ARTICLE

1H R1ρ relaxation dispersion experiments in aromatic side chains

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Abstract

Aromatic side chains are attractive probes of protein dynamic, since they are often key residues in enzyme active sites and protein binding sites. Dynamic processes on microsecond to millisecond timescales can be studied by relaxation dispersion experiments that attenuate conformational exchange contributions to the transverse relaxation rate by varying the refocusing frequency of applied radio-frequency fields implemented as either CPMG pulse trains or continuous spin-lock periods. Here we present an aromatic 1H R1ρ relaxation dispersion experiment enabling studies of two to three times faster exchange processes than achievable by existing experiments for aromatic side chains. We show that site-specific isotope labeling schemes generating isolated 1H–13C spin pairs with vicinal 2H–12C moieties are necessary to avoid anomalous relaxation dispersion profiles caused by Hartmann–Hahn matching due to the 3JHH couplings and limited chemical shift differences among 1H spins in phenylalanine, tyrosine and the six-ring moiety of tryptophan. This labeling pattern is sufficient in that remote protons do not cause additional complications. We validated the approach by measuring ring-flip kinetics in the small protein GB1. The determined rate constants, kflip, agree well with previous results from 13C R1ρ relaxation dispersion experiments, and yield 1H chemical shift differences between the two sides of the ring in good agreement with values measured under slow-exchange conditions. The aromatic 1H R1ρ relaxation dispersion experiment in combination with the site-selective 1H–13C/2H–12C labeling scheme enable measurement of exchange rates up to kex = 2kflip = 80,000 s⁻¹, and serve as a useful complement to previously developed 13C-based methods.

Keywords Conformational exchange · Protein dynamics · Aromatic side chains · Rotating-frame relaxation · Aromatic ring-flip

Introduction

Conformational dynamics in proteins on the microsecond to millisecond time scales are often linked to biological function (Mittermaier and Kay 2009). Transiently populated high-energy states play important roles in enzyme catalysis (Boehr et al. 2006; Cole and Loria 2002; Eisenmesser et al. 2002) or ligand binding by conformational selection (Demers and Mittermaier 2009; Malmendal et al. 1999). Such conformational transitions generally lead to a modulation of NMR parameters as the chemical shift (Gutowsky and Saika 1953), residual dipolar coupling (Igumenova et al. 2007; Vallurupalli et al. 2007) or strong scalar coupling (Weininger et al. 2013b). The conformational dynamics can be probed by NMR relaxation dispersion methods (Palmer 2004; Palmer et al. 1991), such as R1ρ (Akke and Palmer 1996; James et al. 1977) or Carr-Purcell-Meiboom-Gill (CPMG) experiments (Carr and Purcell 1954; Loria et al. 1999a, 1999b; Meiboom and Gill 1958), and chemical exchange saturation transfer (CEST) methods (Forsen and Hoffman 1963; Palmer and Koss 2019; Vallurupalli et al. 2012, 2017) in the case of slower time scales.

Phenylalanine, tyrosine, histidine and tryptophan all have aromatic side chains, which make them an interesting subgroup of amino acids that serve multiple functions in proteins. Aromatic side chains are bulky and constitute a significant proportion of the protein hydrophobic core. They typically form pairs or clusters on the basis of specific
aromatic-aromatic interactions (Burley and Petsko 1985, 1989). Furthermore, aromatic residues are overrepresented in protein binding interfaces, where they contribute a significant part of the binding free energy (Birtalan et al. 2010; Bogan and Thorn 1998; Lo Conte et al. 1999). Finally, tyrosine and especially histidine play critical roles in enzyme catalysis, where they contribute a significant part of the binding free energy (Birtalan et al. 2010; Bogan and Thorn 1998; Lo Conte et al. 1999). Finally, tyrosine and especially histidine play critical roles in enzyme catalysis, where they make up 6% and 18%, respectively, of all catalytic residues (Bartlett et al. 2002). Histidine can exist in three different states, one protonated and two neutral tautomeric forms. Despite their generally tight packing, Phe and Tyr residues undergo intermittent 180° transitions (‘ring flips’) of the $\chi_2$ dihedral angle and thereby provide unique information of transient ‘breathing’ processes of proteins (Dreydoppel et al. 2020; Wagner 1980; Wagner et al. 1976; Weininger et al. 2014b). Each of these properties makes aromatic side chains highly interesting and powerful probes for studying protein dynamics.

