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## Solid-state NMR sequential assignments of the amyloid core of full-length Sup35p

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**Abstract** Sup35p is a yeast prion and is responsible for the [PSI<sup>+</sup>] trait in Saccharomyces cerevisiae. With 685 amino acids, full-length soluble and fibrillar Sup35p are challenging targets for structural biology as they cannot be investigated by X-ray crystallography or NMR in solution. We present solid-state NMR studies of fibrils formed by the full-length Sup35 protein. We detect an ordered and rigid core of the protein that gives rise to narrow and strong peaks, while large parts of the protein show either static disorder or dynamics on time scales which interfere with dipolar polarization transfer or shorten the coherence lifetime. Thus, only a small subset of resonances is observed in 3D spectra. Here we describe in detail the sequential assignments of the 22 residues for which resonances are observed in 3D spectra: their chemical shifts mostly corresponding to  $\beta$ -sheet secondary structure. We suspect that these residues form the amyloid core of the fibril.

Anne K. Schütz, Nina Luckgei, Birgit Habenstein, and Luc Bousset have contributed equally to this work.

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B. Habenstein · N. Luckgei · A. Böckmann (⊠) Institut de Biologie et Chimie des Protéines, UMR 5086 CNRS/ Université de Lyon 1, 7 passage du Vercors, 69367 Lyon, France e-mail: a.bockmann@ibcp.fr

L. Bousset · Y. Sourigues · R. Melki (⊠) Laboratoire d'Enzymologie et Biochimie Structurales, UPR 3082 CNRS, Avenue de la Terrasse, 91198 Gif-sur-Yvette, France e-mail: melki@lebs.cnrs-gif.fr **Keywords** Sup35p · Fibrils · Solid-state NMR · Assignments · Secondary structure

#### **Biological context**

Sup35p is a yeast prion from *Saccharomyces cerevisiae* and is at the origin of the [*PSI*<sup>+</sup>] phenotype (Cox 1965; Wickner et al. 1995). It contains 685 amino acids and consists of three domains. The N-terminal domain (N-domain), comprising residues 1–123, is rich in glutamine and asparagine residues. It has been proposed that the N-domain plays a critical role in prion propagation (DePace et al. 1998). The middle domain (M-domain) with unknown function mainly consists of highly charged residues. The C-terminal domain, spanning residues 254–685, has GTPase activity and is involved in translation termination (Stansfield et al. 1995). Since Sup35p is not infectious for human beings, it is well suited as a model system to gain deeper insight into the structural features of prion proteins.

Until now most of the structural and functional studies concerning Sup35p have been performed on small fragments of the N-domain, as for example Sup35p 7–13, or on the N-domain by itself or together with the M-domain, Sup35pN and Sup35pNM, respectively (Nelson et al. 2005; Krishnan and Lindquist 2005; Shewmaker et al. 2006; van der Wel et al. 2006, 2007; Toyama et al. 2007; van der Wel et al. 2010; Vranken et al. 2005). Sup35p and Sup35pNM both form selfseeding amyloid fibrils in vivo and in vitro under physiological pH and salt concentrations. However, it has been suspected that the two proteins differ in their assembly and infection properties (Krzewska et al. 2007). The assignment of Sup35pNM is described in a companion paper (Luckgei et al. 2013b). A detailed comparison and a biophysical interpretation will be presented elsewhere (Luckgei et al. 2013a). Solid-state NMR spectroscopy has evolved into a powerful tool to structurally investigate protein fibrils (Tycko 2006; Wasmer et al. 2008; Heise et al. 2008; Paravastu et al. 2008; Loquet et al. 2009; van Melckebeke et al. 2010; Böckmann and Meier 2010; Debelouchina et al. 2010; Loquet et al. 2010; van der Wel et al. 2010; Lewandowski et al. 2011; Comellas et al. 2011; Gath et al. 2012; Fitzpatrick et al. 2013). Here we show that even large proteins like Sup35p are amenable to solid-state NMR studies and present the sequential assignments of the resonances visible in 3D correlation spectra, which are all located within the first 30 residues of the protein.

