RIDER distortions in the CODEX experiments

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Abstract. CSA and dipolar CODEX experiments enable obtaining abundant quantitative information on the reorientation of the CSA and dipolar tensors on the millisecond-second time scales. At the same time, proper performance of the experiments and data analysis can often be a challenge since CODEX is prone to some interfering effects that may lead to incorrect interpretation of the experimental results. One of the most important such effects is RIDER (Relaxation Induced Dipolar Exchange with Recoupling). It appears due to the dipolar interaction of the observed X-nuclei with some other nuclei, which causes an apparent decay in the mixing time dependence of the signal intensity reflecting not molecular motion but spin-flips of the adjacent nuclei. This may hamper obtaining correct values of the parameters of molecular mobility. In this contribution we consider in detail the reasons, why the RIDER distortions remain even under decoupling conditions and propose measures to eliminate them. Namely, we suggest 1) using an additional Z-filter between the cross-polarization

- 15 section and the CODEX recoupling blocks that suppresses the interfering anti-phase coherence responsible for the X-H RIDER and 2) recording only the cosine component of the CODEX signal since it is less prone to the RIDER distortions in comparison to the sine component. The experiments were conducted on rigid model substances as well as microcrystalline ²H/¹⁵N-enriched proteins (GB1 and SH3) with a partial back-exchange of labile protons. Standard CSA and dipolar CODEX experiments reveal a fast decaying component in the mixing time dependence of ¹⁵N nuclei in proteins, which can be
- 20 misinterpreted as a slow overall protein rocking motion. However, the RIDER-free experimental setup provides flat mixing time dependencies meaning that the studied proteins do not undergo global motions on the millisecond time scale.

1 Introduction

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CODEX (Cenralband Only Detection of EXchange) (deAzevedo et al., 1999; deAzevedo et al., 2000; Luz et al., 2002; Reichert and Krushelnitsky, 2018) is a powerful NMR tool for studying molecular dynamics in millisecond to second time scale under the magic angle spinning (MAS). It is based on the stimulated echo principle; the simplified pulse sequence is shown in Fig. 1. Depending on the phases of the rf-pulses and receiver, one may record a signal, which is proportional to $\sin(\Phi_1) \cdot \sin(\Phi_2)$ (SIN-component) or $\cos(\Phi_1) \cdot \cos(\Phi_2)$ (COS-component), where Φ_1 and Φ_2 are the phases accumulated by the magnetization vector during the precession under recoupling conditions in the dephasing and rephasing periods, respectively. The sum of the two signals (COS and SIN components) is proportional to $\cos(\Phi_1 - \Phi_2)$. The classical CODEX experiment was

- 30 designed for observing the reorientation of the CSA-tensor: the REDOR-like (Gullion and Schaefer, 1989) train of rotorsynchronised recoupling π-pulses applied with half rotor period spacing on the X-nuclei reintroduce the CSA interaction and thus, the phases Φ₁ and Φ₂ are determined by the precession under the influence of the CSA interaction during the de(re)phasing periods. Potentially interfering dipolar interactions with protons are supposed to be averaged out by proton decoupling during the de(re)phasing periods. However, CODEX can be easily modified for observing motionally modulated 35 dipolar interaction or isotropic chemical shift (i.e. chemical exchange). This can be achieved by a corresponding
- modification of the recoupling pulses (Krushelnitsky et al., 2013; Reichert and Krushelnitsky, 2018).



40 Figure 1. A simplified scheme of the CODEX pulse sequence. Black vertical bars denote $\pi/2$ -pulses, ϕ_{CP} , ϕ_1 , ϕ_2 and ϕ_{Rec} are the phases of the X-channel cross-polarisation pulse, two $\pi/2$ -pulses and the receiver, respectively. The COS-component is recorded when the phase differences are $\phi_{CP}-\phi_1=\pi/2$ and $\phi_2-\phi_{Rec}=\pi/2$, the SIN-component corresponds to $\phi_{CP}=\phi_1$ and $\phi_2=\phi_{Rec}$.

In the CODEX experiment, one can measure the signal intensity as a function of both mixing time and the length of the de(re)phasing periods NT_R (T_R is the MAS period and N is the number of rotor cycles in the de(re)phasing periods), which provides the information on both time scale and geometry of a molecular motion (Luz et al., 2002). Thus, the CODEX experiment enables obtaining more abundant quantitative information on molecular dynamics in comparison to standard NMR relaxation studies. At the same time, CODEX is prone to some interfering effects that may distort the information on molecular dynamics and that should be taken into account in the data analysis. Two most important effects are the proton-driven spin diffusion between X-nuclei and RIDER (Relaxation Induced Dipolar Exchange with Recoupling) (Saalwächter

50 and Schmidt-Rohr, 2000). Spin-diffusion reveals itself as a signal decay in the mixing time dependence, which can be in some cases erroneously attributed to a molecular motion process. Suppressing the spin-diffusion by proton decoupling during the mixing time is in principle possible, but rather difficult and not always reliable and effective (Reichert and

Krushelnitsky, 2018). The most robust way of removing the undesirable spin-diffusion effect is a spin dilution, e.g. using natural abundance 13 C or perdeuterated samples.

- S5 RIDER also leads to an additional decay in the mixing time dependence. Dipolar interaction of X-nuclei (*S*) with either protons or some other magnetic nuclei present in a sample (*I*), adds two terms of the precessing X-nuclei magnetization - inphase $S_x cos(\omega t)$ and anti-phase $2S_y I_z sin(\omega t)$. The last term is the origin of RIDER, which can be simplistically explained as follows: flips of I_z during the mixing time change the sign of the inter-nuclear dipolar interaction (for 1/2-nuclei) and thus change the sign of the dipolar contribution to the precession frequency. This in turn leads to incomplete rephasing of the *S*-
- 60 magnetization at the end of the rephasing period and thus to decrease of the signal. Therefore, the characteristic time of the decay in the mixing time dependence due to RIDER is determined by the timescale of I_z flips, that is, T_1 -relaxation of nuclei *I*. In addition, if the homonuclear dipolar interaction between *I*-spins is significant, spin-diffusion (flip-flops) also contributes to the time scale of RIDER, which can be much shorter than T_1 of *I*-spins. The standard way of suppressing RIDER in the CODEX experiments is heteronuclear *I-S* decoupling during the de(re)phasing periods. For some *I*-nuclei with a large quadrupolar moment, e.g.¹⁴N, decoupling is not effective, and in this case, the only way of removing the undesirable RIDER
- influence is isotopic editing of a sample.

