This is the accepted manuscript of the article:

Barbieri, L., Luchinat, E., Backbone resonance assignment of human DJ-1 in the reduced state and in the cysteine sulfinic acid state. Biomol NMR Assign 13, 371–376 (2019).

This version of the article has been accepted for publication, after peer review, and is subject to Springer Nature's AM terms of use, but is not the Version of Record and does not reflect postacceptance improvements, or any corrections.

The Version of Record is available online at: *https://doi.org/10.1007/s12104-019-09908-8*

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4	¹³ C APSY-NMR for sequential assignment of intrinsically
5	disordered proteins
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1 Abstract

The increasingly recognized biological relevance of intrinsically disordered proteins requires a continuous expansion of the tools for their characterization via NMR spectroscopy, the only technique so far able to provide atomic-resolution information on these highly mobile macromolecules. Here we present the implementation of projection spectroscopy in ¹³Cdirect detected NMR experiments to achieve the sequence specific assignment of IDPs. The approach was used to obtain the complete backbone assignment at high temperature of αsynuclein, a paradigmatic intrinsically disordered protein.

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12 Keywords

13 Intrinsically disordered proteins, IDPs, assignment, NUS, ¹³C detection

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1 Introduction

The functional importance of intrinsically disordered proteins (IDPs) or protein regions 2 (IDPRs) has now been widely recognized (Habchi et al., 2014; Wright & Dyson, 2015; van 3 4 der Lee et al., 2014). Furthermore, a number of key proteins related to the onset of diseases is intrinsically disordered or has large disordered regions (Uversky et al., 2008). Perhaps the 5 most well-known examples include proteins involved in the development 6 of neurodegenerative diseases, such as α -synuclein and Tau for Parkinson's and Alzheimer's 7 diseases. In addition, a vast range of examples of IDPs/IDPRs related to cancer progression 8 9 are emerging, beyond the most well-known example of p53. These include c-myc, p21, AR, BRCA to name just a few recent examples (Uversky et al., 2014). Therefore, novel 10 11 approaches to target IDPs or IDPRs are urgently needed in the field of drug discovery (Tóth et al., 2014; Ambadipudi & Zweckstetter, 2016; Joshi et al., 2016; Heller et al., 2017) and in 12 13 this frame it becomes very important to develop NMR methods to study IDPs/IDPRs in their 14 native state under physiological conditions (Felli and Pierattelli eds., 2015).

The peculiar amino acidic composition of IDPs, often characterized by multiple sequence repetitions and low complexity regions, the high flexibility typical of IDPs/IDPRs and the many conformations sampled in which the protein backbone is largely solvent exposed have an impact on NMR observables that should be considered in the design of effective experimental NMR approaches (Brutscher et al., 2015).

The critical points that need to be faced in order to optimize NMR spectra for the study of 20 highly flexible IDPs/IDPRs include the very low chemical shift dispersion typical of proteins 21 that lack a stable, well defined 3D structure, as well as solvent exchange processes that 22 broaden amide proton resonances when approaching physiological conditions. As known 23 24 since the early studies on protein folding, heteronuclear spins are characterized by a large chemical shift dispersion and thus well suited to characterize disordered proteins (Dyson & 25 Wright, 2001). Indeed, initial studies of unfolded proteins were stimulated by the 26 development of 3D triple resonance experiments (Kay et al., 1990; Sattler et al., 1999). 27 However, to study unfolded/intrinsically disordered proteins of increasing complexity, a 28 further push in resolution was necessary; a major contribution towards this goal derived from 29 the extension of the dimensionality of NMR experiments to more than three dimensions. Of 30 course methods based on uniform sampling of indirect dimensions and on conventional 31 processing strategies were not suitable to achieve a high resolution in all dimensions, a key 32 aspect to study complex IDPs/IDPRs; different approaches needed to be developed to 33

render these experiments feasible in a reasonable amount of time and manageable in terms 1 of processing time and disk space. Many non-uniform sampling approaches and different 2 strategies to process the data were indeed proposed (Kupce & Freeman, 2004; Kim & 3 Szyperski, 2003; Hiller et al., 2005; Hoch & Stern, 2001; Orekhov et al., 2003; Kazimierczuk 4 et al., 2006) and initially implemented in triple resonance multidimensional experiments 5 based on H^N detection (Narayanan et al., 2010; Fiorito et al., 2006; Zawadzka-Kazimierczuk 6 et al., 2010; Motackova et al., 2010) and on H^{α} detection (Yao et al., 2014, Mäntylahti et al., 7 2011) showing a great potential for the characterization of IDPs (Fiorito et al., 2006; 8 9 Zawadzka-Kazimierczuk et al., 2012; Narayanan et al., 2010; Yao et al., 2014).

