Contents lists available at ScienceDirect

Journal of Magnetic Resonance

journal homepage: www.elsevier.com/locate/jmr



Efficient cross-polarization using a composite 0° pulse for NMR studies on static solids

Masashi Fukuchi^{a,b}, Ayyalusamy Ramamoorthy^c, K. Takegoshi^{a,b,*}

^a Division of Chemistry, Graduate School of Science, Kyoto University, Kyoto 606-8502, Japan^b Japan Science and Technology Agency, CREST, 5, Sanbancho, Chiyoda-ku, Tokyo 102-0075, Japan

^c Department of Chemistry and Biophysics, University of Michigan, Ann Arbor, MI 48109-1055, USA

ARTICLE INFO

Article history: Received 6 September 2008 Revised 9 October 2008 Available online 17 October 2008

Keywords: COMPOZER-CP Composite 0° pulse Liquid crystalline materials Lipid membranes

ABSTRACT

In most solid-state NMR experiments, cross-polarization is an essential step to detect low- γ nuclei such as ¹³C and ¹⁵N. In this study, we present a new cross-polarization scheme using spin-locks composed of composite 0° pulses in the RF channels of high- γ and low- γ nuclei to establish the Hartmann–Hahn match. The composite 0° pulses with no net nutation-angle{ $(2\pi)_X - (2\pi)_{-X} - (2\pi)_Y - (2\pi)_{-Y} - \}_n$ applied simultaneously to both high- γ (*I*) and low- γ (*S*) nuclei create an effective heteronuclear dipolar Hamiltonian $H_d^{(0)} = \frac{d}{2}(2I_ZS_Z + I_XS_X + I_YS_Y)$, which is capable of transferring the *Z*-component of the *I* spin magnetization to the *Z*-component of the *S* spin magnetization. It also retains a homonuclear dipolar coupling Hamiltonian that enables the flip-flop transfer among abundant spins. While our experimental results indicate that the new pulse sequence, called *composite zero cross-polarization* (COMPOZER-CP) performs well on adamantane, it is expected to be more valuable to study semi-solids like liquid crystalline materials and model lipid membranes. Theoretical analysis of COMPOZER-CP is presented along with experimental results. Our experimental results demonstrate that COMPOZER-CP overcomes the RF field inhomogeneity and Hartmann–Hahn mismatch for static solids. Experimental results comparing the performance of COMPOZER-CP with that of the traditional constant-amplitude CP and rampCP sequences are also presented in this paper.

© 2008 Elsevier Inc. All rights reserved.

1. Introduction

Cross-polarization is a routinely used solid-state NMR technique to enhance the sensitivity of low- γ nuclei such as ¹³C and ¹⁵N under static and magic-angle spinning (MAS) experimental conditions. There are a number of studies that have well analyzed the theory underlying the CP phenomenon and the structure of spin Hamiltonians under double spin-lock in CP [1-4]. Several studies have shown that the efficiency of the originally reported CP pulse sequence using cw spin-locking RF-irradiation is highly dependent on resonance offset and inhomogeneity of RF field. To overcome these problems, several different types of CP pulse sequences such as CPMOIST [5], VACP [6], rampCP [7], adiabatic-CP [8], and SADIS-CP [9] have been reported in the literatures. There are also other types of CP sequences such as LGCP [10], FFLG-CP [10], CP utilizing offset [11] or time-averaged precession frequency (TAPF) [12] for RF power reduction and WIM-CP [13] that have been demonstrated for additional specific applications. While most of the CP pulse sequences reported so far in the literature have been shown to be useful for studies on a variety of solids, they all have some limitations. By far, the rampCP is the most robust to overcome all of the above-mentioned experimental problems and therefore it is very commonly utilized in solid-state NMR applications. However, it has been noted that the rampCP requires a longer CP contact time, which is not desirable for samples with short T_{10} or samples that deteriorate under RF heating.