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Our interest in the methodological problems of the CODEX experiments was stimulated by the study of slow motions in solid proteins. Recently, it was shown by means of $R_{1\rho}$ relaxometry that proteins in a solid state undergo slow overall rocking motion (Ma et al., 2015; Lamley et al., 2015; Kurauskas et al., 2017; Krushelnitsky et al., 2018). The time scale of this motion is tens of microseconds, which is the limit of the time window accessible with $R_{1\rho}$ relaxation experiments. What happens in the (sub)millisecond time scale up to now remained unclear and the CODEX experiments could answer the question, whether the rocking motion extends to longer correlation times or not.

We have thus conducted CSA and dipolar CODEX experiments on ¹⁵N nuclei in ¹⁵N,²H-enriched microcrystalline proteins (SH3 and GB1) with a partial back-exchange of labile protons. These experiments were conducted with a site-75 specific resolution in 2D ¹H-¹⁵N correlation spectrum using indirect proton detection of a signal (Chevelkov et al., 2006; Krushelnitsky et al., 2009). Surprisingly, all peaks in 2D spectra without exception reveal decays in the mixing time dependencies as shown in Fig. 2. The amplitude of the decay and the apparent correlation time of the fast component (around 20 ms) are the same for all residues. This component cannot be due to the proton driven spin diffusion since the time scale of the spin diffusion between ¹⁵N nuclei even in fully protonated proteins is much longer (Krushelnitsky et al., 2006). In the

- 80 CSA CODEX, this could be the RIDER-effect arising due to the dipolar interaction between ¹⁵N and abundant ²H nuclei. On the other hand, in the dipolar CODEX experiment, we observe very similar shapes of the mixing time dependencies with very similar parameters of the fast component. This was observed both for SH3 and GB1 microcrystalline proteins. In the dipolar CODEX experiment, the recoupling π -pulses are applied on the proton channel and thus, the ¹⁵N-²H dipolar interaction should be effectively averaged out by MAS. From this one could conclude, that the observed fast component of
- the mixing time dependencies is not an artefact and does report on a real overall protein motion in the millisecond time scale.

This would mean that the rocking motion of a protein in a crystal has a very wide correlation times distribution, from microto milliseconds.

However, it turned out that the fast decaying component in the mixing time dependencies is actually a highly non-trivial artefact. Its nature proved to be more complicated than the simple ¹⁵N-²H RIDER-effect. Below we explain the details of the 90 effects responsible for the appearance of this component and suggest some measures for correct conducting CODEX experiments and avoiding misinterpretations of CODEX data in proteins as well as other samples having complex isotopic composition.



Figure 2. Results of the residue-resolved dipolar and CSA CODEX experiments in the SH3 protein microcrystalline sample at ambient temperature, MAS 20 kHz, NT_R=2 ms. The mixing times dependences were measured for each resolved peak of the 2D ¹⁵N-¹H correlation spectrum. On the top, four examples of the mixing time decays (dipolar CODEX) of backbone ¹⁵N's are shown for the residues A11, T32, E45 and Y57. Red solid lines are the fits to the simple equation $I(\tau_m)=I(0)^{-}[(1-S_{\infty})exp(-\tau_m/\tau_c)+S_{\infty}]$, where τ_c is the apparent correlation time and S_{∞} is the decay plateau at long τ_m . At the bottom, τ_c and S_{∞} for dipolar and CSA CODEX mixing time decays are shown as a function of the residue number.

100 **2 Theory**

Here we consider the time evolution of spin coherences in the CSA CODEX experiments using product operator formalism. It is well known that after $I \rightarrow S$ cross-polarization, both in-phase S_x and anti-phase $-2S_yI_z$ terms appear, see e.g.

Schmidt-Rohr and Spiess, 1994. The anti-phase term is usually neglected since in standard CP/MAS experiments it is suppressed by the heteronuclear proton decoupling during the FID acquisition. In the CSA CODEX, it is supposed to be

- 105 suppressed by the proton decoupling during the de(re)phasing periods as well. However, in the CODEX pulse sequence this suppression is much less effective. The train of the X-channel recoupling π -pulses applied during the de(re)phasing periods restores not only CSA, but also dipolar X-¹H interaction. Hence, the proton decoupling affects not just the residual (after MAS) dipolar interaction, but the restored value of this interaction. For this reason, the small but appreciable dipolar X-¹H interaction survives during the de(re)phasing periods, which will be demonstrated experimentally below, and we have to take
- 110 it into account in our analysis.

Let us consider the time evolution of the in-phase and anti-phase terms in the CSA CODEX experiment under the simultaneous influence of the CSA and (not completely suppressed) dipolar interactions during the de(re)phasing periods. The phases acquired during the dephasing period under the influence of the CSA and dipolar interactions we denote as Φ_{CSA} and Φ_D , respectively. We assume for simplicity that Φ_{CSA} remains the same for both dephasing and rephasing periods, but

115 Φ_D can change due to RIDER. Thus, for the rephasing period, the acquired phase will be denoted as $\Phi_D + \Delta \Phi_D$.

In-phase term, dephasing period:

$$S_x \xrightarrow{CSA+DD} S_x \cos(\Phi_{\text{CSA}}) \cos(\Phi_{\text{D}}) - 2S_x I_z \sin(\Phi_{\text{CSA}}) \sin(\Phi_{\text{D}}) + S_y \sin(\Phi_{\text{CSA}}) \cos(\Phi_{\text{D}}) + 2S_y I_z \cos(\Phi_{\text{CSA}}) \sin(\Phi_{\text{D}})$$
(1)

The first two terms are picked up in the COS-component and two second terms in the SIN-component of the CODEX signal. At the end of the rephasing period, we have the COS-component:

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$$S_{x} \cos(\Phi_{CSA}) \cos(\Phi_{D}) - 2S_{x}I_{z} \sin(\Phi_{CSA}) \sin(\Phi_{D})$$
$$\xrightarrow{CSA+DD} S_{x} (\cos^{2}(\Phi_{CSA}) \cos(\Phi_{D}) \cos(\Phi_{D} + \Delta\Phi_{D}) + \sin^{2}(\Phi_{CSA}) \sin(\Phi_{D}) \sin(\Phi_{D} + \Delta\Phi_{D}))$$
(2)
and the SIN-component:

$$S_{y}\sin(\Phi_{\rm CSA})\cos(\Phi_{\rm D}) + 2S_{y}I_{z}\cos(\Phi_{\rm CSA})\sin(\Phi_{\rm D}) \xrightarrow{CSA+DD} S_{y}(\sin(\Phi_{\rm CSA})\cos(\Phi_{\rm CSA})\cos(\Phi_{\rm D})\cos(\Phi_{\rm D} + \Delta\Phi_{\rm D}) - \sin(\Phi_{\rm CSA})\cos(\Phi_{\rm CSA})\sin(\Phi_{\rm D})\sin(\Phi_{\rm D} + \Delta\Phi_{\rm D}))$$
(3)