10 In parallel, the increase of the sensitivity of NMR instrumentation (Kovacs et al., 2005) enabled the development of experimental strategies based on heteronuclear direct detection 11 (Bermel et al., 2006c), which turned out to be an excellent tool to study IDPs. Indeed the 12 valuable chemical shift dispersion of heteronuclear spins can be exploited in all dimensions 13 of NMR experiments, including the directly detected one, and limitations deriving from 14 solvent exchange broadening are obviously reduced. Several methods to overcome the 15 problem of ¹³C homonuclear couplings in the direct acquisition dimension were proposed 16 (Felli & Pierattelli, 2015) and enabled the application of C' direct detection with high 17 resolution. A suite of 3D multidimensional C' detected experiments was proposed (Bermel 18 et al., 2009a; Bermel et al., 2009b; Felli et al., 2009; Bermel et al., 2008; Bermel et al., 19 2006b, O'Hare et al., 2009) and proved to be very useful for sequence specific assignment 20 and characterization of several IDPs. The suite was then expanded to include also C' 21 detected experiments of dimensionality higher than 3 by implementing random non-uniform 22 23 sampling in combination with sparse multidimensional Fourier transform (SMFT) processing of the data showing that experiments of higher dimensionality based on C' detection are 24 25 indeed possible and constitute a valuable tool to study IDPs (Novácek et al., 2011; Bermel et al., 2012; Novacek et al., 2012; Haba et al., 2013; Bermel et al., 2013; Piai et al., 2014; 26 Baias et al., 2017). 27

As a further step facilitating the exploitation of the ¹³C-detection based assignment strategy,
 we present the implementation of projection spectroscopy for C´ detected multidimensional
 NMR experiments. The approach is tested on α-synuclein at high temperature.

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1 Material and Methods

Isotopically enriched α-synuclein was prepared as previously described (Huang, Ren et al.,
2005) A sample of 1.0 mM uniformly ¹³C, ¹⁵N labeled human α-synuclein in 200 mM NaCl,
0.5 mM EDTA, 20 mM phosphate buffer at pH 6.5 was used. 10 % D₂O was added for the
lock.

6 All the 2D NMR experiments were acquired at 285.5 K, 295 K, 305 K and 315 K while the 7 APSY experiments were all acquired at 315 K on a Bruker AVANCE NEO spectrometer 8 operating at 700.06 MHz ¹H, 176.05 MHz ¹³C and 70.97 MHz ¹⁵N frequencies equipped with a cryogenically cooled probehead optimized for ¹³C-direct detection (TXO). Carrier 9 frequencies and RF pulses suitable for triple resonance ¹³C detected experiments were used 10 and are summarized hereafter while specific parameters for the different experiments are 11 reported in Table 1. Q5 and Q3 shapes (Emsley & Bodenhausen, 1992) of durations of 300 12 and 231 µs, respectively, were used for ¹³C band-selective $\pi/2$ and π flip angle pulses, 13 except for the π pulses that should be band-selective on the C^{α} region (Q3, 1200 µs) and 14 for the adiabatic π pulse to invert both C' and C^{α} (smoothed Chirp 500 µs, 20 % smoothing, 15 80 kHz sweep width, 11.3 kHz RF field strength) (Böhlen & Bodenhausen, 1993). The ¹³C 16 band selective pulses on C^{α} and C' were applied at the center of each region, 53 and 173.5 17 ppm respectively; the Chirp pulse was centered at 90 ppm. Carrier frequencies for ¹⁵N and 18 ¹H were 122.5 and 4.7 ppm respectively. Decoupling of ¹H and ¹⁵N was achieved with Waltz 19 20 (2.6 kHz) and Garp-4 (1.0 kHz) decoupling sequences respectively (Shaka et al., 1985). All gradients employed had a smoothed square shape. The parameters used for the acquisition 21 22 of the 5D and 4D experiments as well as the parameters selected to implement the APSY approach are reported in Tables 1 and 2 respectively. FLOPSY-16 (Kadkhodaie et al, 1991) 23 24 was used in the 4D experiment to spin-lock. All the spectra were acquired using Bruker TopSpin 4.0.1 software. Calibration of the spectra was achieved using DSS as a standard 25 for ¹H and ¹³C; ¹⁵N shifts were calibrated indirectly (Markley et al., 1998). 26

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1 Results and discussion

The 2D ¹H-¹⁵N HSQC and 2D ¹³C-¹⁵N CON experiments of α -synuclein recorded at various temperatures are reported in Figure 1. It can be noted that the number of cross peaks that can be observed through 2D ¹H-¹⁵N HSQC spectra decreases with increasing temperature. This is the result of the pronounced exchange processes of amide protons with water protons due to the largely solvent exposed backbone of α -synuclein.

7 Nuclear spins of non-exchangeable atoms should thus be exploited in NMR experiments. 8 However, even if in principle very sensitive, ¹H nuclear spins of aliphatic and aromatic residues are characterized by a guite narrow chemical shift dispersion, in particular when 9 considering amino acids of the same (or similar) type in proteins lacking a 3D structure such 10 11 as in IDPs. In addition the ¹H-¹H homonuclear couplings, which are not easy to remove in the direct acquisition dimension, constitute an additional drawback when really high 12 resolution is needed. Instead, non-exchangeable heteronuclear spins, such as ¹³C and ¹⁵N, 13 provide a large chemical shift dispersion even if at the expense of reduced sensitivity when 14 directly detected. Several strategies can be implemented for C' homonuclear decoupling 15 (Felli and Pierattelli, 2015). Thus, correlations between heteronuclear spins involved in the 16 peptide bond are the most appropriate ones to achieve high resolution (Bermel et al., 2013), 17 as evidenced in the spectra reported in Figure 1 which could be acquired with just a few 18 scans per increment; 2D ¹³C-¹⁵N CON spectra (Bermel et al., 2006a) thus provide a valuable 19 tool to study IDPs at physiological conditions (Gil et al., 2013). 20