Recent developments have enabled straightforward and robust site-selective $^{13}$C labeling of aromatic side chains (Kasinath et al. 2013; Lundström et al. 2007; Schörghuber et al. 2018; Teilum et al. 2006; Weininger 2019), which eliminate unwanted relaxation pathways and coherent magnetization transfer via one-bond $^{13}$C–$^{13}$C couplings. These advancements have made possible advanced studies of protein dynamics involving aromatic side chains, including methods to characterize fast (ps–ns) timescale dynamics via $^{13}$C relaxation rate constants (Weininger et al. 2012a) that enable studies of order parameters (Boyer and Lee 2008; Kasinath et al. 2013, 2015). Furthermore, slower (µs–ms) timescale dynamics have been probed with $^{13}$C relaxation dispersion experiments, either using CPMG (Weininger et al. 2012b) or $R_{1\rho}$ refocusing elements (Weininger et al. 2014a). In particular, these relaxation dispersion methods have been used to measure ring-flip rates (Dreydoppel et al. 2020; Weininger et al. 2014b) and transient histidine tautomeration (Weininger et al. 2017). Complementary $^1$H CPMG relaxation dispersion experiments are applicable to a subset of aromatic sites, namely His δ2, His ε1 and Trp δ1. Other positions are inaccessible due to sizeable $^{3}J_{HH}$ couplings and possibly strong $^{1}J_{CC}$, where $^{1}J$-couplings are equal or greater than the chemical shift difference, can cause severe artifacts (Raum et al. 2018). Site-selective $^{1}H–^{13}C/^{12}C$ labeling addresses both issues and allows to obtain artifact-free $^1$H CPMG relaxation dispersion profiles (Raum et al. 2019).

It is expected that $^1$H $R_{1\rho}$ experiments make it possible to access faster time scales, since the attainable refocusing frequency scales with the gyromagnetic ratio and the RF field amplitude, $\omega_1 = \gamma B_1$. $^1$H $R_{1\rho}$ experiments have successfully been applied to the aromatic position H8 of adenine and guanine and in RNA (Steiner et al. 2016), which is not affected by sizeable $^{3}J_{HH}$ couplings or cross relaxation with remote protons. The situation is more challenging in certain aromatic protein residues, where $^{3}J_{HH}$ is on the order of 7–8 Hz and the chemical shift difference between vicinal ring protons is often small. Here we demonstrate these problems can be solved by site-selective $^1$H–$^{13}$C/-$^{12}$C labeling, which is necessary and sufficient to ensure artifact-free $^1$H $R_{1\rho}$ relaxation dispersion data for aromatic side chains in proteins.

**Materials and methods**

**Protein samples**

The B1 domain of the bacterial antibody-binding protein G (GB1) containing the mutations T2Q, N8D and N37D (QDD variant) was expressed and purified as described previously (Lindman et al. 2006), with three different labeling patterns: (i) site-selective $^{13}$C/-$^{12}$C using 2-$^{13}$C1 glucose with natural abundance $^1$H incorporation (Lundström et al. 2007), (ii) site-selective $^1$H–$^{13}$C/-$^{12}$C labeling protein (Fig. 1) using site-selectively $^1$H/-$^{12}$C and $^{13}$C labeled -ketoacids as precursors (Lichtenecker 2014; Lichtenecker et al. 2013), 80 mg/L for Phe, 350 mg/L for Tyr, 10 mg/L for Trp, (iii) same as (ii) but with additional 70% background deuteration by expression in $D_2$O (implemented only in the case of Phe ε*). NMR samples contained around 1 mM protein in 20 mM HEPES, 90% H$_2$O/10% D$_2$O with addition of small amounts of NaN$_3$. The pH of the NMR samples was adjusted to 7.0.

![Fig. 1](https://example.com/fig1.png)

Labeling patterns in Phe, Tyr and Trp (from left to right), using site-selectively $^1$H/-$^{12}$C and $^{13}$C labeled -ketoacids as precursors (Lichtenecker 2014; Lichtenecker et al. 2013). Protons and deuterons are displayed by H and D, respectively. $^{13}$C is shown in red, other carbon positions are $^{12}$C.
NMR spectroscopy

All experiments were acquired on a Bruker Avance III spectrometer at a static magnetic field strength of 14.1 T, equipped with a room temperature probe. $^1$H $R_1\rho$ relaxation dispersion experiments were performed using the pulse sequence shown in Fig. 2, with spectral widths of 14.0 ppm ($^1$H) and 30.0 ppm ($^{13}$C), by 1024 and 128 points, respectively. Nineteen experiments were performed at temperatures of 15, 20, 25, 30 and 35 °C. $R_1\rho$ rate constants were measured in separate experiments conducted on-resonance with the signal of interest resulting in tilt angles of 90° from the z-axis, using a constant-time relaxation period (Mulder et al. 2001) of 20 ms and spin-lock field strengths varying between 1000 and 9000 Hz; the 4-ms adiabatic ramps used to align the magnetization along the effective spin-lock field do not achieve perfect alignment below 1000 Hz. Spectra were processed with NMRPipe (Delaglio et al. 1995) and analyzed with PINT (Ahlner et al. 2013).