In Eqs. (2) and (3) we left only observable terms that correspond only to COS and SIN components, respectively. Because of the proton decoupling, the phases Φ_D and $\Phi_D + \Delta \Phi_D$ are rather small. Thus, we can reasonably assume that

$$\sin(\Phi_{\rm D})\sin(\Phi_{\rm D} + \Delta\Phi_{\rm D}) \ll \cos(\Phi_{\rm D})\cos(\Phi_{\rm D} + \Delta\Phi_{\rm D}) \tag{4}$$

and
$$\cos(\Phi_{\rm D}) = \cos(\Phi_{\rm D} + \Delta \Phi_{\rm D})$$
 for spin I=1/2, since $\Delta \Phi_{\rm D}$ can be either 0 or -2 $\Phi_{\rm D}$. (5)

which means that for the in-phase term, the effect of the incomplete suppression of the dipolar X-¹H interaction is almost 130 negligible, it leads only to a small decrease of the signal, proportional to $\cos^2(\Phi_D)$.

Let us now consider the time evolution of the anti-phase term. At the end of the dephasing period we have:

$$-2S_y I_z \xrightarrow{CSA+DD} S_x \cos(\Phi_{CSA}) \sin(\Phi_D) + 2S_x I_z \sin(\Phi_{CSA}) \cos(\Phi_D) + S_y \sin(\Phi_{CSA}) \sin(\Phi_D) - 2S_y I_z \cos(\Phi_{CSA}) \cos(\Phi_D)$$
(6)

Analogously to Eq. (1), the first two terms in Eq.(5) correspond to the COS-component and the second two terms to the SIN-component. After the rephasing period, the COS-component reads:

$$S_{x} \cos(\Phi_{\text{CSA}}) \sin(\Phi_{\text{D}}) + 2S_{x}I_{z} \sin(\Phi_{\text{CSA}}) \cos(\Phi_{\text{D}})$$

$$\xrightarrow{CSA+DD} S_{x} \{\cos^{2}(\Phi_{\text{CSA}}) \sin(\Phi_{\text{D}}) \cos(\Phi_{\text{D}} + \Delta\Phi_{\text{D}}) - \sin^{2}(\Phi_{\text{CSA}}) \cos(\Phi_{\text{D}}) \sin(\Phi_{\text{D}} + \Delta\Phi_{\text{D}})\}$$
and the SIN-component is:
$$(7)$$

$$S_{y} \sin(\Phi_{\text{CSA}}) \sin(\Phi_{\text{D}}) - 2S_{y}I_{z} \cos(\Phi_{\text{CSA}}) \cos(\Phi_{\text{D}}) \xrightarrow{CSA+DD} S_{y} \cos(\Phi_{\text{CSA}}) \sin(\Phi_{\text{CSA}}) \{\sin(\Phi_{\text{D}}) \cos(\Phi_{\text{D}} + \Delta\Phi_{\text{D}}) + 140 \cos(\Phi_{\text{D}}) \sin(\Phi_{\text{D}} + \Delta\Phi_{\text{D}})\}.$$
(8)

Again, in Eqs. (7) and (8) only the observable terms are left that correspond to the COS (Eq. 7) and the SIN (Eq. 8) components. It is seen from these equations that for the anti-phase term, the RIDER effect is not negligible and the inequality analogous to Eq. (4) cannot be written if Φ_D is small but appreciable.

- But how can the RIDER effect arising from the anti-phase term be recognized in the analysis of experimental data? This 145 is relatively simple: one may compare the shapes of the mixing time dependence of the COS and SIN components. If these curves, namely the ratio S_{α}/S_0 (S_0 and S_{α} are the signal amplitudes at very short and very long mixing times, respectively), are not similar, than RIDER is relevant. In general, the ratio S_{α}/S_0 for the COS and SIN components should be exactly the same, if only molecular motions and/or spin-diffusion are present in a sample. This can be proved as follows. Let us denote the phases acquired during the dephasing and rephasing periods as Φ and $\Phi+\Delta\Phi$, respectively. At short mixing times, $\Delta\Phi=0$,
- 150 then the ratio S_{∞}/S_0 for different cases would be as follows.

Classical CODEX (COS+SIN components):

$$\frac{S_{\infty}}{S_0} = \langle \cos(\Phi)\cos(\Phi + \Delta\Phi) + \sin(\Phi)\sin(\Phi + \Delta\Phi) \rangle = \langle \cos(\Delta\Phi) \rangle$$
(9)

(10)

COS component:

$$\frac{S_{\infty}}{S_0} = \frac{\langle \cos(\Phi)\cos(\Phi + \Delta\Phi) \rangle}{\cos^2(\Phi)} = \frac{\langle \cos(\Phi)(\cos(\Phi)\cos(\Delta\Phi) - \sin(\Phi)\sin(\Delta\Phi)) \rangle}{\cos^2(\Phi)} = \langle \cos(\Delta\Phi) \rangle - \frac{\sin(\Phi)}{\cos^2(\Phi)} \langle \sin(\Delta\Phi) \rangle$$

155 SIN component:

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$$\frac{S_{\infty}}{S_0} = \frac{\langle \sin(\Phi)\sin(\Phi + \Delta\Phi) \rangle}{\sin^2(\Phi)} = \frac{\langle \sin(\Phi)(\sin(\Phi)\cos(\Delta\Phi) + \cos(\Phi)\sin(\Delta\Phi)) \rangle}{\sin^2(\Phi)} = \langle \cos(\Delta\Phi) \rangle + \frac{\cos(\Phi)}{\sin^2(\Phi)} \langle \sin(\Delta\Phi) \rangle$$
(11)

Next, we have to recall that $\Delta \Phi_{ij} = -\Delta \Phi_{ji}$ (*i* and *j* are the numbers of the exchanging sites) and since it is always assumed that we are dealing with dynamic equilibrium (i.e. the populations of the exchanging sites are constant in time), then obviously $\langle sin(\Delta \Phi) \rangle = 0$. Thus, in all cases $\frac{s_{\infty}}{s_0} = \langle cos(\Delta \Phi) \rangle$, that is the shapes of the COS and SIN components should be the same, although the absolute amplitudes in general case are of course different. Now, let us estimate the ratio S_{∞}/S_0 for the COS and SIN components described in Eqs.(7) and (8) taking into account Eq.(5) and the equation $\langle sin(\Delta \Phi_D) \rangle = \langle sin(\Phi_D + \Delta \Phi_D) \rangle = 0$ (note that this is valid only for I=1/2). COS-component: $\frac{S_{\infty}}{S_0} = \frac{\cos^2(\Phi_{CSA})\sin(\Phi_D)\cos(\Phi_D) - \sin^2(\Phi_{CSA})\cos(\Phi_D) \langle sin(\Phi_D + \Delta \Phi_D) \rangle}{\sin(\Phi_D)\cos(\Phi_D)(\cos^2(\Phi_{CSA}) - \sin^2(\Phi_{CSA}))} = \frac{\cos^2(\Phi_{CSA})}{\cos^2(\Phi_{CSA}) - \sin^2(\Phi_{CSA})}$