The 2D ¹³C-¹⁵N CON spectra can then be expanded into higher dimensional experiments 21 that provide different types of correlations between different nuclear spins in order to achieve 22 sufficient information for sequence specific assignment. Increasing the number of 23 dimensions however, even when exploiting heteronuclear spins to take advantage of their 24 high dispersion in IDPs, is effective only if we are able to achieve a very high resolution in 25 the indirectly sampled dimensions. This is practically not feasible due to time constraints as 26 well as disk usage unless non-uniform sampling strategies are used, which, combined with 27 appropriate processing strategies, have shown to be very effective (Bermel et al., 2013; 28 29 Bermel et al., 2012; Novacek et al., 2012; Novacek et al., 2011). Here we test the applicability of projection spectroscopy and of the relative automated data analysis (Hiller et 30 al., 2005) to the C⁻ detected 5D (HCA)CONCACON experiment (Bermel et al., 2013), which 31 correlates the CON peak of one peptide bond with the preceding one, for the assignment of 32 α -synuclein at high temperature (315 K). 33

The four orthogonal planes that can be collected through this experiment, reported in Figure 1 2, show the excellent resolution that can be achieved in all dimensions thanks to the 2 contribution of heteronuclear chemical shifts, in particular of C' and N that provide 3 correlations across one peptide bond (A) and of C' and N belonging to the same amino acid 4 (B), as well as sequential C'-C' correlations (C). The plane correlating C' and C^{α} (D) is 5 characterized by a somehow reduced resolution but still provides useful information about 6 the amino acid type. These are excellent requisites to implement projection spectroscopy 7 through which not only orthogonal planes but also "transverse" planes cutting the 8 9 multidimensional object at different angles are acquired with very high resolution as shown in panels E and F of Figure 2. Thanks to the excellent resolution that can be obtained in the 10 projections of C' detected exclusively heteronuclear experiments, it is possible to resolve 11 the majority of cross peaks in the projections and to identify essentially all correlations 12 expected in the 5D spectrum just by inspection of a limited set of projections. In the present 13 case, 64 projections were acquired with very high resolution (256 points were set in each of 14 the 4 indirect dimensions); it is worth noting that to achieve this kind of resolution for the 15 analogous 5D experiment would have required a prohibitively long time (hundreds of years!). 16 The analysis of the 64 projections, which can be performed automatically by using the 17 algorithm GAPRO (Geometric Analysis of PROjections) (Hiller, Wider et al., 2008; Hiller & 18 Wider, 2012) as it is implemented in the TopSpin 4.0.1 software, provides 191 5D 19 correlations (131 C'_{i-1}, N_i, C^{α}_i, C'_i, N_{i+1} and 60 C'_{i-1}, N_i, C^{α}_{i-1}, C'_{i-1}, N_i). These correlations 20 constitute an excellent tool to achieve sequence specific assignment by linking the shifts of 21 C' and N involved in a peptide bond with those of the previous one, and also giving some 22 hint on the kind of amino acid through the information of the C^{α} chemical shift. The 23 correlations identified through the GAPRO analysis of the projections are stored in the form 24 of N-dimensional peak-lists that can be manually inspected by loading them on the different 25 projections. In particular, by loading the peak-lists on a 2D ¹³C-¹⁵N CON spectrum the 26 27 completeness of the assignment can be verified and it is possible to perform the sequence specific walk through the backbone just by moving from one cross peak (C'_i, N_{i+1}) to the 28 neighboring one (C'_{i-1}, N_i), as shown schematically in Figure 3 for a few residues. It is worth 29 noting that for most of these residues the H^N resonance was not observable in ¹H detected 30 experiments. 31

A few comments are due on technical details related to the implementation of the APSY approach for the 5D (HCA)CONCACON experiment (Bermel et al., 2013). In contrast to the original sequence, the t₂ period was implemented in a semi-constant time manner in order

to increase to possible resolution in this dimension. As noted above, the version presently 1 used yields two types of correlations: type I (C'_{i-1}, N_i, C^{α}_i, C'_i, N_{i+1}) which contains the 2 information for sequential assignment and type II (C'_{i-1}, N_i, C^a_{i-1}, C'_{i-1}, N_i) which correlate 3 three nuclear spins ($C^{\alpha_{i-1}}$, C'_{i-1} , N_i) giving essentially the same information that can be found 4 in 3D CACON experiments. The experiment is optimized to detect correlations of type I; 5 correlations of type II can in principle be suppressed as suggested for the 3D 6 (HCA)CON(CA)H (Mäntylahti et al. 2011). However, in the present case this subset of peaks 7 could be easily identified directly from the peak list and further used for confirming the 8 9 assignment, so that no further modification of the experiment was required. A few type I correlations were instead missing. Manual inspection of the peak list revealed that most of 10 them involve residues in the known α-synuclein repeats which could not all be safely 11 identified through the automatic algorithm. 12