Data analysis

Relaxation rates in the rotating reference frame were calculated from the signal intensities measured in presence and absence of the spin-lock pulse, $I(\omega_1)$ and $I_0$, respectively, according to (Mulder et al. 2001)

$$R_{1\rho}(\omega_1) = -\frac{1}{T_{SL}} \ln \left( \frac{I(\omega_1)}{I_0} \right)$$

$R_{1\rho}$ relaxation dispersions were fitted to the general equation for symmetric exchange derived by Miloushev and Palmer (2005) using fixed populations, $p_1=p_2=0.5$, and treating $\Delta \delta$ as a free parameter, which was compared to values measured under slow-exchange conditions at –5 °C and 200 MPa. Data modeling utilized the Levenberg–Marquardt nonlinear least-squares optimization algorithm (Press et al. 2002) implemented in MATLAB. Errors in the fitted parameters were estimated using 1000 Monte-Carlo simulations per fit; the reported errors correspond to one standard deviation.

Activation parameters of the ring flips were determined by non-linear regression of the flip rates, $k_{flip} = k_{ex}/2$, on the temperature $T$, using the Eyring equation. The Eyring equation was parameterized as

$$k_{flip} = \left( \frac{k_B T}{h} \right) \times \exp \left[ -\left( \Delta H^{\ddagger} - T \Delta S^{\ddagger} \right)/RT \right]$$

where $k_B$ and $h$ denote Boltzmann’s and Planck’s constants, respectively, and $\Delta H^{\ddagger}$ and $\Delta S^{\ddagger}$ are the activation enthalpy and activation entropy, respectively.

Hartmann–Hahn transfer calculations

A nucleus affected by a spin-lock radio frequency pulse with field strength $\omega_1 = -\gamma B_1$ will perceive an effective

$$\gamma_1 = -\frac{1}{T_{SL}} \ln \left( \frac{I(\omega_1)}{I_0} \right)$$

Fig. 2 Pulse sequence for the $^1$H $R_1\rho$ constant-time relaxation dispersion experiment for measuring conformational exchange of aromatic side chains in specifically $^1$H–$^{13}$C/$^2$H–$^{12}$C labeled proteins. All pulses are applied along the x-axis unless otherwise indicated. Narrow (wide) solid bars indicate rectangular high-power 90° (180°) pulses. The continuous-wave spin-lock relaxation periods $T$ and their flanking 4 ms tan/tanh adiabatic profiles (Mulder et al. 1998) are outlined in gray. The adiabatic sweep is initiated 25 kHz downfield or upfield of the spin-lock frequency. The wide semi-ellipse on $^{13}$C represents a REBURP (Geen and Freeman 1991) pulse with a bandwidth of 40 ppm. The pulse sequence can be modified to accommodate non-constant relaxation periods or off-resonance spin-locks by including a pair of $^{13}$C 180° pulses to mitigate the effects of dipole–dipole/CSA cross-correlated relaxation (Korzhnev et al. 2002; Massi et al. 2004). The delay $\tau$ can be set to 1.6 ms (Phe and Tyr), 1.35 ms (all aromatics) or 1.25 ms (His). The phase cycle is: $\phi_1=(4y, 4-y)$, $\phi_2=(x, -x)$, $\phi_3=(x, x, -x, -x)$, $\phi_{rec}=(x, -x, -x, -x, x, x, x, -x)$. Pulsed field gradients G1–4 are employed to suppress unwanted coherences and artifacts, while GC and GH are encoding and decoding gradients, respectively, for echo/anti-echo coherence selection, obtained by inverting the signs of GH (Kay et al. 1992). The delay $\Delta$ is equal to GC. For every second $t_1$ increment $\phi_2$ and the receiver were incremented. Gradient durations (in ms) and relative power levels (in %) are set to (duration, power level) G1=(1.0, 13), G2=(0.5, 10), G3=(1.0, 30), G4=(1.0, 90), GC=(1.0, 80), GH=(1.0, -20.1)
field strength $\omega_{\text{eff}} = (\Omega^2 + \omega_1^2)^{1/2}$ and will be oriented at an angle $\theta = \tan^{-1}(\omega_1/\Omega)$ from the static magnetic field, with $\Omega = \omega_0 - \omega_c$ denoting the offset between the nuclear precession frequency $\omega_0$ and the carrier frequency $\omega_c$ of the spin-lock pulse. In the case of two spins $I$ and $S$, coupled by a scalar-coupling constant $J$, Hartmann–Hahn matching of the two effective fields causes magnetization transfer between them, according to the coherence transfer function given by (Brath et al. 2006; van de Ven 1995)

$$F_{\text{HfHa}} = A_{\text{HfHa}} \sin^2(DT_{\text{SL}}/2)$$

(3)

where $T_{\text{SL}}$ is the duration of the spin-lock period, $D = (\Delta^2 + J_{\text{eff}}^2)^{1/2}$, $\Delta = \omega_{\text{ef},I} - \omega_{\text{ef},S}$, $J_{\text{eff}} = \frac{1}{2}I(1 + \cos(\theta_I - \theta_S))$ and $A_{\text{HfHa}}$ is the coherence transfer amplitude

$$A_{\text{HfHa}} = \frac{1}{1 + (\Delta^2/J_{\text{eff}}^2)^2}$$

(4)

