 or 1.9 mm rotors presents a strong sensitivity benefit over 0.7 mm/100 kHz MAS experiments.

1 Introduction

- 15 The deuteration of proteins for NMR studies is a very important technique and has greatly expanded the scope of biomolecular NMR. In solution-state NMR, overall Brownian motion modulates the relaxation-active interactions, in particular the strong ¹H-¹H dipolar interactions and chemical-shift anisotropy (CSA), and when this Brownian motion is slow, which is the case for large proteins, the resulting decay of coherences is fast. This rapid relaxation and the associated line broadening represent a fundamental barrier to the size of proteins that can be studied. It can be overcome by using highly deuterated proteins, where
- 20 only a few sites, e.g. amides, methyls or aromatic sites, bear a ¹H spin, while all the rest is deuterated (²H). Together with transverse-relaxation optimised pulse sequences tailored for amides (Pervushin et al., 1997), methyls (Tugarinov et al., 2003) or aromatics (Pervushin et al., 1998), deuteration has alleviated the size limitations of solution-state NMR, and has made proteins in the megadalton range accessible to site-resolved studies of dynamics and interactions.

In magic-angle spinning (MAS) solid-state NMR, overall molecular tumbling, that dominates relaxation in solution is absent,

- 25 and the transverse relaxation is inherently slow. However, the dipole-dipole interactions are not perfectly averaged by magicangle spinning, and these interactions lead to a rapid decay of the observable spin coherences. Traditionally, biomolecular MAS NMR has, thus, focused on observing ¹³C rather than ¹H. The smaller gyromagnetic ratio of ¹³C and accordingly the smaller dipolar couplings lead to a more complete averaging of the dipolar couplings already at lower MAS frequency. However, it also comes with inherently lower detection sensitivity than ¹H. In order to achieve narrow ¹H line widths, diluting the network
- 30 of ¹H-¹H dipolar interactions by deuteration is a well established method. Deuteration with sparse introduction of ¹H nuclei only at e.g. amide (¹H^N), H α , methyl or aromatic sites, has rapidly established itself as a standard technique in biomolecular MAS NMR. The higher detection sensitivity of ¹H translates to smaller sample amounts needed. Pioneering experiments e.g. by the Reif group used perdeuterated proteins in which only a fraction (e.g. 10 %) of the amides are protonated, thereby sacrificing signal intensity, at MAS frequencies of 20 kHz, i.e. with 3.2 mm large rotors and ca. 30 mg material (Chevelkov
- 35 et al., 2006). Faster MAS with smaller-diameter rotor allows to introduce more protons while keeping narrow lines; 100% amide-protonation with MAS frequencies of 40-60 kHz, i.e. smaller diameter rotors of either 1.3 mm (ca. 2.5 mg material) or 1.9 mm (ca. 13 mg) has become of widespread use (Lewandowski et al., 2011; Zhou et al., 2007). Akin to solution-state NMR, amide-¹H-detected MAS NMR experiments are employed for many aspects of protein structure and dynamics studies: they are used to obtain backbone resonance assignment (Barbet-Massin et al., 2014; Fraga et al., 2017; Xiang et al., 2015;
- 40 Stanek et al., 2020; Andreas et al., 2015; Klein et al., 2022), determine structures from ¹H^N-¹H^N distances (Retel et al., 2017; Zhou et al., 2007; Knight et al., 2012; Linser et al., 2011b; Najbauer et al., 2022; Gauto et al., 2019a; Schubeis et al., 2020) or long-distance restraints to paramagnetic (Knight et al., 2012) or ¹⁹F (Shcherbakov et al., 2019) sites, or to characterise protein dynamics (Napoli et al., 2023; Singh et al., 2019; Bonaccorsi et al., 2021; Good et al., 2014; Lewandowski, 2013; Chevelkov et al., 2009) (We refer the reader to excellent recent reviews of ¹H-detected MAS NMR and deuteration (Le Marchand et al., 2019)
- 45 2022; Vasa et al., 2018; Reif, 2021).) Very high MAS frequencies available today with 0.5 mm or 0.7 mm rotors (up to 150 kHz) alleviate the need for deuteration, and protonated proteins are sufficient for structure determination or dynamics studies (Agarwal et al., 2014; Andreas et al., 2016; Lamley et al., 2014; Bougault et al., 2019; Cala-De Paepe et al., 2017). However the absolute detection sensitivity drops substantially due to the smaller rotor volume; for example, using a 0.7 mm rotor instead of a 1.3 mm rotor, each close to its respective maximum MAS frequency, leads to a loss of at least a factor of 2.5 in sensitivity
 50 (or 6 in experimental time) (Le Marchand et al., 2022). Thus, for many practical applications, the use of deuterated proteins
- with MAS frequencies in the 40-60 kHz range (1.9 or 1.3 mm rotors) is still a preferable method.

Deuterated proteins for amide-detected experiments are commonly produced by bacterial overexpression in a minimal growth medium (Gardner and Kay, 1998) (M9 medium (Anderson, 1946)) prepared with D_2O . In the following we refer to this method of deuteration in D_2O based growth media as perdeuteration. As the amide hydrogen atoms are chemically

55 exchangeable, those accessible to the solvent are replaced with the 1 H isotope when the perdeuterated protein is placed in H₂O during purification and subsequent measurement. This amide re-protonation is, however, often incomplete, in particular in the core of large proteins, which leads to lack of information for these parts of the structure. As a consequence of the incomplete

back-exchange of amide sites, many structurally important probes are invisible. For example, in the structure determination of TET2 (Gauto et al., 2019a), for a buried β sheet only very few distance constraints were available.

- 60 One option to circumvent the need for re-protonation of amide sites is to focus on H α sites instead. An elegant approach has been proposed by the Andreas group, which starts from a commercial growth medium, which contains a mix of deuterated amino acids, and makes use of transaminases to protonate the H α positions (Tekwani et al., 2019), in an amino-acid-type dependent manner. Over-expression is then performed with this mix of deuterated H α -protonated amino acids. The method, named "alpha proton exchange by transamination" (α -PET), has been shown to generate proteins that are protonated predominantly at
- 65 the α sites. A potential drawback is, however, that H α sites generally have narrower signal dispersion than amide sites, such that resolution is often lower. Moreover, dynamics studies using the C α site are more complicated, because the ¹³C-¹³C couplings render any quantitative analysis of relaxation complex (if not impossible); dynamics studies using ¹⁵N sites are preferable.

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Denaturation of the perdeuterated protein in H_2O , followed by refolding is one viable route to achieve complete re-protonation of amide sites in perdeuterated proteins (Gardner et al., 1998). However, it is often difficult or even impossible to find suitable conditions in which the protein retrieves its native structure.

An alternative way to obtain (at least partly) deuterated proteins in which all exchangeable hydrogen sites bear a ¹H isotope is to perform the deuteration in H₂O, while providing the building blocks that serve for protein synthesis in a deuterated form. A simple approach is to use deuterated D-glucose as sole carbon source in H₂O-based minimal (M9) medium; this approach has been proposed for MAS NMR and termed inverse fractional deuteration (iFD) (Medeiros-Silva et al., 2016). As the entire

- ⁷⁵ biosynthesis of amino acids from glucose takes place in H₂O, this approach results in relatively low deuteration levels; in particular, the α sites have been reported to be protonated to 88-100 % for all reported amino acids (Table S2 of (Medeiros-Silva et al., 2016)). Due to the low overall deuteration the coherence life times of the amide ¹H spins in the iFD scheme are close to those of protonated samples, i.e. the gain in resolution with this sparse deuteration is limited (Cala-De Paepe et al., 2017).
- Asami *et al.* have proposed another approach, which primarily aims to introduce aliphatic protons in a sparse manner, but which also results in partial amide-protonation. The approach, termed "random adjoining protonation", consists of growing bacteria in a mixture of H_2O and D_2O and protonated ¹³C glucose (Asami et al., 2010, 2012). The aim is not to achieve highest possible deuteration at non-amide sites, but rather some level of aliphatic protonation, for detection of e.g. methyl sites. Thus, the approach is not optimised for amide detection.
- Using deuterated amino acids in the growth medium promises to result in more complete deuteration. Löhr *et al.* grew cultures in H₂O-based medium with a commercial deuterated algal lysate (without adding glucose) for solution-state NMR study of a 35 kDa-large protein (Löhr et al., 2003). A similar strategy has been reported by the Wand group (O'Brien et al., 2018). This approach has not been reported for MAS NMR, to our knowledge. A significant drawback of this method is the price. The lysate that Löhr *et al.* used has a current list price of US\$ 6500 for 1 litre of culture (Dec 2023), which may be
- 90 considered prohibitively expensive in many cases. Moreover, the amino acids are partly re-protonated due to the action of transaminases, and this protonation largely varies depending on the amino acid type; the residual protonation at α sites can reach 80 % (Löhr et al., 2003). Besides bacterial expression, cell-free protein production in H₂O with deuterated amino acids

can be performed. When used with transaminase inhibitors, the level of α protonation can be maintained below ca. 10 % (Imbert et al., 2021; Xuncheng et al., 2011). However, few laboratories have established cell-free production, and this approach is less readily accessible than bacterial over-expression.