A second comment concerns the number of projections that need to be acquired to obtain 13 the information for sequence specific assignment. In the present case we collected 14 projections at 30 different combinations of angles α , β , γ that, considering the issue of 15 quadrature detection in all indirect dimensions, translates into 64 different 2D projections 16 (Table 2). However, processing of the data using a smaller number of projection angles 17 shows that the information content is essentially the same if we use only 26 combinations 18 of angles, as suggested by the software (Hiller & Wider, 2012; Hiller, Wider et al., 2008), 19 and that the number of type I correlations detected in the spectra is only slightly reduced if 20 we consider a smaller subset of angles (Table 3), showing the robustness of this approach. 21 The major issue to be solved was to properly implement in the APSY software the virtual 22 homonuclear decoupling of C' through the IPAP approach. The processing of the data, 23 including the virtual decoupling, can now be handled very easily through TopSpin 4.0.1. 24

The APSY approach can of course be implemented in any kind of C⁻ detected experiment, 25 using the 2D ¹³C-¹⁵N CON spectrum as a reference. For example, the information about the 26 amino acid side chain, can be easily obtained and correlated to each of the cross peaks 27 observed in the 2D ¹³C-¹⁵N CON spectrum, by collecting experiments such as 4D 28 29 HCBCACON (Bermel et al., 2009) or 4D HCCCON (Bermel et al., 2012). As an example the latter was acquired on α-synuclein and provided information about the vast majority of ¹³C 30 chemical shifts of aliphatic side chains. Care was taken to implement the ¹³C evolution in 31 the constant-time mode in order to refocus homonuclear ¹³C couplings of aliphatic carbons 32 33 that easily show up in highly resolved projections and can complicate the analysis of crowded regions. The modified pulse sequence (4D CT-HCCCON) and a few projections to
 illustrate the quality of the spectra are reported in the Supplementary Material (Figures S1
 and S2).

4 A final comment is due about the amide proton resonances that are still observable approaching physiological conditions. Combination of the C[´] detected experiments with H^N 5 detected ones provides a useful tool to achieve unambiguous assignment of the amide 6 proton resonances that are still observable in these conditions. In our case we collected a 7 8 3D HNCO experiment, also in the APSY mode, in order to correlate H^N resonances with the C'_{i-1}-N_i resonances that we have assigned through C[´] detected experiments. Through this 9 approach 93 cross peaks were detected and, thanks to the peak list obtained in a fully 10 automated manner through the GAPRO analysis of the APSY HNCO, the available 11 assignment of heteronuclear spins could easily be extended to the residual H^N resonances 12 that can be detected at this temperature. It is worth noting that a similar result could be 13 14 achieved by following the temperature dependence of the signals. The latter approach however works well only for well resolved H^N cross peaks but it becomes quite ambiguous 15 for cross peaks in crowded regions of the 2D ¹H-¹⁵N HSQC spectra. Therefore the 16 combination of C[´] detected experiments with H^N detected ones, even at high temperature 17 and pH, provides unambiguous information to assign H^N resonances that remain also in 18 presence of fast exchange, without ambiguities resulting from temperature dependence of 19 the signals and/or from incomplete information provided by H^N detected spectra. The 20 temperature dependence of H^N, C['] and N spins, once unambiguous assignment is available, 21 might provide also valuable information to investigate variations of exchange processes of 22 amide protons with the solvent upon increasing temperature as well as to highlight possible 23 24 variations in partially populated local conformations by analyzing heteronuclear chemical shifts. The sequence specific assignment of α-synuclein achieved at 315 K has been 25 26 deposited in the BMRB (ID 27348).

27 Concluding, we demonstrate the implementation of the APSY strategy in C⁻ detected 28 multidimensional NMR experiments. Excellent resolution can be achieved for the 2D cross 29 sections of the high dimensional C⁻ detected exclusively heteronuclear spectra, showing 30 that the two approaches (APSY and C⁻ detection) mutually benefit by their joint 31 implementation.

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1 Acknowledgments

2 The support and the use of resources of the CERM/CIRMMP center of Instruct-ERIC, a

- 3 Landmark ESFRI project, is gratefully acknowledged. This work has been supported in part
- 4 by a grant of the Fondazione CR di Firenze and by MEDINTECH (CTN01_001177_962865).

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1 References

2

Ambadipudi S, Zweckstetter M (2016) Targeting intrinsically disordered proteins in rational drug discovery.
 Expert Opin Drug Discov 11:65-77