Hartmann–Hahn mediated magnetization transfer from the monitored proton of the $^1\text{H}:-^13\text{C}$ moiety to vicinal protons in the aromatic ring scales the intensity by a factor of $1 - F_{\text{HfHa}}(\omega_1)$ for a given spin-lock period $T_{\text{SL}}$. Consequently, the apparent $R_{\text{IP}}$ rate constant is given by:

$$R_{1\rho,\text{app}}(\omega_1) = -\frac{1}{T_{\text{SL}}} \left( \frac{\ln(I(\omega_1)/I_0) + \ln(1 - F_{\text{HfHa}}(\omega_1))}{\omega_0} \right)$$

(5)

where the first term describes the $R_{\text{IP}}$ rate constant according to Eq. (1), i.e., in the absence of Hartmann–Hahn matching. Similarly, as a first approximation, in the case of $n$ protons coupled to the monitored proton, the apparent rate constant is:

$$R_{1\rho,\text{app}}(\omega_1) = -\frac{1}{T_{\text{SL}}} \left( \frac{\ln(I(\omega_1)/I_0) + \sum_{i}^{n} \ln(1 - F_{\text{HfHa}}(\omega_1))}{\omega_0} \right)$$

(6)

$A_{\text{HfHa}}$ and $F_{\text{HfHa}}$ were calculated for scalar-coupled protons in the aromatic rings of GB1 using $J = 3J_{\text{HH}} = 7$ Hz, $T_{\text{SL}} = 20$ ms, and the resonance frequencies measured in the spectrum acquired at $25{^\circ}\text{C}$: 0.22 ppm between Y33 $\varepsilon^*$ and $\delta^*$; 0.65 ppm (0.16 ppm) between F52 $\varepsilon^*$ and $\delta^*(\zeta)$; and 0.98 ppm (0.13 ppm) between W43 $\zeta$ and $\varepsilon^*$ (n2). In case of frequencies averaged by fast ring-flips, i.e. $\delta^*$ and $\varepsilon^*$ in Phe and Tyr, it is sufficient to use them directly.

Results and discussion

We performed aromatic $^1\text{H}$ $R_{\text{IP}}$, relaxation dispersion experiments on the small protein domain GB1, $^1\text{H}:-^13\text{C}$ labeled in three different ways: (i) natural abundance protonation and site selective $^{13}\text{C}/^{12}\text{C}$ labeling; (ii) additional selective deuteration of vicinal hydrogen sites in order to eradicate $^3J_{\text{HH}}$ couplings resulting in site-selective $^1\text{H}:-^13\text{C}/^2\text{H}:-^{12}\text{C}$ labeling (Fig. 1); or (iii) as (ii) with additional non-specific background deuteration at a level of 70%. There are five symmetric aromatic residues (Phe and Tyr) in GB1: Y3, Y33, Y30, Y45 and F52. All but Y33 undergo relatively slow ring-flip processes that have been studied by aromatic $^{13}\text{C}$ $R_{\text{IP}}$, relaxation dispersion experiments (Dreydoppel et al. 2020); in particular Y36, Y3ε, F30δ, Y45ε and F52ε have been studied previously and provide valuable points of reference for comparison with our present work. Individual cross peaks from the two symmetric sites of the aromatic rings could be observed under slow exchange conditions ($-5$ °C, 200 MPa) for Y36, Y3ε, F30δ, Y45ε and F52ε. Based on these results, the following $^{13}\text{C}$ and $^1\text{H}$ chemical shift differences could be extracted: Y36 (2.11 ppm; 0.40 ppm), Y3ε (1.40 ppm; 0.50 ppm), F30δ (5.39 ppm; 0.84 ppm), F30ε (0.00 ppm; 0.56 ppm) and F52ε (1.76 ppm; 0.00 ppm). F52ε, which has a $^1\text{H}$ chemical shift difference of zero, serves as a negative control in the present study, since it should not exhibit any exchange-mediated dependence of the $R_{\text{IP}}$ rate constant on the spin-lock field strength. In addition, Y33ε, which is solvent exposed and undergoing very fast ring flips, is also expected to show an essentially flat dispersion profile. Furthermore, the two labeled sites in tryptophan, W43δ1 and W43ζ3 also serve as useful controls: W43δ1 does not experience sizeable $^3J_{\text{HH}}$ couplings, and neither site undergoes exchange, since this residue does not readily undergo ring flips. In a previous study, site-selective $^1\text{H}:-^{13}\text{C}/^2\text{H}:-^{12}\text{C}$ labeling resulted in 99% correct labeling in case of Phe $\varepsilon^*$ and 75% in case of Trp $\zeta$, whereas Tyr $\varepsilon^*$ labeling yielded only 4% incorporation for reasons that are not clear (Raum et al. 2019). Here, we increased the amount of tyrosine precursor fourfold in the protein expression medium, resulting in 17% of the desired labeling for Tyr $\varepsilon^*$, while the remaining 83% have natural abundance isotope incorporation. Therefore, 0.9% of the sample is $^{13}\text{C}$ labeled at Tyr $\varepsilon$ and protonated at the $\delta$ position. Thus, in the present sample the latter isopomer is diluted by a factor of 20 compared to the desired one, which is sufficient to suppress $^3J_{\text{HH}}$ coupling artifacts.