#### Methods and experiments

Protein expression and purification; sample preparation

Sup35p expression and purification were done as described in Krzewska and Melki (2006), Krzewska et al. (2007). For convenience of the reader, we summarize it in the

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**Fig. 1** Amino-acid sequence of Sup35p, including the purification tag added at the N-terminus (*in grey*). The N domain is shown in *light blue*, and the M domain in *deep blue*, and the C-terminal domain in *black* 

following. The proteins were expressed with an N-terminal His-tag (for the full amino acid sequence see Fig. 1) and purified as described, using M9 medium with <sup>13</sup>C and <sup>15</sup>N labeling. To form fibrillar aggregates, Sup35p (in 20 mM Tris HCl, pH 8.0, 200 mM NaCl, 5 % glycerol, 5 mM  $\beta$ -mercaptoethanol, 1 mM ATP, 1 mM GTP, 10 mM MgCl<sub>2</sub>) was incubated at 10 °C under orbital shaking at 30 rpm, 0.5 cm amplitude, for 3 weeks. The fibrils were spun at 100,000g in a TL-100 tabletop centrifuge for 20 min at 4 °C. The pellets were resuspended in distilled water, washed twice, and filled into 3.2 mm rotors by ultracentrifugation as described (Böckmann et al. 2009).

#### NMR spectroscopy

2D and 3D experiments were used for assignments: 2D DARR and DREAM, 3D NCACB, NCACO, NCACX, NCOCA, NCOCX, CANCO and CCC. The procedure is described in detail in references Habenstein et al. (2011) and Schuetz et al. (2010). 3D spectra were recorded for 2–6 days each (details in Table 1). Resonances were

6 <u>0</u> GYSGYQQGGY	5 <u>0</u> PAGGYYQNYQ	4 <u>0</u> GYQAYNAQAQ	3 <u>0</u> NQQQGNNRYQ	20 QNYQQYSQNG	MGSSHHHHHH 1 <u>0</u> MSDSNQGNNQ
12 <u>0</u>	11 <u>0</u>	10 <u>0</u>	9 <u>0</u>	8 <u>0</u>	7 <u>0</u>
QGYQAGFQPQ	YKNFNYNNNL	QFNPQGGRGN	QYNPQGGYQQ	QQYNPQGGYQ	QQYNPDAGYQ
18 <u>0</u>	17 <u>0</u>	16 <u>0</u>	15 <u>0</u>	14 <u>0</u>	13 <u>0</u>
KEEEKSAETK	VGTKPAESDK	GIKLANATKK	KKTLKLVSSS	KQQKQAAPKP	SQGMSLNDFQ
24 <u>0</u>	23 <u>0</u>	22 <u>0</u>	21 <u>0</u>	20 <u>0</u>	19 <u>0</u>
NVTSADALIK	SESTHNTNNA	ELPKVEDLKI	QTEEKTEEKS	EPVKKEEKPV	Eptkeptkve
30 <u>0</u>	29 <u>0</u>	28 <u>0</u>	27 <u>0</u>	26 <u>0</u>	25 <u>0</u>
TIEKYEREAK	LYLTGSVDKR	AGKSTMGGNL	VSLIFMGHVD	VNDMFGGKDH	EQEEEVDDEV
36 <u>0</u>	35 <u>0</u>	34 <u>0</u>	33 <u>0</u>	32 <u>0</u>	31 <u>0</u>
YVSEMIGGAS	TILDAPGHKM	Ayfetekrry	NDGKTIEVGK	wvmdtnkeer	DAGRQGWYLS
42 <u>0</u>	41 <u>0</u>	40 <u>0</u>	39 <u>0</u>	38 <u>0</u>	37 <u>0</u>
PTVNWSKERY	MVVVVNKMDD	llaktqgvnk	ERGGQTREHA	ARKGEYETGF	QADVGVLVIS
48 <u>0</u>	47 <u>0</u>	46 <u>0</u>	45 <u>0</u>	44 <u>0</u>	43 <u>0</u>
TLLEYLDTMN	PKECPWYTGP	SGANLKDHVD	DVVFMPVSGY	LRAIGYNIKT	DQCVSNVSNF
54 <u>0</u>	53 <u>0</u>	52 <u>0</u>	51 <u>0</u>	50 <u>0</u>	49 <u>0</u>
IQNIYNETEN	LLMPNKTAVE	SGHIKKGQST	LGTIVEGKIE	MLPIAAKMKD	HVDRHINAPF
60 <u>0</u>	59 <u>0</u>	58 <u>0</u>	57 <u>0</u>	56 <u>0</u>	55 <u>0</u>
LKSIIAAGFS	KFVAQIAIVE	SPKNPIKSVT	EDISPGFVLT	VKLRIKGVEE	EVDMAMCGEQ
66 <u>0</u>	65 <u>0</u>	64 <u>0</u>	63 <u>0</u>	62 <u>0</u>	61 <u>0</u>
PVCVETYQDY	KVIAVLETEA	KPPAFAKKGM	LEKGTNRKSK	EVHIVKLLHK	CVMHVHTAIE
			VKIAE	68 <u>0</u> QGTTIAIGKI	67 <u>0</u> PQLGRFTLRD