165 SIN-component:

$$\frac{S_{\infty}}{S_0} = \frac{\sin(\Phi_{\text{CSA}})\cos(\Phi_{\text{CSA}})(\sin(\Phi_{\text{D}})\cos(\Phi_{\text{D}}) + \cos(\Phi_{\text{D}})(\sin(\Phi_{\text{D}} + \Delta\Phi_{\text{D}})))}{2\sin(\Phi_{\text{D}})\cos(\Phi_{\text{D}})\sin(\Phi_{\text{CSA}})\cos(\Phi_{\text{CSA}})} = \frac{1}{2}$$
(13)

(12)

Hence, it is clear that the RIDER effect leads to different shapes of the mixing time dependence of the COS and SIN components. Note that if Φ_D is not small, the ratio S_{∞}/S_0 would be different for the COS and SIN components also for the inphase term, see Eqs. (2) and (3). From the Eqs. (2), (3), (7) and (8) it can also be deduced that the SIN-component is about twice more prone to the RIDER-distortions than the COS-component. This follows from the comparison of the amplitudes of different coherences. The amplitudes of the COS-component of the in-phase and anti-phase terms, see Eqs. (2) and (7), are proportional to $\cos^2(\Phi_{CSA}) \cdot \cos^2(\Phi_D)$ and $\cos^2(\Phi_{CSA}) \cdot \cos(\Phi_D) \cdot \sin(\Phi_D)$, respectively (here we assume Φ_{CSA} to be not large). Hence, the ratio of the anti-phase-term amplitude to the in-phase-term amplitude is proportional to $\tan(\Phi_D)$ or even smaller if the second terms in the parentheses in Eqs. (2) and (7) are taken into account. The same ratio for the SIN-component, see 175 Fas. (3) and (9) is proportional to $2\tan(\Phi_D)$. Thus, the contribution of the anti-phase coherence to the total signal is larger.

175 Eqs. (3) and (8), is proportional to $2 \cdot \tan(\Phi_D)$. Thus, the contribution of the anti-phase coherence to the total signal is larger for the SIN-component.

The analysis presented above is valid only for an isolated *I-S* spin pair. For multinuclear spin systems, the description would be much more complicated since many types of multiple coherences with a complex network of homo- and heteronuclear dipolar interactions should be taken into account. Quantifying this is outside the frames of our work, still we believe

180 that on a qualitative level, two most important points remain valid: first, the anti-phase term appearing after the crosspolarization pulses may cause RIDER distortions of the mixing time dependencies and second, the RIDER effect can be recognized from the comparison of the shapes of the COS and SIN components. This will be proven experimentally below.

3 Materials and methods

3.1 Samples

In our work we used four different samples. Model substances: ¹⁵N enriched BOC-Glycine and ¹⁵N enriched Glycine, which were purchased from Sigma-Aldrich. Proteins: small GB1 and SH3 proteins in a form of microcrystals, ¹⁵N, ²H enriched with a partial back-exchange of labile protons. The GB1 sample was purchased from Giotto Biotech (Florence, Italy), the SH3 sample was prepared in Prof. B. Reif's lab (FMP, Berlin). These are the same samples that were used in our

recent work on R_{10} relaxometry (Krushelnitsky et al., 2018). Both protein samples were prepared according to the protocol ensuring 20% of the back-exchange of labile protons. However, we believe that in reality this percentage is somewhat 190 different: in GB1 it is higher which is indicated by stronger signal and faster proton-driven spin-diffusion between ¹⁵N nuclei (see Figs. 12 and 13 below). The quantitative estimation of this difference is yet rather difficult and uncertain. Since the GB1 sample provides better signal-to-noise ratio, most of the experiments were conducted with this sample.

3.2 NMR experiments

- 195 The experiments were performed on a Bruker AVANCE II NMR spectrometer (600 MHz) with a 3.2 mm MAS probe. In the CODEX experiments with the protein samples, the integral intensity of the entire signal was determined without siteselective specification (except for the data shown in Fig. 2). One-dimensional double cross-polarisation $({}^{1}H \rightarrow {}^{15}N \rightarrow {}^{1}H)$ proton-detected spectra for SH3 and GB1 proteins were shown in Krushelnitsky et al., 2018. For the BOC-Gly and Gly samples the direct ¹⁵N or ¹³C signal detection was employed, and for the protein samples we used indirect ¹H signal detection of the ¹⁵N CODEX mixing time dependencies. This was implemented using back cross-polarisation section (${}^{15}N \rightarrow {}^{1}H$) at the 200 end of the pulse sequence, according to the approach described earlier (Chevelkov et al., 2006; Krushelnitsky et al., 2009). We have checked - the direct ¹⁵N and indirect ¹H signal detections in the protein samples provide the same shape of the CODEX mixing time dependencies, in the latter case the signal-to-noise ratio was however better.
- To exclude the effect of spin-lattice relaxation during the mixing time, each CODEX mixing time dependence was T_1 normalized. For that, for each mixing time dependence two experiments were performed: measuring the mixing time 205 dependence itself and measuring a T_1 -relaxation curve within the same time range. After that, the mixing time dependence was divided by T_1 -relaxation curve. This is a routine procedure described earlier (deAzevedo et al., 1999; deAzevedo et al., 2000; Reichert et al., 2001; Reichert and Krushelnitsky, 2018). Below are shown only the T_1 -normalized mixing time dependencies for all CODEX experiments in protein samples. For BOC-Gly, the T_1 -normalization was not performed since 210 15 N T_1 in this sample was extremely long (800-900 s).

The pulse sequences of the CSA and dipolar CODEX are shown in Figs. 3 and 4. For measuring the mixing time dependence, τ_m was variable and τ_r was fixed at 1 ms; for measuring T_1 -relaxation curve, τ_m was fixed at 1 ms and τ_r was variable. The phase cycle for both COS and SIN components consists of 64 steps: 2x spin-temperature inversion (ensuring that the signal decays to zero (Torchia, 1978)) for T_1 -relaxation, 2x spin-temperature inversion for mixing time, 4x CYCLOPS for the $\pi/2$ -pulse after mixing time, 4x CYCLOPS for the $\pi/2$ -pulse after τ_r delay (Reichert et al., 2001). Typical values for $\pi/2$ pulse for ¹H and ¹⁵N channels were 1.4-1.8 µs and 6.0-6.5 µs, respectively.