In this work, we have proposed a new cross-polarization pulse sequence using a 2π pulse as a building block. This choice is based on its built-in robustness for RF inhomogeneity as well as RF offset. To enhance these features, we may use a pair of 2π pulses with opposite phases X and -X, that we refer to as a composite 0° pulse (Fig. 1a). The pair, however, does not provide the spin-locking field necessary for CP. In the conventional CP sequences mentioned above, transfer of transverse magnetization of heteronuclear spins occurs along the spin-locking RF fields at the X- or Y-axis in the rotating frame. On the other hand, CP along the Z-axis can be achieved either using WIM-24 spin-lock as in the HIMSELF sequence [14] or by the rotor-synchronized 12-rotor sequence under MAS [15]. In both these methods, an isotropic heteronuclear dipolar coupling Hamiltonian is realized by multiple pulses by suppressing the homonuclear dipolar interaction. While this feature has been well utilized in the line-narrowing studies, it is not useful for other experiments like CP to enhance the sensitivity of rare spins. In this work, we have proposed a new cross-polarization



^{*} Corresponding author. Address: Division of Chemistry, Graduate School of Science, Kyoto University, Kyoto 606-8502, Japan. Fax: +81 75 753 4011. *E-mail address:* takeyan@kuchem.kyoto-u.ac.jp (K. Takegoshi).

^{1090-7807/\$ -} see front matter © 2008 Elsevier Inc. All rights reserved. doi:10.1016/j.jmr.2008.10.013



Fig. 1. Radio frequency pulse sequences for COMPOZER-CP: (a) a pair of two 2π pulse, (b) the basic COMPOZER-CP, and (c) the supercycled COMPOZER-CP.

pulse sequence which makes it possible to create the enhanced *Z*magnetization of rare spins. We have also explored the possibility of improving the efficiency of cross-polarization particularly for experimental conditions with poor RF field inhomogeneity and Hartmann–Hahn (HH) mismatch. We have designed a spin-lock sequence based on a composite 0° pulse and implemented in the CP pulse sequence for experiments under static experimental conditions (Fig. 1b and c). The cross-polarization efficiency of this new pulse sequence is demonstrated on adamantane. Theoretical treatment and experimental analyses to evaluate the efficiency of the new pulse sequence are reported in this paper.

2. Experimental

All experiments were performed on a 300 MHz Chemagnetics/ Varian solid-state NMR spectrometer operating at 300.44 and 75.55 MHz for ¹H and ¹³C nuclei, respectively. A 5 mm double-resonance Doty MAS probe was used at room temperature. The rffield intensity for ¹H 90° pulse, ¹³C 90° pulse and CP was 62.5 kHz, and both the nutation and the phase-modulation angles in the TPPM decoupling [16] were optimized for an RF intensity of ≈40 kHz to be 180° and ±7.5, respectively. The ramping of the spin-lock in rampCP was ±5 kHz. Recycle delay was 2 s for all experiments. The ¹³C carrier frequency was chosen to be 33.6 ppm from TMS.

2.1. Theory

There are several studies that reported the spin Hamiltonians and spin dynamics under CP. Therefore, in this paper, we restrict our theoretical treatment to derive the effective Hamiltonians of hetero- and homonuclear dipolar couplings and the effect of offset under the double spin-lock of COMPOZER-CP sequence (Fig. 1).

We consider a heteronuclear two spin system (I and S) under RF-irradiation for a static sample. The dipolar interaction between these two spins is given by the Hamiltonian, $H_d = dI_Z S_Z$, where d is the dipole coupling constant. For simplicity, we neglect the chemical shift anisotropy interaction for both nuclei and assume the frequencies of the applied RF fields are on-resonance to the respective Larmor frequency of I and S spins. The relevant Hamiltonian in a frame rotating at its respective Larmor frequency (i.e., the double rotating frame) is given by

$$H = H_{rf} + H_d \tag{1}$$

where H_{rf} represents the double RF-irradiation along the X-axis and it is written as

$$H_{rf} = \omega_{1I}I_X + \omega_{1S}S_X \tag{2}$$

where ω_{1I} and ω_{1S} are RF field strengths in the *I* and *S* nuclear channels, respectively. The evolution of the spin system during the double RF-irradiation is defined by the propagator U(t) which is given by

$$U(t) = T \exp\left\{-i \int_0^t dt' H\right\}$$
(3)

where T is the Dyson time-ordering operator. It is possible to separate the evolution of the spin system under the heteronuclear dipolar interaction from that under the RF interaction as follows:

$$U(t) = U_0(t)U_1(t)$$
(4)

with

$$U_0(t) = T \exp\left\{-i \int_0^t dt' H_{rf}\right\} = \exp\{-iH_{rf}t\}$$
(5)

$$U_1(t) = T \exp\left\{-i \int_0^t dt' H_d(t')\right\}$$
(6)

and

$$H_d(t) = U_0^{-1}(t)H_dU_0(t)$$
(7)

