Adiabatic TOCSY MAS in liquids.
E. Kupce, P. A. Keifer, M. Delepierre

To cite this version:

HAL Id: pasteur-00164702
https://hal-pasteur.archives-ouvertes.fr/pasteur-00164702
Submitted on 25 Jul 2007

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Adiabatic TOCSY MAS in Liquids

E. Kupce, a,* P. A. Keifer b and M. Delepierre c

a Varian Inc., 28, Manor Road, Walton-on Thames, Oxford, UK
b Varian Inc., 3120 Hansen Way, Palo Alto, CA, USA
c Lab. RMN, Institut Pasteur, 28 rue du Dr Roux, Paris, France

# Presented in part at the 41st Experimental NMR Conference at Asilomar, CA, USA, 10 April 2000.

* To whom correspondence should be addressed; e-mail:
eriks.kupce@varianinc.com
ABSTRACT

The effect of magic angle spinning (MAS) of liquids upon the performance of various isotropic mixing sequences is investigated. Although the mathematical formalism for isotropic mixing under MAS conditions is similar for both liquids and solids, the mechanism through which the coherence transfer is disturbed is different. In liquids, the use of sample spinning in the presence of both RF and magnetic-field inhomogeneities introduces a modulation of the effective field, which compromises the performance of the conventional mixing sequences. This effect is further amplified by supercycles, which normally improve the performance of the mixing and decoupling experiments. It is demonstrated that adiabatic mixing sequences are less susceptible to such modulations and perform considerably better in TOCSY MAS experiments. The best performance of TOCSY MAS is observed at the rotational resonance condition when the sample appears static in the RF reference frame.
INTRODUCTION

The small active volumes and the high sensitivities of Varian Nanoprobes (1) make them ideal for applications where only a limited amount of sample is available. In such probes high sensitivity is achieved by maximizing the filling factor of the receiver coil. The line broadenings caused by the resulting magnetic susceptibility discontinuities are eliminated by using susceptibility matched materials and by magic angle spinning (MAS), typically at 2 to 4 kHz speed. This use of MAS has the additional advantage of suppressing line broadenings in heterogeneous samples that have a non-uniform magnetic susceptibility across the sample, such as solvent-swollen resins (2,3). High resolution and high sensitivity allows routine liquids NMR experiments to be performed on a single bead of solid phase synthesis (SPS) resins with typical sample quantities of under 500 picomoles (4).

Although the vast majority of liquid-state NMR experiments can be recorded in Nanoprobes with no complications, in some cases interference effects have been observed (5). Particularly, performance of the conventional, composite-pulse 2D TOCSY experiment can be extremely poor and unstable (see Fig. 1a). On the other hand, the adiabatic TOCSY
experiment (6) under the same conditions performs considerably better (see Fig. 1b).

In the course of this study we evaluated a number of standard liquid-state experiments at a wide variety of MAS rates. We verified that the signal-to-noise ratio was always practically independent of the MAS rate for those experiments that did not contain a spin-lock, for instance, NOESY, DQF-COSY or simple spin-echo sequences. On the other hand, the performance of experiments employing recursive spin-lock sequences, such as isotropic mixing and spin decoupling, were very sensitive to the sample spinning speed (see Fig. 2a). The use of adiabatic spin-locks typically improved the overall quality of the data (see Fig. 2b). We also found that the magnitude of these effects strongly depended upon the sample volume, the sample geometry, and even the quality of shimming. Surprisingly, experiments that employ continuous wave spin-locks, for instance off-resonance ROESY (7) or selective cross polarization experiments (8,9), seemed to be substantially less sensitive to variation of MAS speed.

In this study we explored the effects of various spin-lock sequences on liquid samples spinning at the magic angle. It is demonstrated that the spin dynamics of experiments that employ repetitive sequences of composite pulses, such as isotropic mixing or spin decoupling, is least disturbed when
the sample spinning speed is comparable to the spin nutation frequency. This condition is known as rotational resonance (10) in solid-state NMR, where it has found many useful applications.