The experimental error in estimation the signal amplitude was: less than 1% for BOC-Gly, 1-2% for GB1, 2-4% for SH3 and 5-10% for natural abundance ¹³C in Gly. On top of the signal noise, a certain contribution to the experimental error in the mixing time dependences comes from the instability of the MAS controller; that was however significant only for BOC-Gly. The final error of the mixing time dependencies for this sample was around 1-2%. For better visual distinguishing

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between the mixing time dependencies shown in Figs. 5, 7 and 8, the adjacent averaging over 5 points filter was applied to the experimental curves in these figures, which significantly decreased the noise spread of the points without the change of the overall shape of the curves.



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Figure 3. CSA CODEX pulse sequence for the direct (¹³C or ¹⁵N, top) and indirect (¹H, bottom) signal detection. Solid and open bars denote $\pi/2$ and π pulses, respectively. The mixing time τ_m is an integer multiple of the MAS period, which is achieved by MAS rotor triggering before and at the end of the mixing time (see details in Reichert and Krushelnitsky, 2018). Rotor synchronization during the τ_r -delay is not necessary. Waltz decoupling in the indirect detection sequence aims to suppress only J-coupling between

230 X and ¹H nuclei, therefore it has low amplitude (few hundred Hz). An additional initial Z-filter and ²H-decoupling (see below) are not shown.

Phase cycle:

 $\phi_1 = x; \ \phi_2 = y; \ \phi_3 = x; \ \phi_4 = x; \ \phi_5 = (y, -y) \ (\text{COS component}); \ \phi_5 = (x, -x) \ (\text{SIN component}); \\ \phi_6 = (x, x, y, y, -x, -x, -y, -y); \ \phi_7 = (y, -y, -x, x, -y, y, x, -x); \\ 235 \ \phi_8 = (-x, -x, -y, -y, x, x, y, y, x, x, y, y, -x, -x, -y, -y) \ (\text{COS component}); \\ \phi_8 = (y, y, -x, -x, -y, -y, x, x, y, y, -x, -x, -y, -y) \ (\text{COS component}); \\ \phi_8 = (y, y, -x, -x, -y, -y, x, x, -y, -y, x, x, y, y, -x, -x) \ (\text{SIN component}); \\ \phi_9 = (x^*16, y^*16, -x^*16, -y^*16); \ \phi_{10} = (y^*16, -x^*16, -y^*16, x^*16); \ \phi_{11} = (x, x, y, y, -x, -x, -y, -y); \\ \phi_{\text{Rec}} = ((y, -y)^*4, \ (-y, y)^*4, \ (-x, x)^*4, \ (-x, x)^*4, \ (-y, y)^*4, \ (x, -x)^*4, \ (-x, x)^*4) \ (\text{direct detection}); \\ \end{cases}$

 $\phi_{\text{Rec}} = (x, -x, y, -y, -x, x, -y, y, -x, x, -y, y, x, -x, y, -y)$ (indirect detection).



Figure 4. Dipolar CODEX pulse sequence for the direct (top) and indirect (bottom) signal detection. The denotations are the same as in Fig. 3. π -pulses applied on X-channel are set in the middle of the de(re)phasing periods, therefore the duration of these periods should be an even multiple of the MAS period. The phase cycle is identical to that shown in Fig. 3. The phases of the π -pulses applied during the de(re)phasing periods on ¹H-channel have no critical significance.

4 Results and Discussion

4.1 CSA CODEX

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4.1.1 Rigid model substances

¹⁵N-enriched BOC-Glycine is a rigid solid sample in which we do not expect any molecular motion on the millisecond

- 250 time scale. Thus, the CODEX mixing time decays can be only due to the RIDER effect since the proton-driven spindiffusion between ¹⁵N nuclei in BOC-Gly is very slow (Krushelnitsky et al., 2006). First, we demonstrate that the anti-phase term discussed above does really cause RIDER distortions in the CSA CODEX mixing time dependence. The anti-phase term appears in the course of cross-polarization; thus, its contribution to the total CODEX signal should depend on the CP contact time. The CSA CODEX mixing time dependencies at various CP times are shown in Fig. 5. These data fully confirm
- 255 the qualitative theoretical analysis presented above. One may see that the amplitude of the RIDER decay depends on the CP time, that COS and SIN components of the mixing time dependencies are different and that the SIN component is more prone to the RIDER distortions than the COS component. It is interesting to mention that the mixing time dependencies shown in Fig. 5 reveal the decays on two different time scales: a few milliseconds and a few hundred milliseconds. Such a two-component shape of the decays reflects two different mechanisms that cause proton spin flips mentioned in the
- 260 Introduction above: spin diffusion (flip-flops) and spin-lattice relaxation.



Figure 5. COS and SIN components of the ¹⁵N CSA CODEX mixing time dependence measured at various cross-polarisation contact times in BOC-Gly. The initial amplitudes of the τ_m -dependencies were normalized to the same value. MAS 20 kHz, NT_R 2 ms, 108 kHz ¹H CW-decoupling during the de(re)phasing periods.

If the heteronuclear proton decoupling during the de(re)phasing periods was effective enough, than the RIDER effect caused by the anti-phase coherence could have been of course avoided. However, this is not always possible for practical reasons because of the hardware limitations for the power of the long proton pulses. We tried to optimize the proton decoupling by the maximum signal at short mixing times. Different decoupling schemes were checked (TPPI, WALZ, SPINAL) at maximum proton power around 100-130 kHz, but none of them provided much better efficiency than simple CW decoupling (which is not the case for ¹H-decoupling during FID, where CW is not the best choice). Therefore, in the experiments shown here we used CW ¹H decoupling during the de(re)phasing periods in the CSA CODEX measurements. We do not claim that CW decoupling is the best option for this purpose. It is quite possible that some other decoupling schemes apecifically designed for the case of the recoupling X-pulses can perform better. However, even having such a decoupling scheme at hand, one should carefully optimize it for different MAS rates and ¹H field strengths. We suggest here another, more simple and robust way of suppressing the undesired RIDER effect.