#### Table 1 Experimental parameters

<u>a</u>						
Experiment	DARR	DREAM	NCA	NCACB	NCACO	NCOCA
MAS frequency (kHz)	18.0	17.5	17.5	17.5	17.5	17.5
Transfer 1	HC-CP	HC-CP	HN-CP	HN-CP	HN-CP	HN-CP
Field (kHz)- <sup>1</sup> H	65.5	72.6	54.4	54.4	58.3	59.0
Field (kHz)-X	55.5	62.3	40.6	41.6	45.6	45.6
Shape	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H
<sup>13</sup> C carrier (ppm)	55.6	55.6	_	_	_	_
Time (ms)	0.5	0.5	0.8	0.8	0.9	0.8
Transfer 2	DARR	DREAM	NC-CP	NC-CP	NC-CP	NC-CP
Field (kHz)- <sup>1</sup> H	5.5	102	102	115	102	102
Field (kHz)- <sup>13</sup> C	_	8.3	10.6	12.5	10.6	10.8
Field (kHz)- <sup>15</sup> N	_	_	7.5	6.5	7.5	8.4
Shape	_	Tangent <sup>13</sup> C				
Carrier (ppm)	_	57	60.2	60.2	60.2	177.6
Time (ms)	20	4.0	5.0	6.0	5.5	5.5
Transfer 3				DREAM	MIRROR	MIRROR
Field (kHz)- <sup>1</sup> H				115.3	21.6	22.1
Field $(kHz)^{-13}C$				8.3	_	_
Field $(kHz)^{-15}N$				_	_	_
Shape				Tangent <sup>13</sup> C	_	_
Carrier (npm)				57	_	_
Time (ms)				4	25	20
t1 increments	768	980	640	112	96	20 96
Sween width (t1) (kHz)	50	70	25	6	5	5
Acquisition time (t1) (ms)	50 7 7	70	12.0	0 3	96	96
t2 ingraments	2.048	7.0	2.048	9.5	9.0	9.0
$t_2$ increments Swaap width $(t_2)$ $(t_1)$	2,048	2,048	2,048	80	144	144
Sweep with $(t_2)$ (KIZ)	10.2	10.2	10.2	5.0	12	12
t2 incremente	10.2	10.2	10.2	J.0 1.526	0	2.048
Successive (2) (111-)				1,550	2,040	2,048
A semisition time (t2) (ma)				100	100	100
Acquisition time (t3) (ms)	100	100	100	/./	10.2	10.2
H Spinal64 decoupling power (kHz)	100	100	100	100	100	100
Interscan delay (s)	2.5	2.0	2.2	2.2	2.3	2.3
Number of scans	20	16	40	8	16	8
Measurement time	ll h	9 h	16 h	2 days	6 days	3 days
D Experiment	NCACY	NCOCY	CANCO	000	INEDT	СИ/СИ
		17.5		17.5	17.5	
MAS freq. (kHz)	17.5	17.5	17.5	17.5	17.5	18.0
Transfer 1	HN-CP	HN-CP	HC-CP	HC-CP	INEPT	HC-CP, $n = 0$
Field (kHz)-'H	54.4	55.5	71.8	72.6	62.5	45
Field (kHz)-X	40.6	45.6	63.0	63.7	50	45
Shape	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Tangent <sup>1</sup> H	Hard pulses	
Carrier (ppm)	-	-	48.5	55.6	0	62.5
Time (ms)	0.8	1.0	0.5	0.5		0.7
Transfer 2	NC-CP	NC-CP	CN-CP	DREAM		HC-CP, $n = 0$
Field (kHz)- <sup>1</sup> H	96	82	121	101.5		45
Field (kHz)- <sup>13</sup> C	11.1	10.6	10.8	8.2		45