This equation signifies the time-dependence of the heteronuclear dipolar interaction due to the RF-irradiations, and the timedependent Hamiltonian can be derived as

$$H_d(t) = d\{I_Z \cos(\omega_{1I}t) + I_Y \sin(\omega_{1I}t)\}\{S_Z \cos(\omega_{1S}t) + S_Y \sin(\omega_{1S}t)\}$$
(8)

Eq. (6) can be expressed by using the Magnus expansion [17] as follows:

$$U_1(\tau) = \exp\{-i(H_d^{(0)} + H_d^{(1)} + ...)\tau\}$$
(9)

$$H_d^{(0)} = \frac{1}{\tau} \int_0^\tau dt H_d(t)$$
 (10)

$$H_d^{(1)} = \frac{-i}{2\tau} \int_0^\tau dt_1 \int_0^{t_1} dt_2 [H_d(t_1), H_d(t_2)]$$
(11)

The zeroth-order average Hamiltonian (Eq. (10)) for an RF-irradiation period t_W may be written as follows:

$$H_{d}^{(0)} = \frac{d}{2t_{W}} \left[\left\{ \frac{\sin(\Sigma t_{W})}{\Sigma} + \frac{\sin(\Delta t_{W})}{\Delta} \right\} I_{Z} S_{Z} + \left\{ \frac{1 - \cos(\Sigma t_{W})}{\Sigma} - \frac{1 - \cos(\Delta t_{W})}{\Delta} \right\} I_{Z} S_{Y} + \left\{ \frac{1 - \cos(\Sigma t_{W})}{\Sigma} + \frac{1 - \cos(\Delta t_{W})}{\Delta} \right\} I_{Z} S_{Z} - \left\{ \frac{\sin(\Sigma t_{W})}{\Sigma} - \frac{\sin(\Delta t_{W})}{\Delta} \right\} I_{Y} S_{Z} \right]$$
(12)

where $\Sigma = \omega_{1l} + \omega_{1s}$ and $\Delta = \omega_{1l} - \omega_{1s}$. For the conventional crosspolarization experiment at the exact Hartman–Hahn matching condition ($\Delta = 0$), we may choose a cycle time t_W to satisfy $\Sigma t_w = 2n\pi(U_0(t_c) = U_0(t = 0) = 1)$, where *n* is an integer. Then, the zeroth-order heteronuclear dipolar coupling Hamiltonian $H_d^{(0)}$ becomes

$$H_d^{(0)} = \frac{d}{2}(I_Z S_Z + I_Y S_Y) \tag{13}$$

Since this form of the dipolar coupling Hamiltonian commutes with H_{rf} , the CP process is defined by $H_d^{(0)}$ to achieve a quasi-stationary spin-state, which is basically the eigen-state of the total

Hamiltonian $H_{rf} + H_d^{(0)}$. A double spin-lock under the regular CP therefore transfers the *X* magnetization of *I* nuclei to *S* nuclei and the S_X magnetization is detected by decoupling *I* nuclei.

In this study, we are interested in designing a CP sequence with the composite 0° pulses $(2\pi)_X - (2\pi)_{-X}$ as a building block, where $(2\pi)_{\phi}$ denotes that the spin-lock pulse has a 360° flip angle with a phase ϕ . Since the $(2\pi)_X - (2\pi)_{-X}$ sequence does not provide the spin-locking field that can build-up the transverse magnetization, the *Z*-axis for trapping the magnetization can be used. For this to happen, the effective Hamiltonian should have a flip–flop term ($I_XS_X + I_YS_Y$). The I_XS_X term in the Hamiltonian can be generated when the phase of the spin-lock is shifted from *X*- to *Y*-axis at $t = t_W$. Therefore, the basic sequence can be $(2\pi)_X - (2\pi)_Y$ and the total zeroth-order average Hamiltonian for a period $t = 0 \sim 2t_W$ becomes

$$H_d^{(0)} = \frac{d}{2} (2I_Z S_Z + I_X S_X + I_Y S_Y)$$
(14)