- Baias M, Smith PE, Shen K, Joachimiak LA, Zerko S, Koźmiński W, Frydman J, Frydman L (2017) Structure
 and dynamics of the Huntingtin exon-1 N-terminus: a solution NMR perspective. J Am Chem Soc 139:11681176
- Bermel W, Bertini I, Csizmok V, Felli IC, Pierattelli R, Tompa P (2009) H-start for exclusively heteronuclear
 NMR spectroscopy: the case of intrinsically disordered proteins. J Magn Reson 198:275-281
- Bermel W, Bertini I, Felli IC, Kümmerle R, Pierattelli R (2006) Novel ¹³C direct detection experiments, including extension to the third dimension, to perform the complete assignment of proteins. J Magn Reson 178:56-64
- Bermel W, Bertini I, Felli IC, Lee Y-M, Luchinat C, Pierattelli R (2006) Protonless NMR experiments for sequence-specific assignment of backbone nuclei in unfolded proteins. J Am Chem Soc 128:3918-3919
- Bermel W, Bertini I, Felli IC, Piccioli M, Pierattelli R (2006) ¹³C-detected protonless NMR spectroscopy of proteins in solution. Progr NMR Spectrosc 48:25-45
- Bermel W, Bertini I, Felli IC, Pierattelli R (2009) Speeding up ¹³C direct detection Biomolecular NMR
 experiments. J Am Chem Soc 131:15339-15345
- 18 Bermel W, Bertini I, Gonnelli L, Felli IC, Koźmiński W, Piai A, Pierattelli R, Stanek J (2012) Speeding up 19 sequence specific assignment of IDPs. J Biomol NMR 53:293-301
- Bermel W, Bruix M, Felli IC, Kumar VMV, Pierattelli R, Serrano S (2013) Improving the chemical shift dispersion
 of multidimensional NMR spectra of intrinsically disordered proteins. J Biomol NMR 55:231-237
- Bermel W, Felli IC, Gonnelli L, Koźmiński W, Piai A, Pierattelli R, Zawadzka-Kazimierczuk A (2013) High dimensionality ¹³C direct-detected NMR experiments for the automatic assignment of intrinsically disordered
 proteins. J Biomol NMR 57:353-361
- Bermel W, Felli IC, Kümmerle R, Pierattelli R (2008) ¹³C direct-detection biomolecular NMR. Concepts Magn
 Reson 32A:183-200
- Böhlen J-M, Bodenhausen G (1993) Experimental aspects of chirp NMR spectroscopy. J Magn Reson Ser A
 102:293-301
- Brutscher B, Felli IC, Gil-Caballero S, Hošek T, Kümmerle R, Piai A, Pierattelli R, Sólyom Z (2015) NMR
 Methods for the study of instrinsically disordered proteins structure, dynamics, and interactions: General
 overview and practical guidelines. Adv Exp Med Biol 870:122
- Dyson HJ, Wright PE (2001) Nuclear magnetic resonance methods for the elucidation of structure and
 dynamics in disordered states. Methods In Enzymology 339:258-271
- Emsley L, Bodenhausen G (1992) Optimization of shaped selective pulses for NMR using a quaternion description of their overall propagators. J Magn Reson 97:135-148
- 36 Felli IC, Pierattelli R (Eds), Intrisically disordered proteins studied by NMR spectroscopy. Springer Switzerland
- Felli IC, Pierattelli R (2015) Spin-state-selctive methods in solution- and solid-state biomolecular ¹³C NMR.
 Prog NMR Spectrosc 84:1-13
- 39 Felli IC, Pierattelli R, Glaser SJ, Luy B (2009) Relaxation-optimised Hartmann-Hahn transfer for carbonyl-
- 40 carbonyl correlation spectroscopy using a specifically tailored MOCCA-XY16 mixing sequence for protonless
- 41 ¹³C direct detection experiments. J Biomol NMR 43:187-196