#### Table 1 continued

Experiment	NCACX	NCOCX	CANCO	CCC	INEPT	CH/CH <sub>2</sub>		
Field (kHz)- <sup>15</sup> N	6.7	9.0	6.6	_		_		
Shape	Tangent <sup>13</sup> C	Tangent <sup>13</sup> C	Tangent <sup>13</sup> C	Tangent <sup>13</sup> C		-		
Carrier (ppm)	57.9	178.6	58.0	57.0		62.5		
Time (ms)	5.0	5.0	8.0	4.0		$0.278~(5 \times \tau_r)$		
Transfer 3	DARR	DARR	NC-CP	DARR		DARR		
Field (kHz)- <sup>1</sup> H	11.8	7.5	110	3.4		11.8		
Field (kHz)- <sup>13</sup> C	-		10.8			-		
Field (kHz)- <sup>15</sup> N	-		8.1			-		
Shape	-		Tangent <sup>13</sup> C			-		
Carrier (ppm)	-		178.6			-		
Time (ms)	80	80	6.0	200		100		
t1 increments	80	72	128	248	192	1,024		
Sweep width (t1) (kHz)	4	4	10	22	8	100		
Acquisition (t1) (ms)	10.0	9.0	6.4	5.6	12.0	5.1		
t2 increments	128	96	96	248	3,072	2,048		
Sweep width (t2) (kHz)	10	8	4.8	22	100	100		
Acquisition (t2) (ms)	6.4	6.0	10.0	5.6	15.4	10.3		
t3 increments	2,048	2,048	2,048	2,048				
Sweepwidth (t3) (kHz)	100	100	100	100				
Acquisition (t3) (ms)	10.2	10.2	10.2	10.2				
<sup>1</sup> H Spinal64 decoupling power (kHz)	100		100	100	77.8	100		
Interscan delay (s)	2.2	2.2	2.35	1.9	2.5	2.2		
Number of scans	16	16	8	4	8	8		
Measurement time	4 days	3 days	3 days	6 days	1 h	13 h		



Fig. 2 2D NCA spectrum of Sup35p with labels for the sequentially assigned residues

connected via the sequential walk in NCACB/NCACO/ NCACX to CANCO and to NCOCA/NCOCX spectra. Side-chain assignments were achieved using NCACB, NCACX, NCOCX and CCC spectra. Experimental details are given in Table 1. The spectra used for assignments were all recorded on a single sample, but several other preparations were checked for reproducibility and yielded identical spectra.