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Here, we propose an approach for producing highly (> 80 %) deuterated proteins in H₂O-based M9 medium with deuterated D-glucose supplemented with deuterated cell lysate. We find that 2 g of cell lysate powder added to the medium suffice to reach 80 % overall deuteration level, and this level is ca. 4 times higher than the one obtained with the previously proposed iFD approach (Medeiros-Silva et al., 2016). While commercial deuterated media can be used, we report the straightforward preparation of a home-made cell lysate from what is usually discarded from perdeuterated bacterial protein production, and

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Moreover, we investigate the ¹H coherence life times of several proteins produced with this H₂O/M9/cell-lysate deuteration scheme, and compare them to those of perdeuterated and fully protonated protein samples over a range of MAS frequencies up to 100 kHz. Most importantly, we report that the amide ¹H line widths of the $H_2O/M9$ /cell-lysate deuterated samples are

- not significantly different from those of perdeuterated samples (although the coherence life times, i.e. the homogeneous line 105 widths(Le Marchand et al., 2022), of the perdeuterated samples are favorable). Using three proteins with molecular weights ranging from 33 to 50 kDa in 1.3 mm or 1.9 mm rotors (38-55 kHz MAS frequency), we show that our approach retrieves many amide signals that were lost in a standard perdeuteration approach. For the 12 x 39 kDa large protein assembly TET2, we obtained 40 % more amide assignments than with the previous perdeuteration approach, most of which in the hydrophobic
- core of the protein. We show furthermore that the residual "unwanted" protonation of α sites can be exploited in 3D and 4D 110 experiments that edit the H α frequency. We obtained 168 H α chemical-shift assignments, which is close to the number obtained from a fully protonated sample at 100 kHz MAS.

2 Results

2.1 Deuteration of proteins in H₂O-based cultures with deuterated algal lysate

show that its properties are indistinguishable from a commercial medium.

- The deuteration level and deuteration pattern (i.e. which sites are deuterated) is essential for the resolution in amide-hydrogen 115 based NMR MAS spectra. We first quantified the overall deuteration level that can be reached using H₂O-based bacterial cultures producing the 33.55 kDa protein MalDH, by determining the intact protein mass with mass spectrometry (Figure 1a). When only deuterated glucose is used (at 2 g per litre of culture; this corresponds to the previously proposed "iFD" approach (Medeiros-Silva et al., 2016)), the resulting protein is deuterated to ca. 25% overall. The addition of deuterated algal lysate-
- derived complex labeling medium increases the deuteration level in a concentration-dependent manner: upon addition of ca. 2 120 g/L of ISOGRO[®] powder to the medium, a plateau level of ca. 80% deuteration is reached. Doubling the amount of ISOGRO[®] added to the medium to 4 g/L has only very small effect, such that we consider 2 g/L as a sufficient amount. For brevity, we refer in the remainder of this text to the deuteration that uses H₂O culture medium with deuterated M9 components (including 2 g/L deuterated D-glucose) and deuterated cell lysate (at 2 g/L) as "H2O/M9/lysate" sample. With lysate we refer either
- 125 to the commercial complex growth medium (such as ISOGRO[®]) or a in-house made variant of it (see below). We refer to

"perdeuteration" as the approach using D_2O -based M9 medium using 2 g/L deuterated D-glucose. In all cases, the final NMR samples are in a buffer composed of H_2O .

Having quantified the overall deuteration level, we investigated more closely the pattern of the residual protonation. To this end, we have produced the 8.6 kDa protein ubiquitin with different labelling patterns: either no deuteration (u-[¹³C,¹⁵N]),
perdeuteration (u-[²H,¹³C,¹⁵N]), or deuteration with the H₂O/M9/lysate approach proposed herein. For the latter, the M9 culture medium was in H₂O and included 2 g/L ²H,¹³C glucose, 1 g/L ¹⁵N ammonium and 2 g/L ²H,¹³C,¹⁵N ISOGRO[®] powder. We measured the deuteration level with ¹H-¹³C HSQC spectra, which provides simultaneously the deuteration levels for all aliphatic sites.

Figure 1b and c show the protonation levels in aliphatic side chains and at H α positions. Marked differences were found 135 for H α deuteration of different amino acid types, qualitatively similarly to previously reported data (Löhr et al., 2003). For example, while Glu and Phe have a high incorporation of ¹H α , Lys and Ala have a higher deuteration level. Positions further out in the side chain have a higher deuteration level. Overall, our data show that there is significant variation in the deuteration level, which we ascribe to the activity of transaminases that differs for the types of amino acids. While the residual protonation may also be of possible use, namely for ¹H detection of aliphatic hydrogen sites, as we will show below, the presence of these

¹H spins in vicinity to amide protons will likely accelerate the coherence decay of amide ¹H spins. This question of coherence life times and line widths is addressed in a later section in this manuscript.

2.2 Straightforward preparation of an isotope-labelled cell lysate as an alternative to commercial media

Having shown that a commercial deuterated algal extract (e.g. ISOGRO[®] from Sigma-Aldrich or Celtone[®] from Cambridge Isotope Laboratories or SILEX[®] media from Silantes) allows obtaining high deuteration levels in H₂O-based bacterial expression, we have investigated the possibility to prepare such a medium in house. In many laboratories, deuterated proteins are prepared from *E. coli* cultures grown in D₂O based M9 media. While the D₂O from such cultures is often reused, after distillation, the contaminant proteins and the insoluble pellet are usually discarded. We have set up a very simple method to make use of this deuterated material, and use it in the same way as the commercial lysate to deuterate proteins.

In principle, different parts may be "recycled" after the protein of interest is purified: either the soluble contaminant proteins or the insoluble debris. We have explored both possibilities. The soluble contaminant proteins were retained during the Niaffinity purification step; the insoluble fraction was simply the part that was in the pellet after a centrifugation following cell lysis. In both cases, the sample was treated with phosphoric acid to hydrolyse the peptide bonds. The pH was neutralised by NaOH and the sample was cleared by ultracentrifugation and then lyophilised. We have obtained ca. 375 mg of lyophilised

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5 debris. We have analysed the amino-acid composition of these powders by amino-acid analysis. Figure 2 compares the relative amino-acid composition of these two samples to the composition of the commercial ISOGRO[®] medium. The composition is similar, which suggests that the cell lysate from bacteria can be used in the same manner as the commercial algal lysate.

powder from 1 L of culture using the soluble fraction and ca. 900 and 1400 mg (two independent samples) from the insoluble

To test this assumption, we have produced samples of ubiquitin with either deuterated commercial algal lysate (ISOGRO[®]) or the home-made bacterial lysate described above (2 g/L each), and recorded NMR spectra (Fig. 2b). The spectra show a very

- 160 similar level of deuteration, supporting that bacterial cell lysates is a viable alternative to commercial ones. Small differences are observed, for example in the methyl region. However, these differences do not seem to be fundamental: in fact, when comparing different productions (even from the same batch of lysate), the intensities of aliphatic sites in 1D spectra vary in a similar manner as the variations seen in Fig. 2b. Given that the deuteration levels of e.g. methyls are of only a few percent (see Fig. 1b), even a factor of two in the deuteration level of a given methyl will hardly change the absolute deuteration level.
- 165 The details will depend on the deuteration level of the culture from which the cell lysate has been made. The bacterial cell lysate comes essentially free of additional cost when perdeuterated proteins are made, considering that it uses fractions that are usually discarded. However, with a yield of ca. 1 g per litre of culture, ca. 2 L of a deuterated culture are needed to prepare 1 L of H₂O/M9/cell-lysate-deuterated sample with good deuteration (cf. Figure 1a). Overall, we have shown that bacterial lysate and commercial algal lysate can be used interchangably; this assessment is also confirmed by an application of the home-made bacterial lysate to the MAS NMR study of the pb6 protein assembly (see below).