Deuteration of vicinal protons is necessary in order to achieve artifact-free aromatic $^1\text{H} R_{\text{IP}}$, relaxation dispersion profiles

Figure 3 shows the resulting $^1\text{H} R_{\text{IP}}$, relaxation dispersion profiles for the $^1\text{H}:-^{13}\text{C}$ labeled $\varepsilon$ sites of residues Y3, Y33, F30, and F52 (a-d), and for the $\delta$1 and $\zeta$3 sites of W43 (e-f). There is a notable variation among the dispersion profiles of the different residues and between the two $^1\text{H}/^2\text{H}$ labeling patterns for each residue. The non-deuterated samples of residues Y33ε, F30ε, F52ε, and W43ζ3 (Figs. 3b–d and
f; red symbols) all exhibit anomalous, ‘inverted’ relaxation dispersion profiles characteristic of sizeable (7–8 Hz) $^3J_{\text{HH}}$ coupling (Raum et al. 2018). In contrast, the dispersion profile of Y3ε (Fig. 3a) does not appear anomalous, because the effect of Hartmann–Hahn matching is in part masked by a substantial chemical exchange contribution. In general, the magnitude of the artifact is more pronounced in the case of Phe (Fig. 3c, d) compared to Tyr (Fig. 3a, b), because the investigated $^1$Hε spins in Phe couple to two vicinal proton spins (δ and ζ) instead of one (δ) in Tyr. In the case of Trp ζ3, the magnitude is closer to the Tyr case. In the non-deuterated sample, W43δ1 displays a flat relaxation dispersion profile (Fig. 3e), as a result of its small coupling constant, $^3J_{\text{HH}} < 2$ Hz, and absence of exchange. This result is in keeping with the expectation outlined above regarding the role of W43δ1 as a negative control.

Site-selective deuteration of the vicinal hydrogen sites efficiently removes the effect of $^3J_{\text{HH}}$ coupling, resulting in ‘normal’-looking relaxation dispersion profiles (Fig. 3; blue symbols). Y33ε, F52ε, and W43ζ3 (Figs. 3b, d, f) show flat dispersion profiles in the deuterated sample, again in agreement with expectations (very fast exchange, no $^1$H chemical shift difference between the symmetric sites, and no exchange, respectively; see above). In the case of F30ε, the site-selectively deuterated sample reveals a fast exchange process that is completely hidden by the strong-coupling effect in the protonated sample (Fig. 3c). Finally, for Y3ε where the relaxation dispersion profile shows clear signs of exchange even in the non-deuterated sample, the profile is more pronounced in case of site-selective deuteration (Fig. 3a). This comparison clearly shows that the Y3ε profile in the non-deuterated sample is compromised and cannot be analyzed to yield reliable exchange parameters. Additional, unspecific 70% background deuteration of the protein does slightly shift the $^1$H $R_{1\rho}$ to lower values, but does not alter the profile shape (Fig. 3c, d, green symbols). A quantitative analysis comparing the results obtained from the samples with or without background deuteration shows that identical exchange parameters are obtained, within the margin of error (see below). This result demonstrates that site-selective deuteration in the aromatic ring alone is sufficient to obtain artifact free relaxation dispersion profiles, and further indicates that long-range ROE effects (Lundström and
Akke 2005; Weininger et al. 2013a) do not cause concern in the present case.

Taken together, the present results illustrate the detrimental effects of $J_{HH}$ couplings on the relaxation dispersion data obtained on non-deuterated samples, and clearly demonstrate that high-quality $^1$H $R_{1\rho}$ relaxation dispersion profiles can only be obtained using samples with site-selective deuteration. Our present approach enables $^1$H $R_{1\rho}$ relaxation dispersion experiments for Phe, Tyr, and Trp $\zeta3$, and thus completes the repertoire of probes to cover all aromatic $^1$H-$^1$C positions. By contrast, only Trp $\delta1$, His $\delta2$ and His $\varepsilon1$ are accessible also in non-deuterated samples, because these positions do not involve sizeable scalar coupling to vicinal protons.