All spectra were recorded on a Bruker Avance II+ 850 MHz spectrometer operating at a static field of 20 T. A 3.2 mm Bruker triple resonance MAS probe equipped with an LLC coil was used to reduce r.f. heating during the experiments. The spectra were recorded at sample temperatures of 7 °C. The pulse sequences were implemented as recently reported (Schuetz et al. 2010). All spectra were processed using TopSpin 2.0 (Bruker Biospin) by zero filling to no more than double the number of acquired points, the time domain signals were apodized with a squared cosine function (SSB 2.2–2.6 depending on spectrum and dimension). Spectra were analyzed and annotated using the CcpNmr Analysis package (Fogh et al. 2002; Vranken et al. 2005; Stevens et al. 2011).

#### Assignment and data deposition

The 2D NCA spectrum of Sup35p is shown in Fig. 2. The spectrum displays considerable overlap as can be expected for such a large protein. Many individual peaks are narrow however, as indicated by the isolated signals. One can already see in the 2D spectrum that there is a remarkably large distribution in peak intensities, and some isolated signals show higher intensities than crowded regions. Even stronger intensity variations are revealed in the 3D spectra (see Fig. 3 as an example), which actually show only a subset of the resonances observed in the 2D spectra; about 45 peaks can be counted in the NCACB spectrum. The linewidth of isolated correlations is mostly between 0.5 and 1 ppm (e.g. 0.6 ppm measured on Ser4 C $\alpha$  and C $\beta$  in the direct dimension), and the 3D spectra show a good resolution and dispersion, allowing for an assignment using the afore-mentioned 3D methods.

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22 of the NCACB signals could be sequentially assigned to N, C $\alpha$ , C $\beta$  triples, and 3 to further side-chain resonances of the assigned residues. Assignments are given on the 2D NCA spectrum in Fig. 2.

To illustrate sequential assignments, a representative plane of the 3D NCACB, NCACO, CANCO and NCOCA experiments is shown in Fig. 3. The 22 assigned residues are all situated in the N-terminal portion of the prion domain. The chemical shifts have been deposited in the BMRB under the accession number 18407.

It might be noted that the C $\beta$  chemical shift of Ser4 is atypically high with 70.9 ppm, which would rather correspond to the C $\beta$  chemical shift of threonine. Nevertheless, several arguments do support this assignment. The sequential assignment of Ser4 is unambiguous in both directions towards Asp3 and Asn5. Furthermore, the signals observed



**Fig. 4** CH versus CH<sub>2</sub> spectral editing for Sup35p. *Blue signals* correspond to positive contours and stem from CH or CH<sub>3</sub> groups (in  $\delta_1$ ). *Red signals* are negative and originate from CH<sub>2</sub> groups (see text). The C $\beta$ -C $\alpha$  and C $\beta$ -C' correlations of Ser4 have been labeled

Fig. 3 Representative planes of the 3D NCC assignment spectra. Experimental details are given in Table 1. NCACB spectra are represented in *purple* (negative signals) and *grey* (positive signals), NCACO spectra in *red*, CANCO spectra in *blue* and NCOCA spectra in *green*. Grey *labels* are used if the resonance has a slightly different <sup>15</sup>N chemical shift than the represented plane





Fig. 5 INEPT spectrum of the Sup35 prion. The signals are assigned to residue types using the random coil chemical shifts (Wang and Jardetzky 2002)



Fig. 6 Assignment graph created using the CCPNmr software (Fogh et al. 2002) of the N-terminal 30 amino acids of Sup35

in the spectra correspond to a serine spin system, since even at long DARR mixing times up to 200 ms no C $\gamma$ 2 shift was observed in the <sup>13</sup>C–<sup>13</sup>C-correlation spectra, although the C $\alpha$ –C $\beta$  correlation is generally one of the stronger peaks. Finally, in the BMRB 115 serine C $\beta$  shifts outliers have been deposited (more than three sigma deviation from the average shift), reaching up to 76.4 ppm. A closer look at the PDBdeposited corresponding structures reveals that out of the 26 structures that show a Ser C $\beta$  chemical shift above 70 ppm, more than 24 of the Ser are located in loops or unstructured regions. However, many of them lie just before or after a  $\beta$ strand. The remaining two Ser are located at the beginning of a  $\beta$ -sheet. Since Ser4 is the first residue of a  $\beta$ -strand, as indicated by secondary chemical shifts (see below), this might be the reason for its unusually high C $\beta$  chemical shift.