Figure 1. Deuteration using H₂O-based medium doped with algal lysate. (a) Deuteration level of *I. islandicus* MalDH (33.55 kDa) expressed in H₂O with either ²H,¹³C glucose only (2 g per liter of culture) and ¹⁵NH₄, or with additional use of ²H,¹³C,¹⁵N algal extract (ISOGRO[®]) at three different concentrations (1, 2, 4 g per liter of culture). The reported molecular-weight values are from intact mass spectrometry. The dashed red line indicates the theoretical molecular weight of fully deuterated MalDH (assuming all exchangeable hydrogens are ¹H). (b,c) Residual protonation level for aliphatic sites, determined by solution-state NMR of a sample of ubiquitin produced in H₂O-based M9 medium supplemented with ²H,¹³C,¹⁵N algal extract (2 g/L). Amino acid types missing in this plot are either not present in ubiquitin, excluding the possibility for quantification (Trp) or are not visible or unresolved (Met, His, Arg).

2.3 Coherence life times in H₂O/M9/cell-lysate deuterated samples

Two key parameters we are interested in are the coherence life times (T_2) and the line widths. The line width contains contributions from the coherence life time (the so-called homogeneous contribution), but additional sample inhomogeneity also contributes (Le Marchand et al., 2022). Which of these contributions dominates depends on the sample and the conditions

- 175 (in particular the MAS frequency). To compare the H₂O/M9/cell-lysate samples with perdeuterated and protonated samples, we have, thus, analysed both the homogeneous contribution and the apparent line widths, in two large test proteins, TET2 (39 kDa monomer size, assembling to a dodecamer) and MalDH (33.55 kDa monomer size, assembling to a tetramer). To quantify the homogeneous contribution we have measured the coherence decay using spin-echo experiments. The experiments applied a spin echo ($\tau 180^{\circ}$ pulse τ) element to either ¹H, ¹⁵N or ¹³CO coherence, embedded into a heteronuclear correlation
- 180 experiment with ¹H detection. We compared samples prepared with full deuteration (produced in D_2O medium) with samples deuterated in H_2O medium doped with deuterated algal lysate, and finally a fully protonated sample. As shown above, these three samples correspond to different levels of deuteration. In all cases, the proteins were in H_2O buffer, i.e. all the exchangeable



Figure 2. Comparison of the amino-acid composition in a home-made bacterial extract with the one in ISOGRO[®]. (a) The amino acid composition of ISOGRO[®], as provided by the manufacturer, is shown as grey bars; the values represent the percentage (weight) of each amino acid in the powder. Data of the home-made preparation from bacterial perdeuterated cultures are shown in blue and red. The preparation made from the soluble-proteins fraction, i.e. contaminant proteins from a protein purification, is shown in red. The preparation made from the insoluble fraction after cell lysis is shown in blue. Asn and Gln are indistinguishable from Asp and Glu with this method. Cys and Trp are destroyed during the processing steps of the analysis. (b) 1D ¹H spectra of ubiquitin deuterated in H₂O based medium supplemented with either commercial algal lysate (black) or the home-made lysate, prepared from the contaminant proteins. The precultures have been made either in H₂O or D₂O, as indicated. The similarity of the spectra confirms that commercial algal lysates and bacterial lysates produce similar labelling patterns, as expected from the very similar distribution of amino acids shown in panel (a).



Figure 3. Investigation of coherence life times and line widths in differently labelled samples of two proteins. (a) Fitted R_2 ' decay rate constants for four differently labelled samples of MalDH as indicated. The decay curves for representative experiments, indicated with an asterisk, are shown in Figure A1. All data were recorded at a B_0 field strength corresponding to 700 MHz ¹H Larmor frequency in a 0.7 mm probe. (b) Similar data for the TET2 protein. (c) ¹H line widths of TET2 in deuterated samples produced in D_2O -based M9 medium (blue), produced in H_2O -based M9 medium doped with 2 g/L deuterated algal extract (red) or fully protonated (black). Spectra are shown in Fig. A2.

sites are protonated to either 100% (for the two types of samples grown in H_2O medium) or to a level that depends on the accessibility of the given amide site. Because the importance of deuteration depends on the MAS frequency (Le Marchand et al., 2022), we have performed the measurements at various MAS frequencies up to 100 kHz (Figure 3).

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As expected from the deuteration level, the ¹H coherence life times of the samples deuterated with algal lysate in H_2O (red in Figure 3) are in between those of perdeuterated samples (blue) and the fully protonated sample (black). Of note, even at the highest MAS frequency used in this study, 100 kHz, the ¹H coherence life time of the fully protonated sample is more than three times shorter than the one of the sample deuterated with algal medium. Moreover, the ¹H coherence life time of the fully

- 190 protonated sample spinning at 100 kHz is significantly (more than 20%) shorter than the one of the algal-medium-deuterated sample spinning at 60 kHz. This is an important realisation, because in order to be able to spin 100 kHz, one needs to use smaller rotors and sacrifice about ca. a factor of 5 in sample amount, which translates to a severe sensitivity penalty of 100-kHz MAS experiments. As expected from the deuteration level study above, the perdeuterated sample (placed in H₂O) has longer ¹H coherence life times (lower R₂'). Moreover, in absolute terms, the MAS dependency is less pronounced for the deuterated
- 195 sample than the protonated one.

The decay of heteronuclear (amide ¹⁵N, carbonyl ¹³C') coherences depends much less on the deuteration level (Figure 3). It depends significantly on the ¹H decoupling scheme and decoupling power used, and the optimum decoupling scheme and decoupling power generally depends on the MAS frequency. To generate a consistent data set, we have used low-power ¹H WALTZ-16(Shaka et al., 1983) decoupling (ca. 10 kHz), which is not the best-performing scheme at 40 kHz (data not shown).

200 Therefore, these data are to be taken with a grain of salt, and the main message from them is that ¹⁵N and ¹³C' decay is much less dependent on the labelling pattern than ¹H coherence life times. Of note, the experiments performed here did not use ²H decoupling; the rationale is that most probes are not equipped with a ²H coil.

2.4 Line widths in H₂O/M9/cell-lysate deuterated samples at 55 kHz MAS are similar to perdeuteration and better than protonated samples at 100 kHz

- 205 The line widths observed in spectra contain additional inhomogeneous contributions due to sample heterogeneity, inhomogeneity of the magnetic field and anisotropic bulk magnetic susceptibility (Le Marchand et al., 2022; Linser et al., 2014). The R₂' rate constants discussed above do not comprise these effects. We have measured the line widths in a series of 2D hNH spectra obtained with TET2 produced either with the H₂O/M9/lysate approach or by perdeuteration (Figures 3c and A2). We find that the amide ¹H line widths in samples deuterated in H₂O with deuterated algal lysate are comparable to those of proteins
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line widths of the same protein in protonated form, spinning at 100 kHz MAS frequency.

Taken together, the above analyses showed that proteins deuterated in H_2O with deuterated cell lysate, and 100% backexchanged in the final sample, have slightly worse spectroscopic properties (shorter ¹H coherence life times) than perdeuterated proteins under comparable conditions; the ¹H line widths, however, are not significantly larger.

produced in D₂O. Importantly, the line widths of deuterated proteins at 55 kHz MAS frequency are significantly smaller than





Figure 4. Deuteration in H₂O enhanced by amino-acid mixtures from cell lysates allows detection of non-exchangeable amide hydrogens. Overlay of (a) hNH, (b) hCANH, (c) hCONH TET2 spectra, (d) H-N and (e) CA-N projections of the hCANH MalDH spectra, and (f) hCANH pb6 spectra. (For the latter, the in-house made bacterial cell lysate was used.) Spectra of samples deuterated by the H₂O/M9/lysate approach are showed in red, while spectra from perdeuterated samples are shown in blue. All samples were ¹³C and ¹⁵N labelled. Peaks that appear to only be present in the perdeuterated sample spectra can be explained by (i) different apparent frequencies of aliased peaks (since the spectral widths and carriers are not identical in the overlayed spectra), (ii) chemical shift perturbations due to small sample-temperature differences, and (iii) different signal/noise ratios of individual peaks.