This zeroth-order heteronuclear dipolar coupling Hamiltonian under the new double spin-lock sequence commutes with $I_Z + S_Z$ but not with $I_Z - S_Z$. Therefore, this Hamiltonian brings quasi-equilibrium longitudinal magnetizations among *I* and *S* spin system. This suggests that it transfers the I_Z magnetization to S_Z . Furthermore, this double spin-lock sequence does not eliminate homonuclear dipolar interactions. In fact, the homonuclear dipolar Hamiltonian $H_D = D(3I_{Z1}I_{Z2} - I_1I_2)$ under the double spin-lock sequence transforms to

$$H_D^{(0)} = \frac{D}{2} (3I_{21}I_{22} + I_1I_2)$$
(15)

at the zeroth-order approximation.

Since RF pulses are not perfect, an offset typically deteriorates the efficiency of an RF pulse. Therefore, it is important to evaluate the role of offset during the new heteronuclear double spin-locks. When the RF-irradiation is at off-resonance, we need to add the offset Hamiltonian, $H_z = \omega_I I_Z + \omega_S S_Z$, for both spins in Eq. (1). Then, evolution operator given in Eq. (6) becomes

$$U_1(t) = T \exp\left[-i \int_0^t dt' \{H_d(t') + H_Z(t')\}\right]$$
(16)

with the time-dependent offset Hamiltonian as

$$H_{Z}(t) = \omega_{I} \{ I_{Z} \cos(\omega_{1I}t) + I_{Y} \sin(\omega_{1I}t) \} + \omega_{S} \{ S_{Z} \cos(\omega_{1S}t) + S_{Y} \sin(\omega_{1S}t) \}$$

$$(17)$$

To simplify the structure of this Hamiltonian, let us omit the offset terms of the *S* spin. Then the zeroth-order average Hamiltonian of the time-dependent *I* spin offset Hamiltonian, $H_Z(t)$ in Eq. (17), for the irradiation period t_W becomes:

$$H_{Z}^{(0)} = \frac{\omega_{I}}{\omega_{11}t_{W}} \{ I_{Z} \sin(\omega_{11}t_{W}) + I_{Y} - I_{Y} \cos(\omega_{11}t_{W}) \}$$
(18)

By applying the above-mentioned cyclic condition of RF spinlock irradiation $\Sigma t_W = 2n\pi$ and $\Delta = 0$ to Eq. (18), we find that $H_Z^{(0)} = 0$ for even values of *n*. Therefore, offset effects can be suppressed during the spin-lock irradiation. Further, to reduce the effects of the RF field inhomogeneity, phases of the spin-lock pulses are inverted to generate a spin-lock sequence $\{(2\pi)_X - (2\pi)_{-X} - (2\pi)_{Y} - (2\pi)_{-Y} - \}_n$. It can be shown that this spin-lock pulse sequence also eliminates the offset and RF field inhomogeneity effects in the *S* spin channel as well. As shown in Fig. 1b, the new spin-lock pulse sequence is applied in both RF channels for the transfer of *Z*-magnetization component from *I* to *S* nuclei without the need for a separate preparation pulse in *I* spin channel. The net combination of pulses in the spin-lock has a 0° nutation-angle and therefore it is named as COMPOZER-CP (*composite zero* cross*p*olarization). In Fig. 1c, we show the COMPOZER-CP sequence which employs the basic COMPOZER-CP sequence (Fig. 1b) as supercycles.

Lastly, we would like to point out that when we interchange the order of the *Y* and -Y pulses in one of the RF channel, the resulting zeroth-order heteronuclear Hamiltonian contains a flop–flop term $(I_XS_X - I_YS_Y)$, leading to the transfer of I_Z to $-S_Z$. This gives us a chance to examine the relative phases between the *I* and *S* channels of our spectrometer. In fact, we realized that the rotational directions of our ¹H and ¹³C pulses, that is, the "naming" the *Y* and -Y are opposite in our spectrometer.