**DISCUSSION**

The presence of rotational resonance in fast rotating liquids can be demonstrated by a simple nutation experiment (see Fig 3). As shown in Fig. 3a the homogeneity of both the RF field and the magnetic field in a static sample is rather poor, partially because of somewhat casual sample shimming. This inhomogeneity is then largely removed by sample spinning at the magic angle (see Fig. 3b). However, if the RF field strength matches the spinning speed, a very sharp peak appears in the indirect dimension, as if the RF field was suddenly extremely homogenous (see Fig 3c). In fact, the magnetization does not de-phase during this continuous wave spin lock period because it is locked by the effective field (11), i.e. the magnetization follows the effective field in a manner similar to the adiabatic following (12). The broad component along the RF field axis in Fig. 3c comes from the parts of the sample where the rotational resonance condition is not met.

It is interesting to compare cross polarization (CP) in solids (13-15) with isotropic mixing in liquids (16-19). In both cases the spin dynamics can
be described using the same formalism. The rotating-frame Hamiltonian is derived for an isolated IS spin pair:

\[
H_{\text{CP}} = -\omega_I I_x - \Delta\omega_I I_z - \omega_S S_x - \Delta\omega_S S_z + \mathcal{J} I_z S_z + H_\mu \tag{1}
\]

where \(\omega_I\) represents the RF field strengths as seen by the I and S spins and the \(\Delta\omega\) terms are the corresponding resonance offsets. In solids the polarization transfer is achieved via the dipolar coupling \((\mathcal{J} = b_{IS})\) whereas in liquids this process is mediated by the indirect (scalar) spin-spin coupling \((\mathcal{J} = J_{IS})\). Assuming that \(\omega_I \gg \mathcal{J}\) and ignoring the homonuclear interactions, the effective Hamiltonian in a tilted reference frame for a static sample is written as:

\[
H_{\text{eff}} = -\omega_{IS} I_x - \omega_{IS} S_x + \mathcal{J}_{\text{eff}} I_z S_z \tag{2}
\]

With the Hartmann-Hahn condition satisfied the first two terms in Eq. [2] vanish and the maximal polarization transfer can be achieved.

When the sample is spun at the magic angle, the heteronuclear dipolar coupling, \(b_{IS}\) becomes time dependent:

\[
b_{IS}(t) = \sum_k b_k e^{ik\omega_\perp t} \tag{3}
\]

This introduces the well known "finger pattern" into the CP spectra as the Hartmann-Hahn condition splits into several sub-conditions:

\[
\Delta = \pm k\omega_\perp, \tag{4}
\]
where $\Delta$ is the RF mismatch parameter, $k$ is an integer (usually $-2 \leq k \leq 2$) and $\omega_r$ is the spinning speed. The efficiency of CP in solids can be substantially improved by employing adiabatic $B_1$ field sweeps (20-23).

In liquids the scalar $J_{iS}$ coupling is not directly affected by sample spinning. However, the effective fields seen by spins $I$ and $S$ are time dependent, in a manner similar to that for $b_{iS}$ (see Eq. [3]). In order to show that such modulations can indeed affect the quality of spin lock experiments, we simulated the influence of RF field modulation on the ability of various mixing sequences to return the starting magnetization to its initial state at the end of a single cycle. As it appears from Fig. 4, the composite pulse sequences are most vulnerable to sample spinning (24). This is hardly surprising, because the frequency spectrum of composite pulse sequences is particularly "noisy" (25) and additional modulation introduced by sample spinning destroys the delicate balance of RF fields created by numerical optimization. Additional simulations (not shown here) indicate that increasing the length of the initial 4 step MLEV-4 supercycle (26) to either MLEV-16 or a 20-step phase cycle (27, 28) makes the performance of mixing sequences in rotating samples even worse. Indeed, supercycles have been designed to work with inversion pulses that are all identical, whereas in the presence of sample spinning the performance of individual (composite)
inversion pulses may vary substantially. As a result, at the end of the cycle the magnetization is not returned back to its initial state, as it would be in a static sample. For the same reason, the overall performance of the TOCSY experiment becomes increasingly sensitive to MAS speed as the length of a supercycle increases.