215 2.5 Application to three large proteins: retrieving the signals of non-exchangeable amide sites

To show the potential of this deuteration of proteins with full amide-protonation, we have recorded three-dimensional correlation spectra of three large proteins, TET2, MalDH and the 50 kDa-large bacteriophage T5 tail protein pb6. Figure 4 shows the overlays of the hNH, hCONH and hCANH spectra of TET2, and hCANH spectra of MalDH and pb6. It is apparent in all three cases that numerous peaks which are undetected in the standard perdeuteration protocol (blue) are visible in the H₂O/M9/lysate-deuterated protein (red).

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To gain more insight, we have analysed in detail which amide signals are visible in TET2 with the two different deuteration approaches. TET2 is an ideal real-world application. With 39 kDa monomer size, it is one of the largest proteins for which extensive assignments have been reported (Gauto et al., 2019a). The assignment of TET2 has been achieved with a combination of 3D and 4D ¹³C-detected experiments on protonated samples, and ¹H-detected 3D correlation experiments on perdeuterated TET2. 85% of the backbone and 70% of the side-chain heavy atoms have been assigned in this way. 140 amide hydrogen frequencies have been assigned using the standard perdeuterated samples (see Table A1). TET2 is a very stable protein, originating from a hyperthermophilic archaeon, which suggests that the protein rarely populates partially unfolded conformations that would be required to re-protonate the amides. Unfolding/refolding comes with large sample losses, such that the H₂O/M9/lysate approach is ideal.

Figure 5. Newly assigned amide hydrogens in TET2 deuterated in H_2O shown on its monomeric structure. Amide ¹H frequencies that were previously inaccessible with perdeuterated samples and that were assigned with the $H_2O/M9/lysate$ deuteration are represented as spheres on the protein structure. Blue spheres correspond to atoms that could be assigned from the spectra of the $H_2O/M9/lysate$ -deuterated sample (600 MHz ¹H Larmor frequency, 55 kHz MAS), while cyan spheres are for atoms that only showed signal in the spectra of the fully protonated sample (950 MHz ¹H Larmor frequency, 100 kHz MAS). Parts for which the backbone is shown in white regions correspond to those for which heavy-atom assignments have previously been reported (Gauto et al., 2019a), while black regions indicate parts with missing heavy-atom assignment. In those parts we did not attempt to get new amide assignments, as the *de novo* assignment of the backbone would likely require more 3D data sets. Taken together, this data highlights the possibility of detecting amide hydrogens of water-inaccessible regions in the protein. Note that the protein forms a dodecamer (Fig. 4a), and for better visibility only the monomeric subunit is shown.

- We have recorded 3D assignment experiments (hCANH, hCONH) with a sample deuterated in H₂O with cell lysate. These experiments allowed us to assign an additional 58 amide resonances (see Table A1). Figure 5 plots the location of these newly assigned amide resonances in the structure and shows that the majority of them are located in the β -sheets of TET2 as well as in several α -helices. Interestingly, also a few resonances in more exposed regions had not been assigned before and have now become assigned.
- With a total of ca. 200 assigned amide hydrogens for a protein of 338 non-proline residues, there are still missing assignments left. This is in part explained by the fact that ca. 15% of the backbone heavy-atom resonances have not been assigned by ¹³C-detected experiments, which is ascribed at least for some parts to extensively fast transverse relaxation due to µs dynamics (Gauto et al., 2022). Moreover, the somewhat limited set of 3D experiments recorded here were not sufficient to unambiguously assign all detected spin systems, as the intention of this study was to investigate the labelling scheme rather than to do an extensive resonance-assignment effort.

We investigated whether an alternative approach, namely the use of fully protonated samples spinning at 100 kHz would allow the assignment of many additional resonances: hCANH and hCONH experiments allowed us to assign only three additional amide frequencies, which brings the number of assigned amide ¹H sites to 201 (Table A1). We note that the sensitivity of these experiments, recorded in 0.7 mm rotors, is much lower than the one of experiments in 1.3 mm rotors, such that the use of samples deuterated in H₂O represents a clear sensitivity advantage.

Overall, these data exemplify how the use of proteins deuterated in H_2O provides access to amide sites that are difficult to exchange, e.g. buried in the hydrophobic core.

2.6 Residual Ha protonation enables Ha-based experiments and Ha assignment

Our investigation of the deuteration pattern (Figure 1) and, similarly, previously reported results (Löhr et al., 2003) show that
there is a significant fraction of the α sites which bear a ¹H spin. While this residual protonation reduces the coherence life times of amide ¹H spins, it can also be turned to an advantage, as the ¹Hα frequency becomes available as an additional reporter. We have exploited the residual ¹Hα protonation in the H₂O/M9/lysate-deuterated sample to assign ¹Hα frequencies in TET2. To this end we have recorded a 4D Hα-Cα-N-H correlation experiment with three cross-polarisation steps (HACANH), at 55 kHz MAS frequency. As the Cα, N and H frequencies are known for most residues, the assignment of the Hα frequency the assignment. Figure 6a shows examples from this 4D experiment. This experiment allowed the assignment of 157 Hα resonances (Table A1).

In addition, we have recorded a complementary assignment experiment with a 13 C, 15 N labelled (protonated) sample at 100 kHz MAS frequency, which correlates the H α to the directly bonded 13 C α and the adjacent 15 N (hNCAHA; Figure 6b, grey). These two experiments both report on the H α resonances. The 3D hNCAHA experiment, also based on three crosspolarisation steps, relies on the ability to resolve 13 C α - 15 N correlations in two dimensions, in order to unambiguously connect the H α frequency, while the 4D HACANH spreads the signal across three previously assigned dimensions and, therefore, is better in providing unambiguous assignments. 11 additional H α assignments were obtained from the experiment on the protonated samples. The assigned H α -C α resonances are indicated on the 2D hCH spectrum in Figure 6c, and the location of the H α assignments along the sequence is shown in Figure 6d and e. H α assignments provide additional secondary-structure information and may help to better define the secondary-structure assignments using programs such as TALOS-N (Shen and

Bax, 2013); the comparison of TALOS-N results with and without the Hα assignments (Fig. A4) shows only small differences.

Taken together, while the partial protonation of α sites is in principle an unwanted by-product of the deuteration in H₂O, we have shown here that the H α nuclei can be used as an additional nucleus with good resolution, already at MAS frequency of 50-60 kHz, where fully protonated samples show poorly resolved aliphatic ¹H peaks.

270 3 Methods

3.1 Protein samples

P. horikoshii TET2 production was achieved by over-expressing pET-41c plasmid in E. coli BL21(DE3) competent cells (Novagen). The plasmid is available at AddGene (deposition number 182428). Culture media based on M9 minimal media were used for the different labeling schemes: (i) $u-[^{2}H,^{13}C,^{15}N]$ labelled (perdeuterated) samples were expressed in M9 minimal media, 99.8% D₂O, 15 NH₄Cl as sole nitrogen source, and deuterated and $^{13}C_6$ -labeled glucose; (ii) H₂O/M9/lysate-deuterated 275 samples were expressed in H₂O M9 minimal media, supplemented with u-[²H,¹³C,¹⁵N] labeled algal lysates (2 g/L of culture unless specified otherwise: Sigma-Aldrich) and ²H.¹³C D-glucose as an additional carbon and energy source. (iii) u- $[^{1}H, ^{13}C, ^{15}N]$ labeled (protonated) samples were expressed in H₂O M9 minimal media, integrating $^{15}NH_{4}Cl$ and protonated ¹³C₆ D-glucose. Cells were grown at 37 °C until OD (600 nm) reached ca. 0.6-0.8. Protein expression was induced using isopropyl-β-D-1-thiogalactopyranoside (IPTG; 1 mM). Cells were harvested by centrifugation and pellets were resuspended 280 in Lysis buffer T (Table A2), and disrupted using a Microfluidizer. Cell lysates were heated to 85 °C for 15 minutes and subsequently centrifuged (17500 rcf, 1 hour, 4 °C). The supernatant was dialyzed overnight against Dialysis buffer at room temperature and re-centrifuged as in the previous step. Protein purification was conducted using a Resource O column (GE Healthcare), eluting TET2 using a linear gradient of Elution buffer over 10 column volumes. Fractions were analysed by SDS-285 PAGE (12.5 % polyacrylamide) and TET2 was identified by its monomeric molecular weight (39 kDa) and concentrated using an Amicon 30 kDa concentrator (Millipore). The concentrated solution was loaded onto a HiLoad 16/60 Superdex 200 column (GE Healthcare) equilibrated with Dialysis buffer. Samples for MAS NMR measurements were prepared as described earlier (Gauto et al., 2019a), by concentrating TET2 to 10 mg/mL in NMR buffer. The solution was then mixed (1:1 vol/vol) with 2-methyl-2.4-pentanediol (MPD). Protein precipitates were packed into 0.7/1.3/1.6/3.2 mm MAS rotors by ultrecentrifugation (50000 g, at least 1 hour). 290