- Fiorito F, Hiller S, Wider G, Wüthrich K (2006) Automated resonance assignment of proteins: 6D APSY-NMR.
 J Biomol NMR 35:27-37
- Gil S, Hošek T, Solyom Z, Kümmerle R, Brutscher B, Pierattelli R, Felli IC (2013) NMR studies of intrinsically
 disordered proteins near physiological conditions. Angew Chem Int Ed 52:11808-11812
- Haba NY, Gross R, Nováček J, Shaked H, Židek L, Barda-Saad M, Chill JH (2013) NMR determines transient
 structure and dynamics in the disordered C-terminal domain of WASp interacting protein. Biophysical J
 105:481-493
- Habchi J, Tompa P, Longhi S, Uversky VN (2014) Introducing protein intrinsic disorder. Chem Rev 114:6561 6588
- Hiller S, Fiorito F, Wüthrich K, Wider G (2005) Automated projection spectroscopy (APSY). Proc Natl Acad Sci
 USA 102:10876-10881
- 12 Hiller S, Wider G (2012) Automated projection spectroscopy and its applications. Top Curr Chem 316:21-47
- Hiller S, Wider G, Wüthrich K (2008) APSY-NMR with proteins: practical aspects and backbone assignment.
 J Biomol NMR 42:179-195
- 15 Hoch JC Stern AS (2001) Nuclear Magnetic Resonance of Biological Macromolecules. pp 159-178
- Huang C, Ren G, Zhou H, Wang C (2005) A new method for purification of recombinant human alpha-synuclein
 in Escherichia coli. Protein Expr Purif 42:173-177
- Joshi P, Chia S, Habchi J, Knowles TPJ, Dobson CM, Vendruscolo M (2016) A fragment-based method of creating small-molecule libraries to target the aggregation of intrinsically disordered proteins. ACS Comb Sci 18:144-153
- Kadkhodaie M, Rivas O, Tan M, Mohebbi A, Shaka AJ, (1991) Broadband homonuclear cross polarization
 using flip-flop spectroscopy, J Magn Reson. 91:437–443
- Kay LE, Ikura M, Tschudin R, Bax A (1990) Three-dimensional triple-resonance NMR spectroscopy of
 isotopically enriched proteins. J Magn Reson 89:496-514
- Kazimierczuk K, Zawadzka A, Koźmiński W, Zhukov I (2006) Random sampling of evolution time space and
 Fourier transform processing. J Biomol NMR 36:157-168
- Kim S, Szyperski T (2003) GFT NMR, a new approach to rapidly obtain precise high-dimensional NMR spectral
 information. J Am Chem Soc 125:1385-1393
- Kovacs H, Moskau D, Spraul M (2005) Cryogenically cooled probes a leap in NMR technology. Prog NMR
 Spectrosc 46:131-155
- Kupce E, Freeman R (2004) Projection-reconstruction technique for speeding up multidimensional NMR
 spectroscopy. J Am Chem Soc 126:6429-6440
- Mäntylahti S, Hellman M, Permi P (2011) Extension of the HA-detection based approach: (HCA)CON(CA)H
 and (HCA)NCO(CA)H experiments for the main-chain assignment of intrinsically disordered proteins. J Biomol
 NMR 49:99-109
- Markley JL, Bax A, Arata Y, Hilbers CW, Kaptein R, Sykes BD, Wright PE, Wüthrich K (1998)
 Recommendations for the presentation of NMR structures of proteins and nucleic acids. IUPAC-IUPMB-IUPAB
 inter-union task group on the standardization of data bases of protein and nucleic acid structures determined
 by NMR spectroscopy. Eur J Biochem 256:1-15
- Motackova V, Nováček J, Zawadzka-Kazimierczuk A, Kazimierczuk K, Židek L, Sanderová H, Krásný L,
 Koźmiński W, Sklenář V (2010) Strategy for complete NMR assignment of disordered proteins with highly
 repetitive sequences based on resolution-enhanced 5D experiments. J Biomol NMR 48:169-177

Narayanan RL, Duerr HN, Bilbow S, Biernat J, Mendelkow E, Zweckstetter M (2010) Automatic assignment of
 the intrinsically disordered protein Tau with 441-residues. J Am Chem Soc 132:11906-11907

Nováček J, Haba NY, Chill JH, Židek L, Sklenář V (2012) 4D Non-uniformly sampled HCBCACON and ¹J_{NCα} selective HCBCANCO experiments for the sequential assignment and chemical shift analysis of intrinsically
 disordered proteins. J Biomol NMR 53:139-148

Novácek J, Zawadzka-Kazimierczuk A, Papoušková V, Zídek L, Sanderová H, Krásný L, Koźmiński W, Sklenář
 V (2011) 5D ¹³C-detected experiments for backbone assignment of unstructured proteins with a very low signal
 dispersion. J Biomol NMR 50:1-11

- 9 Nováček J, Zawadzka-Kazimierczuk A, Papoušková V, Židek L, Sanderová H, Krásný L, Koźmiński W, Sklenář
- V (2011) 5D ¹³C-detected experiments for backbone assignment of unstructured proteins with a very low signal
 dispersion. J Biomol NMR 50:1-11
- 12 O'Hare B, Benesi AJ, Showalter SA (2009) Incorporating ¹H chemical shift determination into ¹³C-direct 13 detected spectroscopy of intrinsically disordered proteins in solution, J Magn Reson, 200:354-358
- Orekhov VY, Ibraghimov I, Billeter M (2003) Optimizing resolution in multidimensional NMR by three-way decomposition. J Biomol NMR 27:165-173
- Piai A, Hošek T, Gonnelli L, Zawadzka-Kazimierczuk A, Koźmiński W, Brutscher B, Bermel W, Pierattelli R,
 Felli IC (2014) "CON-CON" assignment strategy for highly flexible intrinsically disordered proteins. J Biomol
 NMR 60:209-218
- Sattler M, Schleucher J, Griesinger C (1999) Heteronuclear multidimensional NMR experiments for the
 structure determination of proteins in solution employing pulsed field gradients. Progr NMR Spectrosc 34:93 158
- Shaka AJ, Barker PB, Freeman R (1985) Computer-optimized decoupling scheme for wideband applications
 and low-level operation. J Magn Reson 64:547-552
- Tóth G, Gardai SJ, Zago W, Bertoncini CW, Cremades N, Roy SL, Tambe MA, Rochet JC, Galvagnion C,
 Skibinski G, Finkbeiner S, Bova M, Regnstrom K, Chiou SS, Johnston J, Callaway K, Anderson JP, Jobling
 MF, Buell AK, Yednock TA, Knowles TP, Vendruscolo M, Christodoulou J, Dobson CM, Schenk D,
 McConlogue L (2014) Targeting the intrinsically disordered structural ensemble of α-synuclein by small
 molecules as a potential therapeutic strategy for Parkinson's disease. Plos ONE 9:e87133
- Uversky V, Oldfield CJ, Dunker AK (2008) Intrinsically disordered proteins in human diseases: introducing the
 D2 concept. Annu Rev Biophys 37:215-246
- Uversky VN, Davé V, lakoucheva LM, Malaney P, Metallo SJ, Pathak RR, Joerger AC (2014) Pathological
 unfoldomics of uncontrolled chaos: intrinsically disordered proteins and human diseases. Chem Rev 114:6844 6879
- van der Lee R, Buljan M, Lang B, Weatheritt RJ, Daughdrill GW, Dunker AK, Fuxreiter M, Gough J, Gsponer
 J, Jones DT, Kim PM, Kriwacki RW, Oldfield CJ, Pappu RV, Tompa P, Uversky VN, Wright PE, Babu MM
 (2014) Classification of intrinsically disordered regions and proteins. Chem Rev 114:6589-6631
- Wright PE, Dyson HJ (2015) Intrinsically disordered proteins in cellular signalling and regulation. Nat Rev Mol
 Cell Biol 16:18-29
- Yao X, Stefan B, Zweckstetter M (2014) A six-dimensional alpha proton detection-based APSY experiment
 for backbone assignment of intrinsically disordered proteins. J Biomol NMR 60: 231-240
- Zawadzka-Kazimierczuk A, Kazimierczuk K, Koźmiński W (2010) A set of 4D NMR experiments of enhanced
 resolution for easy resonance assignment in proteins. J Magn Reson 202:109-116