The anti-phase term can simply be suppressed by an additional Z-filter between the CP pulses and the dephasing period, as illustrated in Fig. 6. The delay of this Z-filter should be short compared to 15 N T_1 and long compared to T_2 . Thus, after such Z-filter one would have only in-phase component. Fig. 7 shows the mixing time dependencies of the COS and SIN components at different delays of the Z-filter. It is clearly seen that the Z-filter fully removes the contribution of the antiphase coherence.



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Figure 6. Initial part of the CODEX pulse sequence (Figs. 1 and 3) with the additional Z-filter between the cross-polarization section and the dephasing period. ϕ_{CP} - ϕ_{Z1} =± $\pi/2$, ϕ_{Z2} =- ϕ_{Z1} .

Still it is seen that even at long delays of the Z-filter, the mixing time dependencies are not completely flat, as they 290 should be. The observed distortions are obviously the RIDER effect of the in-phase coherence. The Z-filter eliminates the anti-phase coherence (Eqs. 7 and 8), but it does not improve the efficiency of the proton decoupling during the de(re)phasing periods and thus the phase Φ_D remains non-zero. If the second terms in the parentheses in Eqs. (2) and (3) are not negligibly small in comparison to the first terms, then the RIDER is present also in the in-phase coherence and the Z-filter obviously cannot remove it. Fig. 8 presents the COS and SIN components of the mixing times dependencies at different durations of 295 the de(re)phasing periods measured with the additional Z-filter. It is clearly seen: the longer NT_R , the larger the RIDER distortions. This is reasonable since Φ_D is proportional to NT_R. Note that the COS component is less prone to distortions not only in the case of the "anti-phase", but also of the "in-phase" RIDER. We are not able at the moment to explain the unusual bell-shaped form of the mixing time dependencies. It is likely that the network of multi-nuclear dipolar interactions should be taken into account and thus the explanation will not be simple. We also cannot exclude that transient NOE effects may 300

play a certain role.

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However, in any case this is the unwanted distortion and regardless of the exact nature of this distortion it should be maximally suppressed in real experiments. For this, the efficiency of the ¹H-decoupling during the de(re)phasing periods must be optimized as far as possible. As mentioned above, the standard heteronuclear decoupling schemes used for FID detection do not help much for the case of de(re)phasing periods. This is illustrated by the example of SPINAL sequence, see Fig. 8. Still one may minimize the "in-phase" RIDER effect by keeping NT_R as short as possible and by recording only the COS component of the mixing time dependence. Anyway, the "in-phase" RIDER is much smaller than the "anti-phase" one and in most real experiments it can be safely neglected, as we will see below by the example of the protein samples.

At the end of this section, we demonstrate the application of the additional Z-filter to the natural abundance ¹³C CSA CODEX experiment performed on carbonyl carbons in ¹⁵N-enriched glycine (¹⁵N enrichment is necessary to avoid the ¹³C-³¹⁰ ¹⁴N RIDER effect). We see the same effect as in the case of ¹⁵N CSA CODEX (Fig. 9). The dependencies measured with the Z-filter (red points in Fig. 9) are not completely flat, however, this is hardly due to the "in-phase" RIDER since the shapes of the SIN and COS components are very similar (in the case of RIDER they should be different) and the time constant of the decay (about 50-60 s) is obviously too long compared to the proton T₁ (few seconds). We suspect this decay is a manifestation of the proton-driven spin diffusion between natural abundance ¹³C nuclei. Its time constant is roughly of the

315 same order of magnitude as spin diffusion times between natural abundance ¹³C nuclei measured in other organic solids, see e.g. Reichert et al., 1998. Spin diffusion however has no direct relevance to the topic of this work and we did not analyze this in detail.

In summary, the theoretical and experimental results presented above show that the proton decoupling under the influence of the recoupling π -pulses in the CSA CODEX is not fully efficient. This leads to the evolution of both in-phase and anti-phase coherences during the de(re)phasing periods under the influence of the residual ¹⁵N(¹³C)-¹H dipolar coupling, that is, to the RIDER effect. The dominant contribution to the RIDER distortions of a mixing time dependence arises from the anti-phase term. This contribution can be suppressed by the additional Z-filter between CP and dephasing period. The RIDER distortion of the in-phase term is smaller but still appreciable at long de(re)phasing periods. This interfering effect cannot be suppressed completely, but it can be significantly minimized if only the COS-component of the mixing time

325 dependence is measured and analyzed since this component is less prone to RIDER in comparison to the SIN-component.



Figure 7. COS and SIN components of the ¹⁵N CSA CODEX mixing time dependence in BOC-Gly at different Z-filter delays. All the dependencies were normalized to the same initial amplitude. MAS 20 kHz, 108 kHz CW ¹H decoupling during the de(re)phasing periods, NT_R 2 ms, CP contact time 3 ms.



Figure 8. Normalized SIN and COS components of the ¹⁵N CSA CODEX mixing time dependence in BOC-Gly at different NT_R. MAS 20 kHz, 108 kHz CW or SPINAL ¹H decoupling during the de(re)phasing periods, Z-filter delay 100 ms, CP contact time 3 ms.



Figure 9. ¹³C (carbonyls, nat.abundance) CSA CODEX mixing time dependencies in ¹⁵N-enriched Glycine measured with and without additional Z-filter (Fig. 6). All decays were normalized to the same initial amplitude, the real ratio between the amplitudes of SIN and COS components is 0.7 for both experiments. 80 kHz CW ¹H decoupling and 35 kHz CW ¹⁵N decoupling during the de(re)phasing periods were applied. MAS 22 kHz, NT_R 2 ms, Z-filter delay 20 ms, CP contact time 3 ms.

4.1.2 Protein samples

In the protein samples we have three types of nuclei that we need to take into account - ¹⁵N, ¹H and ²H. The direct and inverse ¹H-¹⁵N cross-polarisation sections employed in the CODEX pulse sequence ensure that in the experiment we observe

345 only those nitrogens that have protons attached, and all ¹⁵N's coupled to ²H in the protein backbone remain invisible. Still,

the interactions between protonated ¹⁵N's and many remote ²H's can be sufficient to induce RIDER-type distortions in the CODEX experiment. To demonstrate the hierarchy of the inter-nuclear interactions in our samples, we measured ¹⁵N Hahn-echo decays (Fig. 10) and the initial signal S_0 (the signal at short mixing time) in the CSA and dipolar CODEX experiments

as a function of NT_R (Fig. 11) for various combinations of ¹H and ²H decoupling schemes. Note that the S₀ vs NT_R
dependence is in fact the analogue of the Hahn-echo experiment, the only difference is that either CSA or dipolar interaction is reintroduced by means of recoupling pulses during the transverse relaxation.