I. islandicus malate dehydrogenase was produced by over-expression of pET-21a(+) plasmid in *E. coli* BL21(DE3) competent cells (Novagen). M9 minimal-based media were used as for TET2; H₂O/M9/lysate-deuterated cultures were supplemented with 1, 2 or 4 g/L of culture of u-[²H,¹³C,¹⁵N] labeled ISOGRO[®], as well as the standard 2 g/L of ²H,¹³C D-glucose and 1

g/L ¹⁵NH₄Cl. Cells were grown at 37 °C until OD (600 nm) reached ca. 0.6-0.8. Protein expression was induced using IPTG
(1 mM). Cells were harvested by centrifugation after 3-5 hours of expression, and pellets were resuspended in Lysis buffer
M. Lysis was performed by sonication, using a Q700 Ultrasonic Processor (Qsonica), at 40% amplitude and for a total oper-



Figure 6. Protein deuteration in H₂O allows assignment of H_α hydrogens

Figure 6. (continued) (a) Example strips from 4D HACANH and (b) 3D hNCAHA, hCANH correlation experiments exploited for the assignment of H α hydrogens. (c) Overlay of the H α region of the hCH spectra for the fully protonated (grey) and deuterated in H₂O (red) samples. The same color scheme applies to panels a and b. (d) Newly assigned H α hydrogens are represented as spheres on the protein structure. Green spheres correspond to atoms that could be assigned from the spectra of the sample deuterated in H₂O, while orange spheres are for atoms that only showed signal in the fully protonated sample's spectra. Black regions indicate missing H α assignment. (e) Sequencebased representation of TET2 resonance assignments. Newly assigned amide and H α hydrogens are shown in blue and green, respectively. Lost assignments correspond to sites for which the automated-assignment software FLYA (Schmidt and Güntert, 2012) did not converge after adding the peak list from the H₂O/M9/lysate-deuterated sample to the ones used previously for automatic assignment (Gauto et al., 2019a). Protein deuteration in H₂O allows assignment of Ha hydrogens.

ating time of 6 minutes. Cell lysates were heated to 70 °C for 20 minutes and subsequently centrifuged (40000 rcf, 1 hour, 4 °C). The supernatant was fitered and protein purification was conducted using a Resource O column (GE Healthcare) equilibrated in Buffer A, eluting the protein using a linear gradient of Buffer B over 10 column volumes. Fractions were analysed by SDS-PAGE (12.5 % polyacrylamide) and the protein was identified by its monomeric molecular weight (33.55 kDa) and concentrated using an Amicon 10 kDa concentrator (Millipore). The concentrated solution was therefore loaded onto a HiLoad

26/600 Superdex 200pg column (Sigma-Aldrich) equilibrated with Buffer A. MAS rotors were filled for NMR measurements

by ultracentrifugation (68300 rcf, over night).

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Bacteriophage T5 tail tube protein pb6 modified in C-ter with TEV cleaving site and hexahistidine-tag was produced by 305 overexpression in E. coli BL21(DE3) (SingleTM competent cells Merck) of a pb6-pLIM13 plasmid. Cells were grown either (i) from deuterated pre-culture in D₂O-based minimal buffer (M9) or (ii) with home-made $u-[^{2}H, ^{13}C, ^{15}N]$ cell lysate from a deuterated bacterial culture added at a concentration of 2 g/L to a H₂O/M9 culture medium. ¹³C-labeled glucose (2 g/L) and ¹⁵NH₄ (1 g/L) were used in the medium in both cases. The purification protocol is reported in ref. (Arnaud et al., 2017). Briefly, cells lysis was performed by 6 passages through a Microfluidizer (M-110 P Microfluidics). Cell lysates were centrifuged for 310 30 minutes at 3000 rpm (Type 45 Ti Rotor) and 4°C, and pellets were resuspended and incubated for 1 hour at 37°C in

- Resuspension buffer P. They were then applied to a sucrose cushion gradient (50, 40, 35 and 35% sucrose cushion, Sucrose buffer), and spun for 30 minutes at 30000 rpm (SW41 rotor) and 4°C. The obtained pellet was energetically resuspended in Buffer P and pelleted/resuspended several times by low-speed centrifugation, to remove sucrose contamination. Ubiquitin samples, prepared to estimate the labelling amount, were expressed in *Escherichia coli* BL21(DE3) cells, trans-
- formed with a pET-21b plasmid carrying the human Ubiquitin gene. For the production of perdeuterated ubiquitin, transfor-315 mants were adapted progressively in four stages over 48 h to M9/D₂O media containing 1 g/L ¹⁵ND₄Cl, 2 g/L ²H, ¹³C D-glucose as the sole nitrogen and carbon sources. In the final culture, the bacteria were grown at 37 °C. When the optical density at 600 nm (OD₆₀₀) reached ca. 0.8-0.9, protein expression was induced by addition of IPTG to a final concentration of 1 mM and cells were incubated for another 3 h at 37 °C. For the ubiquitin samples prepared with cell lysate (either ISOGRO[®] or in-house 320 made cell lysate, see below) we used either the same adaptation protocol and a D_2O -based preculture, or a shortened protocol
 - 15

without adaption state and with a H₂O-based preculture. The D₂O preculture does not have an effect on the final labelling (Fig.

2b). In all cases, cells were harvested by centrifugation, resuspended in 20 mL of Lysis buffer U, and lysed by sonication. The lysate was centrifuged for 30 minutes at 46,000 g (JA25-50 Beckman rotor), and the supernatant was dialyzed against two times 300 mL of Buffer U. After dialysis the sample was centrifuged for 30 minutes at 46,000 g and loaded on a 40 mL O-

325 Sepharose column. Ubiquitin was recovered in the flow-through fractions, which were subsequently concentrated and injected on a HiLoad 16/60 Superdex 75 column equilibrated with 1 column volume of Buffer U. The buffer was exchanged to pH 6.5 for solution-state NMR.

3.2 NMR spectroscopy

MAS NMR spectra of the proteins TET2 (except those of Fig. 3) and pb6 were acquired on a Bruker Avance 3 HD (600 MHz) equipped with a 1.3 mm probe tuned to ¹H, ¹³C, ¹⁵N with an auxiliary coil tuned to ²H. The effective sample temperature of 330 MAS NMR experiments was kept at ca. 28 °C, measured from the bulk water frequency, using the relationship T[° C]=255 - 90 $\cdot \delta_{H20}$, where δ_{H20} is the bulk water (Böckmann et al., 2009) frequency in ppm. The chemical shift was referenced with respect to the signal of MPD at 4.1 ppm, which is not significantly dependent on temperature (Fig. S1 of ref. (Gauto et al., 2019b)). Additional spectra of TET2 at 100 kHz MAS frequency (Fig. 6) were recorded on a Bruker Avance 3 HD (950 MHz) spectrometer equipped with a 0.7 mm probe tuned to ¹H, ¹³C, ¹⁵N. Experiments for T₂' measurements (Fig. 3) were recorded 335

on a Bruker NEO spectrometer (700 MHz) equipped with a 0.7 mm HCN probe (Fig. 3). Experiments with MalDH were recorded on Bruker NEO spectrometers operating at 600 MHz or 700 MHz as specified in Fig. 4d, e. using 1.9 mm HXY probes tuned to ¹H, ¹³C, ¹⁵N.

All experiments reported herein (2D hNH, 2D hCH, 3D hCANH, 3D hCONH, 4D HCANH) were recorded with pulse sequences available in the NMRlib library (Vallet et al., 2020). All transfers in these experiments used cross-polarization 340 transfer steps. The 4D HCANH (Fig. 6a) used the following CP parameters: 40 kHz (¹³C) and 92 kHz (¹H, linear ramp 90-100) for the H-CA transfer (4 ms), 12.5 kHz (¹³C) and 42 kHz (¹⁵N, linear ramp 70-100) for the CA-N transfer (8 ms) and 40 kHz (¹⁵N) and ca. 90 kHz (¹H), linear ramp 90-100) for the N-H transfer (1 ms). The indirect dimensions were as follows: CA 34 ppm, 88 points; N 32 ppm, 120 points; HA 12 ppm, 58 points. The recycle delay was set to 0.65 s, and the total duration was

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ca. 6 days. Similar CP parameters were used for hCONH, hCANH and hNH and hCH experiments recorded at 55 kHz. Typical experimental times for the 3D experiments were 2 days.