Therefore, the optimum performance of isotropic mixing and spin decoupling can be expected if the length of individual inversion pulses, $T_p$, matches the rotor period, $T_r$. This is indeed confirmed by experiment (see Fig. 5). Under these circumstances, in the RF reference frame the sample looks as if it is static and the individual contributions from $\omega_1$ and $\Delta \omega$ modulation can be regarded as local distortions of the mixing field. As long as these distortions can be tolerated by the applied spin-lock sequence, the performance of the experiment is acceptable. Adiabatic pulses are known to be particularly insensitive to RF field distortions (29). Provided the rotational resonance condition is met, the efficiency of spin mixing is further improved by supercycles.

**CONCLUSIONS**

Although the appearance of interference between the Hartmann-Hahn polarization transfer and sample spinning is similar in solids and in liquids,
the mechanisms affecting the experimental performance are quite different. The conventional composite-pulse mixing sequences are particularly vulnerable to such interference effects. The performance of TOCSY experiments can be substantially improved by using adiabatic (WURST-2) mixing sequences. The optimum performance in 2D TOCSY experiments is achieved at the rotational-resonance condition, when the repetition rate of the adiabatic pulses in a mixing sequence matches the MAS speed. Although both the experimental data and simulations indicate existence of other areas where the spin lock performance is close to the optimal (see Figs. 4 and 5), they are typically less efficient and the relation to the MAS speed is not as obvious.

ACKNOWLEDGEMENTS

We thank Tom Barbara, Knut Mehr, and Dave Rice from Varian Inc. (Palo Alto) for many useful discussions and suggestions.

REFERENCES


Figure Captions

**Figure 1.** Comparison of 2D H-H correlated xy-TOCSY spectra of a small protein (5) from scorpion venom (1 mM in D$_2$O) recorded at 500 MHz using a 40 µl sample in Varian's Nano-NMR probe at 2.5 kHz spinning speed; a) adiabatic (WURST-2) mixing, $B_{1\text{max}} = 6.44$ kHz, length of a single adiabatic inversion pulse $T_p = 0.4$ ms; b) composite pulse mixing (DIPSI-2), $B_{1\text{max}} = 6.94$ kHz, length of a single composite inversion pulse $T_p = 1.036$ ms. Both spectra were acquired using a 60 ms mixing time, 128 increments in $t_1$ and 4 scans per increment. The total experiment time for each dataset was approximately 36 minutes.

**Figure 2.** Matrix plots of the first trace from H-H z-TOCSY spectra of 6 mg quinine in 40 µl CD$_2$Cl$_2$ recorded on INOVA 500 with gHX Nanoprobe; a) using DIPSI-2 spin lock and b) using WURST-2 spin lock. The vertical axis represents MAS spin rates ranging from 1100 Hz to 3200 Hz (from bottom to top; in 22 steps of 100 Hz each). The horizontal axis shows increasing mixing times ranging from 0 to 100 ms (from left to right; in 21 steps of 5 ms each). No effort was made to synchronize the beginning of the experiment with the rotor position.
**Figure 3.** Nutation experiment recorded at 500 MHz in gHX Nanoprobe using H$_2$O sample. (a) static sample, $B_1$ field strength of 25 kHz; (b) as in (a) but with sample spinning at the magic angle at 2.5 kHz speed; (c) the same as in (b) but using 2.5 kHz $B_1$ field strength. The sharp peak along the RF dimension (vertical) is due to the rotational resonance (see text). The broad component comes from parts of the sample where the rotational resonance condition is not met.

**Figure 4.** Simulation of the effect of MAS upon the performance of (a) adiabatic (WURST-2) and (b) composite pulse (DIPSI-2) mixing. The 2D plots reveal the ability of the particular mixing sequence to return the magnetization to its initial state ($M_z$) after a complete MLEV-4 phase cycle. The amplitude and frequency of the RF field was modulated according to Eq. [3]. The modulation depth was 10 % for the first harmonic and 5 % for the second harmonic. The contour plots show residual $M_z$ magnetization. The contour levels were increased linearly from 0 to 1 in steps of 0.1. Ideally, very little residual magnetization should be observed (see text).
**Figure 5.** An illustration of the effect of the spinning speed upon the performance of adiabatic xy-TOCSY experiment as in Fig. 1b. The strongest response is observed at the rotational resonance condition when the MAS rate matches the repetition rate of individual adiabatic inversion pulses. Note that this experiment is more sensitive to the MAS rate than z-TOCSY experiment shown in Fig. 2. Poor shimming further enhances the effect.