- Zawadzka-Kazimierczuk A, Koźmiński W, Sanderová H, Krásný L (2012) High dimensional and high resolution
- pulse sequences for backbone resonance assignment of intrinsically disordered proteins. J Biomol NMR
- 52:329-337

1 Figures

Figure 1. ¹H-¹⁵N HSQC (top) and ¹³C-¹⁵N CON-IPAP spectra (bottom), acquired at different
temperatures (285.5 K, 295 K, 305 K, 315 K), showing the temperature dependence for αsynuclein signals. Experiments were acquired at 16.4 T on α-synuclein at pH 6.5.

Figure 2. A subset of the 2D projections acquired with the 5D (HCA)CONCACON 5 experiment are reported to show the quality of the data and to provide an intuitive picture of 6 how the APSY approach works with C' detected experiments on IDPs (Bermel et al., 2013). 7 Orthogonal projections (A - D) and non-orthogonal ones (E - F) are shown. The spectra 8 9 were recorded at 16.4 T on α -synuclein at 315 K. The four orthogonal projections correspond to C'(i)-N(i+1), C'(i)-N(i), C'(i)-C'(i±1), and $C^{\alpha}(i)$ -C'(i). In the two non-orthogonal projections 10 the y-coordinate units are arbitrary, as the frequencies in the indirect dimensions are a linear 11 combination of frequencies, that is (E) $\omega_2 \sin(45^\circ) + \omega_4 \cos(45^\circ)$ and (F) $\omega_1 \sin(55^\circ) + \omega_2 \sin(45^\circ)$ 12 ω₄cos(55°). 13

Figure 3. As an example the assignment strategy using the peak lists resulting from the GAPRO analysis of the 5D (HCA)CONCACON projections is shown for residues 65-67 of α -synuclein (Bermel et al., 2013). Four five-dimensional "type I" correlations (C'_{i-1}, N_i, C^{α}_i, C'_i, N_{i+1}) are shown. It is worth noting that the amide proton resonances of these residues were not detectable.

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20 Tables

Table 1. Acquisition parameters for 5D ^{APSY}-(HCA)CONCACON (Bermel et al., 2013) and
 4D ^{APSY}-HCCCON experiments (Bermel et al., 2012).

Table 2. Values of the projection angles α , β , γ , and of the spectral widths in the dimensions ω_{1-4} used here for recording the 2D-projections of the 5D ^{APSY}-(HCA)CONCACON experiment (Bermel et al., 2013). The resulting linear combinations of frequencies are given in the right column.

Table 3. Analysis of the 5D ^{APSY}-(HCA)CONCACON processed considering a reduced
 number of projection angles as described in the text.

1 Figure 1





176 174 δ(¹³C)/ppm

178 176 174 δ(¹³C)/ppm





Table 1

EXPERIMENT	Spectral width (Hz) & Maximal evolution times				NS	Inter scan delay	Dimension of acquired data				data	Duration of the	Number of projections
	F1	F2 / F4	F3	F5		(s)	F1	F2	F3	F4	F5	experiment	
5D (HCA)CONCACON	1695 (¹³ C') 28.4 ms	2404 (¹⁵ N) 53.2 ms	4505 (¹³ C ^α) 75.5 ms	9090 (¹³ C') 112.6 ms	8	0.8	256	256	256	256	2048	4 days 18 hours	64
	F1	F2	F3	F4	4	0.8	512	512	512	2048	/	2 days 23 hours	
4D HCCCON	3840 (¹ H) 66.6 ms	10870 (¹³ C ^{ali}) 23.6 ms	2404 (¹⁵ N) 106.5 ms	9090 (¹³ C') 112.6 ms									41