**Hartmann–Hahn transfer explains artifacts in aromatic $^1$H $R_{1\rho}$ relaxation dispersion experiments**

The artifacts described above for the non-deuterated samples (Fig. 3) can be rationalized as increased homonuclear Hartmann–Hahn transfer from the proton of interest ($I$) to a scalar coupled vicinal proton ($S$) with increasing spin-lock field strength ($\omega_1$). Figure 4 shows the expected values of the amplitude ($A_{\text{HaHa}}$) of the transfer function and the relative loss of magnetization for spin $I$ ($1-F_{\text{HaHa}}$), as a function of the spin-lock field strength and the difference in offset between spins $I$ and $S$; see Eqs. (3–5). The magnetization loss expected for the case where spin $I$ has two $J$-coupled vicinal protons, as for Phe $\varepsilon$, is included in Fig. 4c; see Eq. (6). We calculated the amount of magnetization lost from $I$ during the spin-lock period for those residues that do not show exchange contributions to $R_{1\rho}$ ($Y33\varepsilon$, $F52\varepsilon$, and $W43\zeta3$), and estimated their resulting apparent $R_{1\rho}$ rate constant as a function of $\omega_1$. To do so, we took the average $R_{1\rho}$ rate constant measured on site-selectively $^1$H–$^1$C/$^2$H–$^1$C labeled samples to represent the artifact-free first term of Eq. (6), and added the second term involving the calculated value of ($1-F_{\text{HaHa}}$). As shown in Figs. 4d–f, in all three cases the experimental data points acquired on the uniformly protonated sample are reproduced well by the calculated function describing the apparent $R_{1\rho}$ values. Thus, Hartmann–Hahn transfer quantitatively explains the observed artifacts in aromatic
Comparison of ring flip rates of $^1$H and $^{13}$C based aromatic $R_{1\rho}$ relaxation dispersion experiments

We acquired relaxation dispersion profile for Y3ε and F30ε in the site-selectively deuterated sample at three different temperatures (Fig. 5). The artifact-free relaxation dispersions could be fitted to the general equation for symmetric exchange, resulting in flip rates at each temperature together with a global chemical shift difference for each residue (Table 1). The derived chemical shift differences are $\Delta \delta = (0.46 \pm 0.03)$ ppm for Y3ε and $(0.47 \pm 0.02)$ ppm for F30ε, in good agreement with the values measured in the spectrum under slow exchange conditions (0.50 ppm and 0.56 ppm, respectively). In the case of Y3ε, the resulting flip rates are virtually the same, within the margin of error, as those determined from $^{13}$C based experiments (Fig. 6a, Table 1). Consequently, the activation enthalpy ($\Delta H^\ddagger$) and activation entropy ($\Delta S^\ddagger$) determined from the temperature dependence of the flip rates are identical within the margin of error to previous estimates (Fig. 6a): $\Delta H^\ddagger = (82 \pm 8)$ kJ mol$^{-1}$ for $^1$H versus $(87 \pm 14)$ kJ mol$^{-1}$ for $^{13}$C; and $\Delta S^\ddagger = (111 \pm 26)$ J mol$^{-1}$ K$^{-1}$ versus $(126 \pm 46)$ J mol$^{-1}$ K$^{-1}$. In case of F30ε, we cannot directly compare flip rates determined by $^1$H and $^{13}$C $R_{1\rho}$ experiments, because its $^{13}$Ce chemical shift difference is zero. However, ring flip rates have been measured for F30ε using $^{13}$C $R_{1\rho}$ relaxation experiments at two higher temperatures, although the flip rate is far outside the optimal range of the $R_{1\rho}$ experiment in this case and should only be interpreted semi-quantitatively (Fig. 6b). At lower temperatures F30ε becomes broadened beyond detection, because of its large $^{13}$Cδ chemical shift difference. The rates derived from the $^1$H $R_{1\rho}$ experiments seem to be a bit higher (as gauged from the extrapolated line in Fig. 6b), although the two data points from the $^{13}$C $R_{1\rho}$ experiments seem to follow the temperature dependence determined from the $^1$H $R_{1\rho}$ experiments. Background deuteration in the F30ε sample leads to a minor improvement of the fit in the region of the highest rephasing frequencies (Fig. 5 bc), where the ROE effect is the most severe. The derived values of $\Delta H^\ddagger$ and $\Delta S^\ddagger$ with and without background deuteration are also identical, within the margin of error (Fig. 6b): $\Delta H^\ddagger = (47 \pm 4)$ kJ mol$^{-1}$ for $^1$H with protonated background versus $(50 \pm 1)$ kJ mol$^{-1}$ with deuterated background; and $\Delta S^\ddagger = (0 \pm 13)$ J mol$^{-1}$ K$^{-1}$ versus $(10 \pm 4)$ J mol$^{-1}$ K$^{-1}$. In summary, aromatic $^1$H $R_{1\rho}$ experiments allow the determination of correct exchange parameters, provided that the aromatic ring is site-selectively $^1$H/$^2$H labeled; further deuteration of the background is not needed.