The assignment of Ser4 was verified independently by spectroscopic means using spectral editing techniques. In a Thr residue, the C $\beta$  atom is part of a CH group, whereas it is part of a CH<sub>2</sub> group in a Ser residue. An approach for spectral editing of CH versus CH<sub>2</sub> groups in the solid state has been proposed (Wu and Zilm 1993; Sangill et al. 1994), which relies on the presence of second-order cross terms

between  ${}^{1}H{-}^{1}H$  and  ${}^{1}H{-}^{13}C$  dipolar couplings in CH<sub>2</sub> groups. We have set up such an element followed by a DARR mixing time to resolve the signals in a 2D spectrum. An excerpt of the aliphatic region is shown in Fig. 4. The C $\beta$ -C $\alpha$  and C $\beta$ -C' correlations of Ser4 are inverted as would be expected for the CH<sub>2</sub> groups of Ser residues. One should note that the signal observed around 64/58 ppm corresponds to Ser random-coil chemical shifts, and thus stems from flexible residues. Their dipolar couplings are scaled and thus do not follow the editing rules for rigid residues. The inverted sign of the Ser4 correlations is another proof that despite the exotic C $\beta$  chemical shift, the residue in question must be a Ser, and cannot be a Thr.

As mentioned above, parts of the protein show either static disorder or flexibility. Highly dynamic residues generally can be detected in INEPT spectra. All amino-acid types with the exception of Arg, Cys, Trp and Tyr could be observed in this scalar-coupling based spectrum as shown in Fig. 5. The absence of these residues in the spectra coincides with their absence in the M-domain. In addition, all residue-types present in the M-domain show signals in the INEPT spectrum. **Fig. 7** Extract of *NCACB*, *NCACO*, and *CANCO* 3D spectra for Gly7 and an unassigned glycine residue. The absence of peaks in the *NCACO* and *CANCO* spectra shows that a sequential walk is not possible for this residue due to weak signal strength



Fig. 8 Secondary chemical shifts for Sup35p. Secondary shifts have been calculated as described in (Wishart and Sykes 1994). Three or more negative values in a row suggest  $\beta$ -strand conformation. Isolated positive or negative values for a given residue suggest turn- or random coil-conformation. For Gly only the secondary shift of C $\alpha$  is plotted

Assigned atoms are shown in the assignment graph in Fig. 6. Most strong cross peaks in the 2D NCA spectrum can be explained based on this assignment.

As mentioned above, assignments could not be obtained for all signals observed in the 3D spectra. A typical case of a signal present in the NCACB spectrum but not assignable due to absence of the corresponding peaks in the remaining 3D spectra is shown in Fig. 7, using the example of two glycine residues. In the more sensitive NCACB experiment, signals are observed for both glycines, however significantly weaker already for the upper one (labeled Gly). In the NCACO and CANCO spectra, shown in red and blue respectively, peaks are only observed for Gly7 and its connections to Gln6 (in the CANCO spectrum). For the other glycine, no signals allowing for sequential walk were detected.

The secondary chemical shifts for the assigned residues are given in Fig. 8. The assigned residues mostly display secondary chemical shifts typical for  $\beta$ -strand conformation, with the exception of residues Asp3 and Gly20 and Gly25. To summarize, our data suggest that Sup35p possesses an amyloid core formed by two  $\beta$ -strands running from residues 2–11 and 19–30. Large parts of the remaining regions of the protein are statically disordered or show mobility on time scales that interfere with dipolar polarization transfer or that shorten the coherence lifetime. The sequential assignment of the strong signals observed in the Sup35p spectra constitutes a first step towards a high-resolution structure of its hypothetical amyloid core.

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