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Enhancing the resolution of multi-dimensional heteronuclear NMR spectra of intrinsically disordered proteins by homonuclear broadband decoupling†‡

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Limited spectral resolution in the proton dimension of NMR spectra is a severe problem for intrinsically disordered proteins. Here we show that homonuclear broadband proton decoupling of the direct and indirect dimensions of multi-dimensional NMR spectra significantly enhances their resolution.

One-dimensional NMR spectra are rarely sufficient for the structure analysis of small molecules, since they can provide only limited information about through-bond and through-space connectivities and their resolution is not adequate for natural products and even less for biomacromolecules, like proteins. In contrast two- and multidimensional heteronuclear-correlated spectra contain not only important information about connectivities but also show increased signal dispersion. The large majority of two- and multidimensional NMR spectra contain one or more proton dimensions, which typically show the lowest resolution. This results from a combination of their relatively small spectral range (~10 ppm) together with the extensive proton-proton scalar couplings. Limited spectral resolution is a particularly severe problem for intrinsically disordered proteins (IDPs) 1, whose chemical shifts are often close to random coil values. The low resolution encountered in NMR spectra of intrinsically disordered proteins can be partially overcome by using carbon-detected experiments 2 or by employing higher-dimensionality (> 3D) NMR spectra, typically in combination with fast acquisition methods like non-uniform sampling 3, GFT 4, projection reconstruction 5 or APSY 6. However, these experiments are often time-consuming and/or require more elaborate data processing schemes. Broadband homonuclear decoupling has been suggested to be a solution for the resolution problem of proton NMR spectra for small molecules 7. Broadband homonuclear decoupling can be achieved by several approaches: constant-time experiments (applicable for defined, typically heteronuclear coupling constants) 8,9, time reversal experiments 10, the diagonal signals of z-COSY spectra 11, projections of J-resolved 2D spectra 12, BIRD-based decoupling 13,14 and frequency-selective decoupling during a weak gradient field (also called Zangger-Sterk or ZS method) 7,15,16. All of these methods enable decoupling of the indirect dimension of multidimensional spectra. If 1D spectra or decoupling in the acquisition dimension were desired the actual decoupled spectra had to be constructed by different post-processing algorithms. This additional “decoupling dimension” significantly reduces the achievable signal/noise ratio per unit time. Very recently it has been shown that BIRD-based decoupling of protons bound to carbon at natural abundance as well as slice-selective decoupling during a weak gradient field can also be obtained during acquisition in a single scan 17,18. All other methods for decoupling are not amenable for decoupling during acquisition as they do not physically decouple the protons but only extract the decoupling information from the special appearance of the respective spectra or after combining spectra obtained with different pulse phases. Decoupling during acquisition allows the spectra to be processed like regular NMR data and it significantly increases the sensitivity per unit time. So far pure shift spectra were only reported for one- and two-dimensional experiments on small molecules. The use of conventional homonuclear decoupled experiments on biomacromolecules was typically prevented by low sensitivity. The enormous sensitivity gain per time achieved by homonuclear decoupling during acquisition finally lifts this shortcoming. For protons in uniformly 13C-labeled molecules, the frequency-selective decoupling during a weak gradient field is the only option to achieve broadband homonuclear proton decoupling during the acquisition. For this method a selective pulse is applied during a weak pulsed field gradient, which results in the excitation of different signals in different slices of the NMR sample 16,19,20. Homonuclear broadband decoupling is then achieved by a combination of hard and selective 180° pulses, which again are applied during the same weak gradient. In each slice, the signal that was originally excited by the selective pulse is then decoupled from all other protons (for more detailed explanations see 2,16,18,21). Here we show that by using this type of broadband decoupling both in the direct and indirect dimensions of multidimensional NMR spectra, the resolution in the proton-dimensions of intrinsically disordered proteins (IDPs) can be significantly enhanced. The number of known IDPs is rapidly growing and it is currently believed that 45-50% of all eukaryotic proteins contain long ID regions 22,23. They are involved in various physiological functions, like signalling, cell-cycle control or molecular recognition, but they are also implicated in various
Assignment of side chain resonances of well-structured proteins can be readily achieved by triple resonance experiments. The HCCCONH experiment is widely used to correlate side chain carbon and proton or carbon dimension, were used. With this approach we investigated the interaction of the two intrinsically disordered proteins α-synuclein and SERF1a. The pathogenic amyloidogenesis of the Parkinson-associated protein α-synuclein is accelerated by SERF1a through transient interaction with the positively charged C-terminal region of α-synuclein.

Fig. 1. Pulse scheme of the HD $^1$H, $^{13}$C HSQC experiment. The pulse sequence is based on the sequence for a constant time (CT) $^1$H, $^{13}$C HSQC, which has been described earlier, and has been modified to accomplish homonuclear decoupling in the acquisition dimension as follows: The first 90° pulse has been replaced by a slice selective 90° pulse. Gradients 2, 3, and 4 are used to explicitly select magnetization which has been slice-selectively excited and subsequently transferred to $^{13}$C. Acquisition is interrupted every 10-20 ms for decoupling, which is achieved by a combination of a hard and a slice selective 180° pulse as described elsewhere. $^{13}$C decoupling during slice-selective excitation and decoupling is achieved with a GARP decoupling scheme. Narrow and wide rectangles represent 90° and 180° pulses. Filled and open half-ellipsoids denote shaped 90° and 180° pulses respectively. SL represents a spin-lock pulse to suppress residual HDO resonance.