Concluding remarks

There are multiple positions available in aromatic side chains that are suitable for studying exchange dynamics. However, it is not uncommon that certain sites do not experience sufficiently large $\Delta \delta$ for a given nucleus, or experience
exchange rates outside of the optimal relaxation dispersion window for a given nuclide (e.g., $^1$H or $^{13}$C). Thus, it is highly advantageous to be able to measure exchange using different nuclides, so as to probe as many sites as possible in a given protein. For example, our studies of ring flip dynamics in GB1 highlight the advantage of using both $^1$H and $^{13}$C based experiments: Y3 can be probed using all four positions ($^1$H$_\delta$, $^{13}$C$_\delta$, $^1$H$_\varepsilon$ and $^{13}$C$_\varepsilon$), while F30 and F52 can each be probed using only a single probe ($^1$H$_\varepsilon$ and $^{13}$C$_\varepsilon$, respectively). Our present approach extends the available toolbox to make all aromatic positions accessible by NMR relaxation dispersion experiments using either CPMG or $R_1\rho$ refocusing elements. The higher effective field strength available in $^1$H $R_1\rho$ experiments is clearly of advantage in this context, and allows to reliably study exchange processes up to an exchange rate of about 80,000 s$^{-1}$ (equal to a flip rate of 40,000 s$^{-1}$).

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Data availability All data generated or analysed during this study are included in this published article.

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References

Ahlner A, Carlsson M, Jonsson BH, Lundström P (2013) PINT: a software for integration of peak volumes and extraction of relaxation rates. J Biomol NMR 56:191–202. https://doi.org/10.1007/s10858-013-9737-7

Akke M, Palmer AG (1996) Monitoring Macromolecular motions on microsecond-millisecond time scales by $R_1\rho$ constant-relaxation-time NMR spectroscopy. J Am Chem Soc 118:911–912

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### Table 1 Ring flip rates and chemical shift differences

|          | Y3$^a$   | Y3$^b$   | F30$^a$  | F30$^b$  | F30$^c$  |
|----------|----------|----------|----------|----------|----------|
| $\Delta\delta$(ppm) | 0.46 ± 0.03 | 0.50     | 0.47 ± 0.02 | 0.49 ± 0.02 | 0.56     |
| $k_{flip}$($10^3$ s$^{-1}$) | 15 °C | 18 ± 1   | 20 ± 1    | 15 ± 1    | 12 ± 2   |
|          | 20 °C | 28 ± 1   | 29 ± 1    | 26 ± 2    | 22 ± 3   |
|          | 25 °C | 15 ± 2   | 12 ± 2    | 37 ± 1    | 42 ± 1   |
|          | 30 °C | 26 ± 2   | 22 ± 3    | 38 ± 4    | 53 ± 4   |
|          | 35 °C | 46 ± 3   | 38 ± 4    |          | 75 ± 8   |
|          | 40 °C |          |          |          |          |

$^a$Derived from aromatic $^1$H $R_1\rho$ relaxation dispersion experiments on position $\varepsilon$ of site-selective deuterated samples with protonated background.

$^b$Taken from (Dreydoppel et al. 2020); chemical shift difference of position $\varepsilon$ are derived from spectra under slow exchange conditions; ring flip rates are derived from aromatic $^{13}$C $R_1\rho$ relaxation dispersion experiments on Y3 $^{13}$C$\delta$ and F30 $^{13}$C$\delta$.

$^c$In the case of F30 the flip rates are clearly outside the optimal range of the experiment and should only be interpreted semi-quantitatively.

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**Fig. 6** Temperature dependence of ring-flip rates. $k_{flip}$ is plotted as a function of $1/T$. A Y3 and B F30 flip rates determined from aromatic $^1$H $R_1\rho$ relaxation dispersion measurements are shown in blue (protonated background) and green (deuterated background), and from previous $^{13}$C $R_1\rho$ relaxation dispersion measurements in grey (Dreydoppel et al. 2020). Solid lines represent non-linear regression of $k_{flip}$ on $T$ according to the Eyring equation, displayed on a logarithmic y-axis to show the expected linearity. Derived activation enthalpies ($\Delta H^\ddagger$) and entropies ($\Delta S^\ddagger$) are: Y3 $^1$H (protonated background), (82 ± 8) kJ mol$^{-1}$ and (111 ± 26) J mol$^{-1}$ K$^{-1}$; Y3 $^{13}$C (87 ± 14) kJ mol$^{-1}$ and (126 ± 46) J mol$^{-1}$ K$^{-1}$ (A, blue); F30 $^1$H (protonated background), (47 ± 4) kJ mol$^{-1}$ and (0 ± 13) J mol$^{-1}$ K$^{-1}$ (B, blue); F30 $^{13}$C (deuterated background), (50 ± 1) kJ mol$^{-1}$ and (10 ± 4) J mol$^{-1}$ K$^{-1}$ (B, green).