The conclusions that can be deduced from these data are as follows. First, despite the proton dilution, the ¹⁵N-¹H dipolar line broadening at the MAS frequency 20 kHz remains quite appreciable and strong proton decoupling is necessary to suppress the ¹⁵N-¹H dipolar interaction. The comparison of the S_0 vs NT_R dependences of dipolar CODEX in fully protonated BOC-Glycine and the deuterated protein shows that the proton dilution reduces of course the inter-proton interaction (flip-flops) and thus, the rate of the ¹⁵N decay: slower ¹H-flips ensure slower ¹⁵N-¹H coupling modulation and hence, better refocusing the signal after the end of the rephasing period. Still the rate of the proton flip-flops in the protein sample remains in the millisecond range. This is an important point, which will be discussed below. This result corresponds

360 Second, it is clearly seen that the 130 kHz CW-decoupling performs much worse in comparison to the SPINAL scheme (Fig. 10). As mentioned above, under the influence of the ¹⁵N recoupling π -pulses during the de(re)phasing periods, SPINAL does not provide significant advantage in comparison to CW. This confirms our previous statement that the proton decoupling efficiency under the influence of the X-channel recoupling pulses is much different in comparison to FID detection.

well to the proton line width estimations made by Reif and co-workers (Chevelkov et al., 2006).

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Figure 10. ¹⁵N Hahn-echo decays measured in GB1 protein sample at different ¹H and ²H decoupling schemes. MAS 20 kHz, t 13 °C, ¹H decoupling strength (both for CW and SPINAL) 130 kHz, duration of ²H π -pulses 10.5 μ s.



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Figure 11. Signal intensity (COS component) at short mixing time (1 ms) in ¹⁵N CODEX experiments as a function of NT_R in GB1 protein and BOC-Gly samples. MAS 20 kHz, t 13 °C, ¹H and ²H CW decoupling strengths during the de(re)phasing periods 130 kHz and 45 kHz, respectively.

Third, the ¹⁵N-²H interaction is non-negligible. ²H decoupling does not lead to longer the Hahn-echo decays since it is effectively (but not completely, see below) reduced by MAS even without decoupling. However, the reintroduction of the ¹⁵N-²H dipolar coupling by the REDOR pulse train applied on deuterons appreciably shortens the decays, see Fig. 10. In the CSA CODEX experiment, the ¹⁵N-²H interaction is initially reintroduced by means of REDOR pulse train applied on ¹⁵N's. In this case, the ²H decoupling has the effect and makes the decay slower (Fig. 11).

- Now the recipe for a methodologically correct CSA CODEX experiment is evident. In deuterated proteins, there are two 380 simultaneous RIDER effects arising from the ¹⁵N-¹H and ¹⁵N-²H dipolar interactions and one has to take care of both of them. Coincidentally, both RIDERs have similar, although not exactly the same, time constants. The time constant for the proton flip-flops can be estimated directly from the Hahn-decay which gives the value of about 10 ms (Fig. 10). As for the ²H T_1 , it has a value of 25 ms for aliphatic deuterons in the SH3 protein sample, which was measured by a simple inversionrecovery method. Both these values are quite close to the time constant of the short component of the CODEX mixing time 385 dependences observed in proteins (Fig.2).
- 385 dependences observed in proteins (Fig.2).

The ¹⁵N-¹H and ¹⁵N-²H RIDER effects can be suppressed by the additional Z-filter between CP and dephasing period (see above) and the rf-decoupling, respectively. We remind that the Z-filter suppresses only the "anti-phase" ¹⁵N-¹H RIDER, but not the "in-phase" one. However, the "in-phase" RIDER distortion of the COS component at reasonably short NT_R is practically negligible, as our data demonstrate. Figs. 12 and 13 present the mixing time dependences of the CSA CODEX at various combinations of the ¹⁵N-¹H and ¹⁵N-²H suppression tools for GB1 and SH3 protein samples. It is seen that the dominant contribution to the short component in the mixing time dependence (Fig.2) comes from the ¹⁵N-²H RIDER. Still ²H-decoupling alone cannot ensure the artefact-free experiment, and only combination of Z-filter and ²H-decoupling provides the flat mixing time dependence in the millisecond time scale for both proteins. This demonstrates that both SH3 and GB1 proteins in microcrystalline form do not undergo global motions in the millisecond time scale and the overall

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395 rocking motion is limited to the microsecond range only.



Figure 12. COS component of the ¹⁵N CSA CODEX mixing time dependence in linear (right) and logarithmic (left) time scale measured in GB1 with/without Z-filter and with/without ²H CW decoupling during the de(re)phasing periods. MAS 20 kHz, t 13 °C, CP contact time 1.5 ms, NT_R 2 ms, ¹H and ²H CW decoupling strengths 130 kHz and 45 kHz, respectively.



Figure 13. The same data at the same conditions as shown in Fig. 12 for SH3.



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Figure 14. ¹⁵N CSA CODEX mixing time dependencies measured in GB1 at different MAS rates and temperatures. In all cases Z-filter and ²H CW decoupling during the de(re)phasing periods were applied (the parameters are the same as mentioned in the caption to Fig. 12). NT_R 2 ms.

- The mixing time dependences in Figs. 12 and 13 also reveal a rather slow decay with a time constant in the second range. 410 This is spin diffusion between ¹⁵N nuclei, which is easy to prove. The spin diffusion rate should not depend on temperature and should depend on the MAS rate (Reichert et al., 2001; Krushelnitsky et al., 2006). We measured the mixing time dependence for the GB1 sample at two temperatures and two MAS rates, see Fig. 14. The results shown in this figure leave no doubts that this is the ordinary proton driven spin-diffusion. The rate of these decays is approximately 3-4 times slower than the spin-diffusion rate in a fully protonated protein (Krushelnitsky et al., 2006) still it is quite appreciable. Spin
- 415 diffusion rate in SH3 protein is noticeably slower; we believe this is due to the lower proton density in this sample, which is confirmed by a somewhat weaker signal from SH3 compared to GB1.

4.2 Dipolar CODEX

The principal problem of the dipolar CODEX is that the ¹⁵N-¹H interaction cannot be decoupled for obvious reasons and thus the anti-phase term responsible for the RIDER effect emerges explicitly during the de(re)phasing periods even without CP. To solve this problem, in our first paper on dipolar CODEX (Krushelnitsky et al., 2009) we suggested to measure only the COS component of the mixing time dependence. The COS component must be RIDER-free, which directly follows from the Eq. (2). In the dipolar CODEX $\Phi_{CSA}=0$, and since $\cos(\Phi_D)=\cos(\Phi_D+\Delta\Phi_D)$ (we repeat again that this is valid only for I=1/2), the COS-component of the dipolar CODEX mixing time dependence should not be affected by the ¹⁵N-¹H RIDER. However, this is only true under the condition that we did not pay a proper attention to at that time. This condition is: the longer than NT_R. If this is not so, then $\cos(\Phi_D)\neq\cos(\Phi_D+\Delta\Phi_D)$ since Φ_D and $\Phi_D+\Delta\Phi_D$ are randomly modulated by I-spin flips within the de(re)phasing periods. Thus, the COS component at this condition is not RIDER-free anymore.