The hNCAHA experiment performed at 100 kHz (950 MHz) used the following CP parameters. H-N CP at 22 kHz (¹⁵N) and 122 kHz (¹H, ramp 90-100; 1 ms), N-CA 38 kHz (¹⁵N, ramp 90-100, 4 ms) and 70 kHz (¹³C), and CA-HA CP at 30 kHz (¹³C) and 125 kHz (¹H, ramp 90-100, 1 ms); the total experimental time was 2 days. The CP setting used for the hCH experiment was the same as the CA-HA setting above.

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The hCANH experiment performed at 38 kHz for MalDH (600/700 MHz) used the following CP parameters. H-N CP at 24/48 kHz (¹⁵N) and 63/85 kHz (¹H, ramp 90-100; 1 ms) H-CA CP at 23/50 kHz (¹³C) and 63/82 kHz (¹H, ramp 90-100; 4 ms), N-CA 28/30 kHz (¹⁵N, ramp 90-100; 10 ms) and 8/7 kHz (¹³C). The experimental time was 2 days.

Solution-state NMR of ubiquitin was performed on a 700 MHz Bruker Avance 3 HD spectrometer equipped with a cryoprobe to quantify the amount of deuteration. ¹³C-HSQC experiments were recorded on samples of ca. 0.3-0.5 mM concentration 355

with a ¹³C spectral width of 70 ppm (512 complex points) using a 3 s recycle delay. The intensity was normalized by the sample concentration (confirmed by 1D ¹H spectral intensity), and the deuteration level was obtained by comparing the peak intensities of aliphatic signals with those of the fully protonated sample. The results for the H α sites were confirmed using an alternative method based on HNcoCA experiments omitting ¹H decoupling, as done by Löhr *et al.* (Löhr et al., 2003).

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For all NMR spectra, spectral inspection, manual peak-picking and chemical shifts assignment were performed using the CCPNMR software (Vranken et al., 2005). Routines for fits of coherence decays (Fig. A1) were written in Python.

3.3 Mass spectrometry

The deuteration level of MalDH produced with different culture-media compositions (Fig. 1a) was determined by intact mass spectrometry at the MS facility of the Max Perutz Labs. Briefly, samples were diluted to 20 ng/μL in H₂O, and 30 - 40 ng
were loaded on a XBridge Protein BEH C4 Column, 300 Å, 2.5 μm, 2.1 mm x 150 mm (Waters), using a Dionex Ultimate 3000 HPLC system (Thermo Scientific). The proteins were eluted with an acetonitril gradient from 12 to 72 % in 0.1 % formic acid at a flow rate of 250 μL/min. Mass spectra were recorded in the resolution mode on a Synapt G2-Si mass spectrometer equipped with a ZSpray ESI source (Waters). Glu[1]-Fibrinopeptide B (Glu-Fib) was used as a lock mass, and spectra were corrected on the fly. Data were analyzed in MassLynx version 4.1 using the MaxEnt 1 process to reconstruct the uncharged average protein mass.

3.4 Preparation of in-house made cell lysate

We prepared a substitute for the commercial algal medium by using unused parts from cultures prepared under perdeuteration conditions, i.e. from proteins usually considered as contaminants. Standard perdeuteration conditions were used for the cultures $(D_2O, 2 \text{ g/L} ^2H, ^{13}C \text{ glucose}, 1 / L ^{15}N \text{ ammonium chloride})$. We prepared such cell lysates from two independent cultures. The

- 375 proteins produced in these cultures were either TET2 or ClpP (as reported in ref. (Felix et al., 2019)). In both cases, the proteins are expressed in the soluble fraction (rather than in the pellet of the membrane/inclusion bodies). In one experiment, performed with ClpP, the protein of interest was purified by Ni-affinity chromatography. All wash fractions from this affinity chromatography step (prior to eluting ClpP) with significant amount of protein were pooled and dialysed against water to eliminate the imidazol contained in the wash steps. The solution containing the contaminant proteins was then acidified with 1 M phosphoric
- 380 acid (1 M final) and left at 80°C in an incubation oven for 5 days to allow for acid-catalysed peptide-bond hydrolysis. The rationale for using phosphoric acid is that after neutralisation the sample contains phosphate, a buffer component used in M9 media. Thereafter, it was neutralised with NaOH, cleared by centrifugation and lyophilised.

The second approach, performed with ClpP and TET2, used the debris after lysis of the cells and centrifugation; rather than using the soluble fraction that was taken above, the insoluble fraction was subjected to the same acid hydrolysis treatment described above. The amino-acid composition from all three samples was very similar.

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3.5 Amino-acid analysis

For the determination of the amino-acid composition of the in-house made lysate, the lyophilised powder of the samples described in the previous section, doped with a precisely known amount of norleucine, were dried and then hydrolyzed for 24 hours at 110 °C in 6 N HCl containing 1 % (w/v) phenol. Samples were then dried and resuspended in analysis buffer and injected into an ion-exchange column of a Biochrom 30 amino acid analyzer, using measurements of the optical density at 570 and 440 nm. Quantification was performed with respect to the internal norleucine standard.

Conclusions 4

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For the deuteration of proteins by bacterial over-expression, several approaches have been proposed which differ in the nature of the growth-media components added, and the pattern and amount of labelling obtained in the final protein. As a rough 395 guide of what labelling can be expected, it is useful to keep in mind that protons from glucose tend to end up in side chain positions, α protons tend to come from the solvent (H₂O vs D₂O) due to transaminases activity, and amides are protonated to a level the corresponds to the H₂O content. If bacteria do not need to synthesise amino acids from glucose because these are provided in deuterated form, then the side chains tend to be deuterated to a level close to the one of the added amino acids. With these rough guidelines in mind, one can understand that the "iFD" approach (Medeiros-Silva et al., 2016), which

- 400 uses 100% H_2O and deuterated glucose, has a lower deuteration level than the H_2O /lysate approach used here. The "random adjoining protonation" appraoch (Asami et al., 2010, 2012), which uses protonated glucose and ca. 75 to 95 % D₂O, yields a higher protonation level at side chain positions, in particular methyls. The approach we advocate here which combines H_2O_2 deuterated glucose and deuterated amino acids from a lysate, yields highest deuteration level among these approaches (overall ca. 80 % deuteration), and 100 % amide protonation. Compared to the iFD method (Medeiros-Silva et al., 2016) our approach 405 enhances the deuteration ca. 4-fold.

The deuteration level is lower for α sites and varies depending on the amino-acid type, presumably due to the efficiencies of the transaminases acting on different amino acid types. Despite the substantial residual H α protonation, which leads to somewhat shorter amide-¹H coherence life times (Fig. 3), the line widths, which include inhomogeneous contributions, are similar to those of perdeuterated samples. We have demonstrated the utility of the approach for high-resolution ¹H-detected

410 MAS NMR using two large soluble protein complexes and a tube-forming protein, all of which have subunit sizes in the 33-50 kDa range. The approach will likely also become useful for membrane proteins, which have solvent-occluded parts that are often difficult to reprotonate at the amide sites (Medeiros-Silva et al., 2016).

The residual protonation of aliphatics, and in particular of α sites, can be used to obtain assignments of H α frequencies. One may also use these residual protons for proton-proton distance measurements. A possible advantage over fully protonated

415 samples is that the aliphatic protons are more dilute, thereby reducing dipolar-truncation effects that represent a challenge for such measurements (Jain et al., 2017). A drawback of the sparseness of the aliphatic protons, however, is that the probability of observing a contact depends on the product of the populations of the two involved protons. In this respect, a specific labelling to near 100%, such as methyl labelling, is certainly more sensitive (Huber et al., 2011; Linser et al., 2011a).

Given that the protonation level at the Hα sites varies for different amino-acid types (Fig. 1), one may use Hα-detected or
-filtered experiments for amino-acid type editing. We have not explored this possibility at this stage. A possible difficulty that needs to be addressed in this direction is the fact that even with a uniform protonation level, peak intensities often differ vastly because of e.g. differences in transfer efficiency.