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$R_1\rho$ refocusing elements. The higher effective field strength available in $^1$H $R_1\rho$ experiments is clearly of advantage in this context, and allows to reliably study exchange processes up to an exchange rate of about 80,000 s$^{-1}$ (equal to a flip rate of 40,000 s$^{-1}$).
Press WH, Teukolsky SA, Vetterling WT, Flannery BP (2002) Numerical recipes in C++: the art of scientific computing, 2nd edn. Cambridge University Press, Cambridge

Raum HN, Dreydoppel M, Weininger U (2018) Conformational exchange of aromatic side chains by (1)H CPMG relaxation dispersion. J Biomol NMR 72:105–114. https://doi.org/10.1007/s10858-018-0210-5

Raum HN, Schorghuber J, Dreydoppel M, Lichtenecker RJ, Weininger U (2019) Site-selective (1)H(2)H labeling enables artifact-free (1)H CPMG relaxation dispersion experiments in aromatic side chains. J Biomol NMR 73:633–639. https://doi.org/10.1007/s10858-019-00275-z

Schorghuber J et al (2018) Late metabolic precursors for selective aromatic residue labeling. J Biomol NMR 71:129–140. https://doi.org/10.1007/s10858-018-0188-z

Steiner E, Schlagnitweit J, Lundstrom P, Petzold K (2016) Capturing excited states in the fast-intermediate exchange limit in biological systems using (HNMR)-H-1 spectroscopy. Angew Chem Int Edit 55:15869–15872. https://doi.org/10.1002/anie.201609102

Teilum K, Brath U, Lundström P, Akke M (2006) Biosynthetic C-13 labeling of aromatic side chains in proteins for NMR relaxation measurements. J Am Chem Soc 128:2506–2507

Vallurupalli P, Bouvignies G, Kay LE (2012) Studying “Invisible” excited protein states in slow exchange with a major state conformation. J Am Chem Soc 134:8148–8161. https://doi.org/10.1021/ja3001419

Vallurupalli P, Hansen DF, Stellor E, Meirovitch E, Kay LE (2007) Measurement of bond vector orientations in invisible excited states of proteins. Proc Natl Acad Sci USA 104:18473–18477

Vallurupalli P, Sekhar A, Yuwen TR, Kay LE (2017) Probing conformational dynamics in biomolecules via chemical exchange saturation transfer: a primer. J Biomol NMR 67:243–271. https://doi.org/10.1007/s10858-017-0099-4

van de Ven FJM (1995) Multidimensional NMR in liquids: basic principles and experimental methods. Wiley, New York

Wagner G (1980) Activation volumes for the rotational motion of interior aromatic rings in globular-proteins determined by high-resolution H-1-Nmr at variable pressure. FEBS Lett 112:280–284

Wagner G, Demarco A, Wüthrich K (1976) Dynamics of aromatic amino-acid residues in globular conformation of basic pancreatic trypsin-inhibitor (Bpti) 1 H-1 Nmr-studies. Biophys Struct Mech 2:139–158

Weininger U et al (2013) Protein conformational exchange measured by H-1 R-1 rho relaxation dispersion of methyl groups. J Biomol NMR 57:47–55

Weininger U (2019) Optimal isotope labeling of aromatic amino acid side chains for NMR studies of protein dynamics. Methods Enzymol 614:67–86. https://doi.org/10.1016/bs.mie.2018.08.028

Weininger U, Diehl C, Akke M (2012a) C-13 relaxation experiments for aromatic side chains employing longitudinal- and transverse-relaxation optimized NMR spectroscopy. J Biomol NMR 53:181–190

Weininger U, Respondek M, Akke M (2012b) Conformational exchange of aromatic side chains characterized by L-optimized TROSY-selected C-13 CPMG relaxation dispersion. J Biomol NMR 54:9–14

Weininger U, Respondek M, Löw C, Akke M (2013) Slow aromatic ring flips detected despite nearly-degenerate NMR frequencies of the exchanging nuclei. J Phys Chem B 117:9241–9247

Weininger U, Brath U, Modig K, Teilum K, Akke M (2014a) Off-resonance rotating-frame relaxation dispersion experiment for C-13 in aromatic side chains using L-optimized TROSY-selection. J Biomol NMR 59:23–29

Weininger U, Modig K, Akke M (2014b) Ring flips revisited: C-13 relaxation dispersion measurements of aromatic side chain dynamics and activation barriers in basic pancreatic trypsin inhibitor. Biochemistry 53:4519–4525

Weininger U, Modig K, Geitner AJ, Schmidpeter PAM, Koch JR, Akke M (2017) Dynamics of aromatic side chains in the active site of FKBP12. Biochemistry 56:334–343. https://doi.org/10.1021/acs.biochem.6b01157

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