The comparability of NT_R and the time scale of proton spins filps is exactly our case. We have estimated above the characteristic time of the protons flip-flops, which is about 10 ms (Fig. 10). The duration of the de(re)phasing periods in the

- 430 CODEX experiments is usually from few hundred microseconds to several milliseconds. This is shorter than 10 ms but still comparable, which is enough for the RIDER effect. From this we pessimistically conclude that the X-H dipolar CODEX experiment even in proton-diluted samples like deuterated proteins with a partial back-exchange of labile protons is not suitable for studying slow molecular dynamics there will always be RIDER distortions. This means that the decay in the dipolar CODEX mixing time dependencies of backbone nitrogens in SH3 protein that we observed earlier (Krushelnitsky et
- 435 al., 2009) is not due to molecular motions but due to the RIDER effect and that these data were misinterpreted. The dipolar CODEX experiment, however, can be implemented using other nuclei pairs, e.g. ¹³C-¹⁵N (McDermott and Li, 2009), ensuring that the flip-flop time is much longer than the duration of the de(re)phasing periods.

The last point that deserves to be discussed here is the influence of the ¹⁵N-²H interaction on the dipolar CODEX results. At first sight, there should be no influence, since this interaction is not reintroduced in the dipolar CODEX and it should be simply suppressed by MAS. However, this is not the case. Fig. 15 presents the mixing time dependences in GB1 measured at different powers of the CW-²H-decoupling during the de(re)phasing periods. The data demonstrate that in spite of MAS, the ¹⁵N-²H interaction has a small but well visible contribution to the short component, i.e. RIDER effect, of the mixing time

dependence. The residual ¹⁵N-²H interaction is rather small since only few kHz of CW decoupling is enough to suppress it completely. So, why MAS does not do its job alone, without the rf-decoupling? The reason is the protein mobility in the

- 445 microsecond time scale. If the ¹⁵N-²H interaction is modulated by a molecular motion on a time scale of the MAS period (for 20 kHz it is 50 μs), then MAS cannot suppress this interaction completely, which leads to the increased linewidths of the MAS centerband (Suwelack et al., 1980). As we know, the correlation time of the protein rocking motion is few tens microseconds (Krushelnitsky et al., 2018). On top of that, there can be interaction of protein nitrogens with deuterons of solvent molecules, and these molecules can also reveal a mobility in the microsecond time scale. Thus, the appearance of the
- 450 residual ¹⁵N-²H interaction after MAS can be reasonably explained.



Figure 15. ¹⁵N dipolar CODEX mixing time dependencies (COS component) measured in GB1 protein sample at various ²H CW decoupling strengths during the de(re)phasing periods. Z-filter 0.1 s between the cross-polarization section and the dephasing period was applied, MAS 20 kHz, t 13 °C, NT_R 2 ms.

In summary, we have shown that both ${}^{15}N{}^{-2}H$ and ${}^{15}N{}^{-1}H$ RIDER effects contribute to the short component of the mixing time dependencies of both CSA and dipolar CODEX experiments in the protein samples. However, the dominant contributions in these two experiments are different: in the CSA CODEX the dominant source of the short component is the ${}^{15}N{}^{-2}H$ interaction, and in the dipolar CODEX it is the ${}^{15}N{}^{-1}H$ interaction. As estimated above, the time constants of the two

- 460 RIDER effects are similar but not the same: ²H spin-lattice relaxation is somewhat slower than the proton flip-flop rate. Therefore, the apparent decay rate of the short component in the CSA and dipolar CODEX experiments should also be somewhat different. This is illustrated in Fig. 16, which presents the fast RIDER-components of the CSA and dipolar CODEX experiments after subtraction of the spin-diffusion component and normalization of the decay amplitudes to the same value. The direct comparison of these decays is in a perfect agreement with the findings described above. Interesting to
- 465 note that in SH3, the difference of the apparent correlation times of the short component for the CSA and dipolar CODEX is much smaller, see Fig. 2 (τ_c as a function of the residue number). This can also be reasonably explained by the different proton density in the GB1 and SH3 samples: the lesser the proton density, the slower the flip-flop rate and thus, the smaller the difference between the rates of proton spin diffusion and deuteron spin-lattice relaxation.



470 Figure 16. Direct comparison of the CSA and dipolar CODEX data in GB1 protein sample. Left: the mixing times dependencies taken from Fig. 12 (CSA CODEX, without Z-filter and without ²H decoupling) and Fig. 15 (dipolar CODEX, no ²H decoupling). Red solid lines - the exponential fits of the initial parts of the spin-diffusion components. Right: Fast initial components of the decays after subtraction the spin-diffusion components and normalization to the same initial amplitude.

Conclusions.

475 1) The comparison of the shapes of SIN and COS components of the mixing time dependences is a simple and robust way of detecting the presence/absence of the RIDER effect in the CODEX experiments. The COS component is less prone to the RIDER distortion (appearance of the short component) and for minimising this distortion, it is advisable to record and to analyse in experiments only the COS component.

2) Proton decoupling under the influence of the recoupling π-pulses applied on X-channel is not as effective as in the case of normal X-nuclei FID detection. Thus, the suppression of the anti-phase coherence emerging after the cross-polarisation section can be incomplete in CSA CODEX. This may lead to the RIDER distortion in mixing time dependences. This problem can be effectively resolved by inserting additional Z-filter between the cross-polarization section and the dephasing period.

3) In ¹⁵N CODEX experiments in deuterated proteins with a partial back-exchange of labile protons one has to consider two different RIDER effects arising from ¹⁵N-¹H and ¹⁵N-²H dipolar interactions. CSA and dipolar CODEX are affected predominantly by ¹⁵N-²H RIDER and ¹⁵N-¹H RIDER, respectively. A combination of Z-filter and ²H-decoupling during the de(re)phasing periods enables suppression of both effects in the CSA CODEX, however for the dipolar CODEX this is not possible.

4) GB1 and SH3 proteins in their microcrystalline form do not reveal global motion in the millisecond time scale.

Author contributions

AK designed the project, conducted the experiments and the data analysis, wrote the paper; KS took part in planning the experiments and discussing the results, and edited the paper.

Competing interests

495 The authors declare no conflict of interests.

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