Assignment of side chain resonances of well-structured proteins can be readily achieved by triple resonance experiments. The HCCCONH experiment is widely used to correlate side chain $^{13}$C and $^1$H resonance frequencies with the amide $^1$N and $^1$H resonances. Two 3D experiments, namely a $^1$H($^{13}$C($^{15}$N)) and a ($^1$H($^{13}$C)($^{15}$N)) experiment are frequently used for resonance assignment. However, resolution in the indirect dimension of these experiments is limited to >20 Hz for $^1$H and >60 Hz for $^{13}$C due to homonuclear couplings. Since resonance frequencies of many methyl groups only differ by a few Hz, reliable, unambiguous assignment cannot be obtained with the conventional 3D pulse sequences. In order to obtain high resolution methyl assignments we recorded broadband homonuclear decoupled versions of the experiments. Homonuclear decoupling in the ($^{1}$H($^{13}$C)($^{15}$N)) experiment was achieved by placing a ZS-decoupling scheme in the middle of the $^1$H evolution period. (ESI Fig. 1) To allow for $^{13}$C homonuclear decoupling in the ($^{1}$H($^{13}$C)($^{15}$N)) a constant time evolution period was used instead of the commonly implemented real-time evolution. For those amino acids which have more than one methyl group (Val, Leu, Ile) the $^1$H, $^{13}$C HSQC was used to identify the correct $^1$H, $^{13}$C resonance pairs. In the very few cases where the HD $^1$H, $^{13}$C HSQC spectra do not resolve either of the two methyl groups, a 3D ($^1$H($^{13}$C))H acquisition decoupled in both the indirect carbon and proton dimensions can be utilized. With this approach >95% of the methyl groups of α-synuclein could be assigned unambiguously. Since the linewidths of the decoupled dimension is less than 4 Hz, assignments can be easily transferred to a HD $^1$H, $^{13}$C HSQC spectrum. Thus, the use of high definition spectra for resonance assignment renders the exhaustive acquisition and analysis of 4D and higher dimensional spectra unnecessary. In cases where the resolution in such decoupled 3D spectra is not sufficient, broadband homonuclear decoupling can of course also be combined with fast NMR methods like non-uniform sampling (NUS) or reduced dimensionality experiments. To ensure optimal resolution in all indirect dimensions we used a NUS scheme with 5-20% of the regular sampling. Spectra were processed using MDDNMR.

Despite the high resolution in the indirect dimension of a CT $^1$H, $^{13}$C HSQC most of the resonances of α-synuclein are overlapped and hardly any peak can be assigned unambiguously (Fig. 2). The HD $^1$H, $^{13}$C HSQC, which is decoupled in both frequency dimensions, displays much less signal overlaps. Slice selective decoupling is associated with a loss of sensitivity determined by $\Delta \omega/\gamma G^*s$, with the excitation bandwidth $\Delta \omega$, the gyromagnetic ratio $\gamma$, the gradient strength $G$ and the length of the active sample volume $s$. Note, that this equation is only valid for $\Delta \omega \approx 2G^*s$. The loss of sensitivity is thus proportional to the spectral window which is covered in the decoupled dimension and is partially compensated by the collapse of multiplets into singlets due to decoupling. To restrict the sensitivity loss we recorded two spectra covering the Ha and the H-methyl region of the HSQC spectrum. To achieve this, the carrier frequency was set to 1 or 4 ppm and selective pulses with 80 Hz bandwidths in combination with a gradient of 0.2 or 0.4 Gauss/cm were used for excitation and decoupling of H-methyl or Ha respectively. The actual loss in the sensitivity of the HD HSQC is therefore on the order of 80%. Fig. 2 shows the Val methyl region of α-synuclein, which is rather crowded in the CT $^1$H, $^{13}$C HSQC with only very few peaks being resolved. In contrast the HD $^1$H, $^{13}$C HSQC eliminates most of the overlap: out of 19 Val present in α-synuclein more than 70% show at least one of the methyl groups being resolved. In addition, overlap was greatly reduced for Ala-methyl groups as well so that 9 out of 19 Ala methyl groups are baseline separated in the high definition HSQC. This high level of peak separation, which can very well compete with that of a $^{15}$N-HSQC, enables the HD $^1$H, $^{13}$C HSQC to serve as the basis for NMR titration experiments. In contrast to the $^1$H, $^{13}$C HSQC which is very well suited to monitor interactions involving the protein backbone, $^1$H, $^{13}$C HSQCs can detect in particular hydrophobic interactions which often involve methyl groups. Upon addition of SERF1a, methyl-groups of residues located at the C-terminus of α-synuclein show the most prominent chemical shift perturbations. (CSP, Fig. 3, ESI Fig. 4).
The binding of SERF1a to the C-terminus of \(\alpha\) synuclein enhances its signal dispersion inherent to the direct and/or indirect dimensions greatly simplifies assignment of side chain dimensions. It is perfectly suited for field decoupling by frequency-selective pulses during a weak gradient.

In conclusion we have shown that broadband homonuclear proton decoupling by frequency-selective pulses during a weak gradient can be applied both in the direct and indirect dimensions of two- and multidimensional NMR spectra. It is perfectly suited for intrinsically disordered proteins which suffer from low resolution. Using 2D \(\text{\textsuperscript{1}H}, \text{\textsuperscript{13}C}\) HSQC as well as 3D HCHCOSY spectra, homonuclear broadband proton decoupled in the direct and/or indirect dimensions greatly simplifies assignment of side chain resonances. Furthermore, methyl chemical shift perturbations traced by HD \(\text{\textsuperscript{1}H}, \text{\textsuperscript{13}C}\) HSQC can give valuable information which might not be picked up by conventional NMR titration based on \(\text{\textsuperscript{1}H}, \text{\textsuperscript{13}C}\) HSQCs.

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Notes and references

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