α	β	γ	Spectral width (Hz)	Linear combination
0°	0°	0°	2404	ω ₄
0°	0°	90°)° 1695 ω ₁	
0°	90°	0°	2404	ω ₂
90°	0°	0°	4505	ω3
±28°	0°	0°	4238	$\omega_4 \cos(28^\circ) \pm \omega_3 \sin(28^\circ)$
0°	± 45°	0°	3400	$\omega_4 \cos(45^\circ) \pm \omega_2 \sin(45^\circ)$
0°	0°	± 55°	2767	$\omega_4 \cos(55^\circ) \pm \omega_1 \sin(55^\circ)$
90°	± 62°	0°	4238	$\omega_3 \cos(62^\circ) \pm \omega_2 \sin(62^\circ)$
90°	0°	± 69°	3197	ω₃cos(69°) ± ω₁sin(69°)
0°	90°	± 55°	2767	$\omega_2 \cos(55^\circ) \pm \omega_1 \sin(55^\circ)$
±28°	± 41°	0	4775	$\omega_4 \cos(28^\circ)\cos(41^\circ) \pm \omega_2 \sin(41^\circ) \pm \omega_3 \sin(28^\circ)\cos(41^\circ)$
±28°	0°	± 51°	3984	$\omega_4 \cos(28^\circ)\cos(51^\circ) \pm \omega_1 \sin(51^\circ) \pm \omega_3 \sin(28^\circ)\cos(51^\circ)$
0°	± 45°	± 45°	3603	$\omega_4 \cos(45^\circ) \cos(45^\circ) \pm \omega_1 \sin(45^\circ) \pm \omega_2 \sin(45^\circ) \cos(45^\circ)$
90°	± 62°	± 51°	3984	$ω_3$ cos(62°)cos(51°) ± $ω_1$ sin(51°) ± $ω_2$ sin(62°)cos(51°)
±15°	0°	0°	3488	$\omega_4 \cos(15^\circ) \pm \omega_3 \sin(15^\circ)$
0°	± 27°	0°	3233	ω ₄ cos(27°) ± ω ₂ sin(27°)
0°	0°	± 35°	2941	ω ₄ cos(35°) ± ω ₁ sin(35°)
90°	± 43°	0°	4934	$\omega_3 \cos(43^\circ) \pm \omega_2 \cos(43^\circ)$
90°	0°	± 53°	4065	ω₃cos(53°) ± ω₁sin(53°)
0°	90°	± 35°	2941	ω₂cos(35°) ± ω₁sin(35°)
±47°	0°	0°	4934	ω₄cos(47°) ± ω₃sin(47°)
0°	± 63°	0°	3233	ω₄cos(63°) ± ω₂sin(63°)
0°	0°	± 71°	2385	ω ₄ cos(71°) ± ω ₁ sin(71°)
90°	± 75°	0°	3488	ω₃cos(75°) ± ω₂cos(75°)
90°	0°	±79°	2523	ω₃cos(79°) ± ω₁sin(79°)
0°	90°	± 71°	2385	₀₂cos(71°) ± ₀₁sin(71°)
±15°	0°	0°	3488	⊛₄cos(15°) ± ω₃sin(15°)
0°	±20°	0°	3081	ω₄cos(20°) ± ω₂sin(20°)
±70°	0°	0°	5056	
0°	±70°	0°	3081	ω₄cos(70°) ± ω₂sin(70°)

Table 3

Angle	Projections	Experimental	Identified correlations		
sets		time	(Type I)		
30	64	4d, 18h	131		
26	56	4d, 5h	131		
20	44	3d, 9h	129		
10	16	1d, 2h	116		

¹³C APSY-NMR for sequential assignment of intrinsically disordered proteins

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Supporting material

Figure S1. Pulse scheme for the 4D CT-HCCCON. The following phase cycling was employed: $\phi_1 = x, -x$; $\phi_2 = 2(y), 2(-y)$; $\phi_3 = 4(x), 4(-x)$; $\phi_4 = 16(x), 16(-x)$; $\phi_5 = 8(x), 8(-x), \phi_{rec} = x, 2(-x), x, (-x), 2(x), (-x), 2(x), (-x), x, 2(-x), x$ The quadrature detection was achieved through the States-TPPI approach by incrementing the phase of the $\pi/2$ pulse prior to the evolution period; C´ homonuclear decoupling was achieved using IPAP scheme. The length of the delays were: $\delta = 0.475$ ms, $\Delta = 56$ ms, $\Delta_1 = 9$ ms, $\Delta_2 = 25$ ms – p₁₂, $\epsilon = t_3(0) + p_8$. t_a, t_b and t_c were used to achieve the semi-constant time mode for the ¹H indirect dimension. The ¹H carrier was placed in the middle of the ¹H aliphatic region (2.5 ppm) during the evolution block while it was set at 4.7 ppm in the remaining parts of the pulse sequence.

Figure S2. Examples of 2D projections acquired with the 4D CT-HCCCON experiment to show the quality of the data. Three orthogonal projections (A – C) and a non-orthogonal one (D) are shown. The spectra were recorded at 700 MHz on α -synuclein at 315 K. The three orthogonal projections correspond to C'(i)-N(i+1), C'(i)-Hali(i), C'(i)-Cali(i). The non-orthogonal projection has, as indirect dimension, the combination of $\omega_1 \sin(-32^\circ) + \omega_3 \cos(-32^\circ)$.