Our approach is very similar to (and conceptually the same as) the previously proposed method of deuteration in H₂O which uses a bacterial growth medium (such as SILEX[®] or ISOGRO[®]) without the other M9 medium components, as proposed for solution-state NMR (Löhr et al., 2003) or for MAS NMR (Aucharova et al., 2024), published simultaneously to this manuscript by the Linser group. There are slight variations in the amount of components used in these reports and thus the prices of the samples. It is interesting to obtain an estimate of the sample costs by considering competitive prices we obtained from different providers at the end of 2023 (D₂O ca. 400 €/L, ²H, ¹³C D-glucose ca. 300-400 €/g, ¹³C D-glucose ca. 300 €/g, algal lysate ²H, ¹³C, ¹⁵N ca. 200-350 €/g). These numbers result in a cost of ca. 1500 €/L of culture for a sample deuterated in H₂O with cell lysate as proposed here, and ca. 1200 €/L for a perdeuterated sample. Using more of the complex growth media (SILEX or ISOGRO), as proposed elsewhere, leads to somewhat higher prices (Löhr et al., 2003; Aucharova et al., 2024). The cost of the samples can further be reduced, compared to what has been used in our study: O'Brien *et al.* have obtained deuteration levels approaching 80 % using 2 g of algal cell extract and 1 g of unlabeled glucose per litre of culture (as opposed to 2 g/L deuterated glucose used here). They assumed that this quantity of glucose may be required to maintain some cellular functions

- 435 of *E. coli* (O'Brien et al., 2018). Accordingly, the cost of the algal-lysate/H₂O approach would be further reduced to 400-700 €/L, which is of the order of two-fold cheaper than a perdeuterated sample. Of course, these numbers are to be taken with caution as they vary. An additional advantage of the use of cell lysates, compared to perdeuteration, is that they often also boost protein production significantly, effectively lowering the cost of this approach further, compared to perdeuteration. The details depend very much on the protein and expression conditions.
- 440 Compared to the use of protonated samples used at very high MAS frequencies, the deuteration approach proposed here has significant advantages in terms of NMR parameters. Amide-¹H line widths obtained with protonated samples spun at 100 kHz (0.7 mm rotor) are significantly larger than those of H₂O/M9/lysate-deuterated samples spun at 55-60 kHz (1.3 mm rotor), and similar to those obtained when spun at 40 kHz (1.9 mm rotor; see Fig. 3c). The larger volume of 1.3 mm and 1.9 mm rotors (2.5 µL and 13.1 µL, respectively) represents a significant sensitivity advantage over a 0.7 mm rotor (0.59 µL). Le Marchand *et*
- *al.* estimated a factor of ca. 2.5 lower detection sensitivity for a 0.7 mm probe, due to the smaller rotor (Fig. 9 of (Le Marchand et al., 2022)). An experimental study compared 1.3 and 1.9 mm rotors and reported a sensitivity gain of the 1.9 mm rotor by a factor of ca. 1.2-1.9 (Nieuwkoop et al., 2015). The sensitivity ultimately also depends on the losses during the pulse sequence and the relaxation properties. Given the apparent decay rate constants in H₂O/M9/lysate-deuterated and protonated samples (Fig. 3a, b), the approach proposed here likely provides superior sensitivity also for complex experiments with many transfer
- steps. Lastly, while protonated samples are cheaper than the ones we used here, 0.7 mm rotors are more expensive (ca. 3800 €) than 1.3 mm (ca. 2600 €) or 1.9 mm rotors (2000 €), and in our hands the process of opening and re-using 0.7 mm rotors is more prone to damage, in particular of caps (a cap also costs ca. 250-300 €). Again, these prices are to be taken with a grain of salt, but illustrate that currently the proposed H₂O/M9/lysate approach is also economically advantageous.

Our approach can of course also be used with a H₂O/D₂O mixture, which will result in even higher deuteration levels. For example, using 50 % D₂O in the growth medium will result in 50 % protonation of the non-exchangeable sites and 100 % for the exchangeable sites (provided the final sample is in H₂O). Using this approach enables the detection of all amide sites without lengthy exploration of refolding protocols with unsure outcome. In new projects in our group we apply this approach right from the start whenever the involved protein is likely to have non-exchangeable amides, thereby saving precious time and ensuring high quality samples.



Figure A1. Examples of coherence decays in differently labelled samples. The decay curves correspond to the data marked with an asterisk in Figure 3a. The solid lines are fits to the data, using a mono-exponential decay function. The data have been normalised such that the fitted curve at a relaxation delay of zero is at unity.



Figure A2. Line widths in differently labelled samples. These spectra have been used to extract the ¹H line width statistics reported in Figure 3c. Samples 1 and 2 were deuterated in a D₂O based culture; they have been used for previous studies(Kurauskas et al., 2016; Gauto et al., 2019a), for which additional labelling has been used as indicated. Note that this labelling introduces protons only at the terminal methyl of Ile (sample 1) or a single hydrogen at the terminal methyls of Ile, Leu, Val and the ζ -position of Phe (sample 2), far from the amide sites, and does not lead to significant changes of amide ¹H line widths. Sample 3 has been deuterated in H₂O based medium supplemented with deuterated ISOGRO[®]. Sample 4 is fully protonated; note that the latter has been measured at 950 MHz ¹H Larmor frequency, while all other spectra were measured at 600 MHz. Crosses indicate the peaks that have been used for the analysis, and the plots at the bottom indicate the binned line width distribution.



Figure A3. Ultra-fast MAS of a fully protonated and deuterated TET2 samples indicate the usefulness of deuteration. (a) 2D hNH overlay of perdeuterated (1 H, 13 C, 15 N; final sample in H₂O; blue) and protonated (13 C, 15 N) TET2 samples. Newly assigned amino-protons are labeled. (b) Overlay of 2D hCH spectra of the protonated sample (black) with a sample that is 13 CHD₂ labeled at Ile (δ 1), Leu (δ 1) and Val (γ 1) as well as the *para*-CH site of Phe (Gauto et al., 2019b). The resolution gain with the specifically methyl-labelled sampled is obvious.



Figure A4. TALOS-N analysis of the previously-deposited (BMRB 27211) and newly-determined chemical shift assignments. (a) TALOS-N prediction of the secondary-structure (helix, strand or loop) probabilities per residue. The predictions for the two data sources is almost identical, and both are in agreement with the secondary-structure elements from the crystal structure (printed over the bar graphs). (b) Predicted RCI-S² backbone order parameters highlight an overall decrease of the protein flexibility and improvement of the assignment self-consistency. (c) TALOS-N evaluation of the assignment's quality per residue. The new data show an improvement in Strong (228 \rightarrow 238) and Generous (16 \rightarrow 20) assignments, and a decrease of Warning ones (41 \rightarrow 31). (d) TALOS-N confidence of the 3-state (helix, sheet or coil) secondary-structure prediction per residue.

Table A1. Residues for which the chemical shifts of amino- and α -protons were assigned. Residues that could not be assigned in the sample above are highlighted in bold.

Labeling	Assignment
DCN	$ \begin{array}{c} \underline{\mathrm{H}}: 14, 17, 18, 20, 21, 23, 24, 25, 26, 28, 29, 35, 36, 38, 39, 41, 42, 43, 44, 46, 47, 48, 50, 55, 56, 57, 62, 70, 71, \\ 73, 78, 79, 80, 81, 82, 83, 84, 87, 90, 91, 92, 93, 94, 96, 100, 101, 102, 108, 109, 110, 111, 112, 113, 114, 118, \\ 135, 146, 147, 148, 149, 150, 152, 153, 155, 157, 158, 159, 161, 165, 166, 168, 170, 171, 172, 174, 179, 181, \\ 182, 184, 185, 186, 195, 203, 210, 211, 214, 219, 221, 223, 224, 225, 227, 229, 231, 235, 236, 238, 239, 240, \\ 241, 242, 244, 245, 247, 248, 249, 251, 252, 253, 255, 262, 263, 264, 265, 269, 270, 272, 273, 275, 276, 283, \\ 288, 290, 291, 292, 298, 301, 302, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 321, 326 \end{array}$
ISOGRO®	$ \begin{array}{l} \underline{\mathrm{H}}: 14, 17, 18, 20, 21, 23, 24, 25, 26, 28, 29, 35, 36, 38, 39, 41, 42, 43, 44, 46, 47, 48, 50, 55, 56, 57, 62, 70, 71, 73, 78, 79, 80, 81, 82, 83, 84, 87, 90, 91, 92, 93, 94, 96, 97, 100, 101, 102, 108, 109, 110, 111, 112, 113, 114, 118, 135, 146, 147, 148, 149, 150, 152, 153, 155, 157, 158, 159, 161, 165, 166, 168, 170, 171, 172, 174, 179, 181, 182, 184, 185, 186, 195, 203, 210, 211, 214, 219, 221, 223, 224, 225, 227, 229, 231, 235, 236, 238, 239, 240, 241, 242, 244, 245, 247, 248, 249, 251, 252, 253, 255, 262, 263, 264, 265, 269, 270, 272, 273, 275, 276, 283, 288, 290, 291, 292, 298, 301, 302, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 321, 326, 16, 22, 27, 32, 33, 45, 49, 51, 52, 54, 64, 66, 67, 69, 74, 75, 85, 98, 104, 105, 106, 115, 117, 133, 144, 145, 162, 163, 169, 173, 192, 193, 196, 204, 205, 207, 215, 216, 217, 222, 226, 230, 233, 257, 258, 261, 274, 284, 289, 314, 315, 316, 317, 319, 325, 329, 330, 334 \\ \underline{\mathrm{HA}}: 16, 18, 20, 22, 25, 26, 27, 28, 32, 33, 35, 38, 41, 45, 46, 47, 49, 50, 51, 52, 53, 56, 57, 62, 63, 64, 65, 66, 67, 69, 70, 73, 74, 75, 78, 79, 81, 82, 84, 85, 87, 90, 91, 92, 93, 94, 96, 98, 101, 102, 105, 106, 110, 111, 113, 115, 117, 133, 135, 144, 145, 146, 147, 148, 149, 150, 157, 158, 159, 161, 162, 163, 165, 166, 168, 169, 171, 173, 174, 179, 181, 185, 192, 193, 196, 203, 204, 205, 207, 210, 211, 212, 214, 215, 216, 221, 222, 224, 225, 226, 230, 231, 233, 235, 236, 241, 242, 244, 245, 247, 248, 250, 251, 252, 253, 255, 257, 258, 260, 262, 263, 264, 265, 269, 272, 273, 274, 276, 279, 284, 288, 289, 290, 298, 301, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 314, 315, 316, 317, 319, 321, 325, 326, 320, 330, 331, 334 \\ \hline \begin{array}{l} \underline{\mathrm{A}}: & 16, 18, 20, 21, 22, 23, 23, 23, 235, 236, 241, 242, 244, 245, 247, 248, 250, 251, 252, 253, 255, 257, 258, 260, 262, 263, 264, 265, 269, 272, 273, 274, 276, 279, 284, 288, 289, 290, 298, 301, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 314, 315, 316, 317, 319, 321, 325, 326, 329, 330, 331, 334 \\ \end{array} $
CN	$ \begin{array}{l} \underline{\text{H}:} 14, 17, 18, 20, 21, 23, 24, 25, 26, 28, 29, 35, 36, 38, 39, 41, 42, 43, 44, 46, 47, 48, 50, 55, 56, 57, 62, 70, 71, 73, 78, 79, 80, 81, 82, 83, 84, 87, 90, 91, 92, 93, 94, 96, 100, 101, 102, 108, 109, 110, 111, 112, 113, 114, 118, 135, 146, 147, 148, 149, 150, 152, 153, 155, 157, 158, 159, 161, 165, 166, 168, 170, 171, 172, 174, 179, 181, 182, 184, 185, 186, 192, 195, 203, 210, 211, 214, 219, 221, 223, 224, 225, 227, 229, 231, 235, 236, 238, 239, 240, 241, 242, 244, 245, 247, 248, 249, 251, 252, 253, 255, 262, 263, 264, 265, 269, 270, 272, 273, 275, 276, 283, 288, 290, 291, 292, 298, 301, 302, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 321, 326, 16, 22, 27, 32, 33, 45, 49, 51, 52, 54, 64, 66, 67, 69, 74, 75, 85, 97, 98, 104, 105, 106, 115, 117, 133, 144, 145, 162, 163, 169, 173, 192, 193, 196, 204, 205, 207, 215, 216, 217, 222, 226, 230, 233, 257, 258, 261, 274, 284, 289, 314, 315, 316, 317, 319, 325, 329, 330, 334, 72, 208, 218 \underline{\text{HA}:} 16, 18, 20, 22, 25, 26, 27, 28, 32, 33, 35, 38, 41, 45, 46, 47, 49, 50, 51, 52, 53, 56, 57, 62, 63, 64, 65, 66, 67, 69, 70, 73, 74, 75, 78, 79, 81, 82, 84, 85, 87, 90, 91, 92, 93, 94, 96, 98, 101, 102, 105, 106, 110, 111, 113, 115, 117, 133, 135, 144, 145, 146, 147, 148, 149, 150, 157, 158, 159, 161, 162, 163, 165, 166, 168, 169, 171, 173, 174, 179, 181, 185, 192, 193, 196, 203, 204, 205, 207, 210, 211, 212, 214, 215, 216, 221, 222, 224, 225, 226, 230, 231, 233, 235, 236, 241, 242, 244, 245, 247, 248, 250, 251, 252, 253, 255, 257, 258, 260, 262, 263, 264, 265, 269, 272, 273, 274, 276, 279, 284, 288, 289, 290, 298, 301, 303, 304, 305, 306, 307, 308, 309, 310, 312, 313, 314, 315, 316, 317, 319, 321, 325, 326, 329, 330, 331, 334, 72, 208, 218, 43, 54, 97, 104, 170, 217, 249, 261$

Table A	2.1	Buffers	used	during	sample	preparations
				<u> </u>		1 1

Protein, Buffer	Composition		
TET2, Lysis buffer T	50 mM Tris, 150 mM NaCl, 20 mM MgSO ₄ , 0.1 % Triton X-100,		
	0.025 mg/ml lysozyme, 0.05 mg/ml deoxyribonuclease, 0.2 mg/ml ribonuclease (pH 8)		
TET2, Dialysis buffer	20 mM Tris, 100 mM NaCl (pH 7.5)		
TET2, Elution buffer	20 mM Tris, 1 M NaCl (pH 8)		
TET2, NMR buffer	20 mM Tris, 20 mM NaCl (100 % H ₂ O, pH 7.6)		
MalDH, Lysis buffer M	$50\ mM$ Tris, $50\ mM$ NaCl, $2\ mM$ MgCl_2, $0.05\ mg/ml$ deoxyribonuclease, $0.25\ mg/ml$		
	ribonuclease, cOmplete EDTA-free protease inhibitor (pH 7)		
MalDH, Buffer A	50 mM Tris, 50 mM NaCl (pH 7)		
MalDH, Buffer B	20 mM Tris, 1 M NaCl (pH 7.5)		
Ubiquitin, Buffer U	50 mM Tris (pH 8)		
Ubiquitin, Lysis buffer U	50 mM Tris, 2 µg/mL leupeptine, 2 µg/mL pepstatine (pH 8)		
pb6, Resusp. buffer P	50 mM Tris, 100 mM NaCl, 0,5% Triton X-100, 1 mM EDTA, 100 µg/ml lysozyme		
	(pH 8)		
pb6, Sucrose buffer	20 mM Tris, 100 mM NaCl, 0,05% Triton X-100 (pH 8)		
pb6, Buffer P	20 mM Tris, 100 mM NaCl (pH 6.9)		

Code and data availability. All pulse sequences are available in the NMRlib library and from the authors. Analysis scripts for coherencedecay experiments and line widths analysis are available from the authors. The new resonance assignments are submitted to the BioMagRes-Bank (https://bmrb.io/; accession number pending). Other data are available from the authors upon request.

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Author contributions. F. N. prepared all MalDH samples, and TET samples for T₂' measurements, recorded and analysed all T₂' and line width data, recorded all multidimensional experiments of MalDH, performed TET2 resonance assignment and TALOS analysis. J.-Y. G. prepared ubiquitin samples and analysed the labelling pattern, and a TET2 sample. C.-A. A. and C. B. prepared pb6 samples. P. M. and H. F. prepared cell lysates from bacterial cultures. P. S. designed the study and recorded NMR experiments. F. N. and P. S. prepared all figures and wrote the manuscript.

Competing interests. At least one of the (co-)authors is a member of the editorial board of Magnetic Resonance.

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