3. Results and discussion

Fig. 2 shows the nutation-angle (θ_w) dependence of the ¹³C signal intensity of adamantane (stational powder) at a contact time of 1.2 ms, where θ_w is given as $\omega_1 t_w$ in the supercycled COMPOZER-CP (Fig. 1c). Since we have not suppressed the ¹³C longitudinal magnetization before the application of the COMPOZER-CP sequence, the intensities at $\theta_w = 0$ are not zero. It is shown that the application of the supercycled COMPOZER-CP sequence leads the -Z-magnetization (the open circles and triangles), showing the definition of the pulse rotational axis of our ¹H and ¹³C RF channels are opposite. Exchanging the Y and -Y pulses in the ¹³C channel, the transferred ¹³C magnetization becomes positive (the closed circles and triangles). These results show that the theoretical treatment presented above is appropriate and both of the flip-flop and flop-flop transitions can be used for CP. Fig. 2 also shows that the CP efficiency becomes large, when θ_w is equal to $n\pi$ (n = integer), which is also in agreement with the theoretical prediction. In the experimental results presented below, we used $\theta_w = 2\pi$ pulses.

In Fig. 3, we compared the effect of HH mismatch for the conventional CP, the rampCP, and the COMPOZER-CP. Fig. 3a and b shows the normalized intensities of the CH₂ and CH peaks of adamantane, respectively, against the HH mismatch ($\Delta = \omega_{1C} - \omega_{1H}$). In this experiment, we suppressed the ¹³C thermal-equilibrium magnetization before the application of the CP sequence. Clearly, the COMPOZER-CP sequences are superior to the CP and rampCP







Fig. 3. Performance of CP sequences against the HH mismatch ($\Delta = \omega_{1C} - \omega_{1H}$) using a static adamantane sample: (a) CH₂ peak and (b) CH peak. Crosses, open triangles, closed squares, and closed circles indicate the intensities obtained from CP, rampCP, COMPOZER-CP (Fig. 1b), and the supercycled COMPOZER-CP (Fig. 1c) experiments, respectively. The signal intensities were normalized by the maximum intensity of the CH₂ or CH peak obtained in a CP. A contact time of 7.92 ms was used in all experiments.

sequences in the compensation of the HH mismatch. Fig. 3 shows that the COMPOZER-CP sequences are capable of suppressing HH mismatch over Δ/ω_{1C} = ±20%. In Fig. 4, we compared the contacttime-dependence of the ¹³C magnetization for these sequences. Fig. 4a and b shows the normalized intensities of the CH₂ and CH peaks, respectively. It is shown that the build-up rates for COMPO-ZER-CP sequences are faster than that of the rampCP sequence whereas they are slower than that of the CP sequence. The term "spin-lock" is used in this paper as the magnetization of both I and S nuclei are locked along the Z-axis. This is strictly not the same as the "spin-lock" used in regular CP experiments where transverse magnetization is locked along the RF field axis. Fig. 4, however, shows that the COMPOZER-CP does achieve spin-locking by retaining the magnetization of ¹H and ¹³C nuclei along the Z-axis, and that there is no significant difference between the spin-locking efficacies of rampCP and COMPOZER-CP.

Effects of MAS on COMPOZER-CP have not been examined theoretically, however, our preliminary results indicate that its efficiency is deteriorated under MAS. For example, Fig. 5 shows the effect of the HH mismatch under MAS ($v_r = 10$ kHz) for the CH₂ carbon of adamantane. For the rampCP and the conventional CP, the intensity becomes large at the sideband HH matching condition of $\Delta = n\omega_R$ (ω_R is the spinning frequency). On the other hand, the strong RF mismatch cancellation by COMPOZER-CP leads to an efficient CP at $\Delta = 0$, however, the signal intensity is much smaller than the maximum signal achieved under rampCP. We also evaluated the efficiency of a ramped version of the COMPOZER-CP sequence (data not shown), however, our results suggest that ramping does not improve its efficiency. Nevertheless, the ramped version of the COMPOZER-CP sequence could be useful in situations where RF field inhomogeneity is very poor.



Fig. 4. Cross-polarization build-up curves for a static adamantane sample: (a) CH_2 peak and (b) CH peak. Crosses, open triangles, closed squares, and closed circles indicate the intensities obtained from CP, rampCP, COMPOZER-CP, and the supercycled COMPOZER-CP experiments, respectively. The signal intensities were normalized by the maximum intensity of the CH_2 or CH peak obtained in a CP.



Fig. 5. Performance of COMPOZER-CP against the Hartmann–Hahn mismatch under MAS ($v_r = 10 \text{ kHz}$) for an adamantane sample: CH₂ peak. Crosses, open triangles, closed squares, and closed circles indicate the intensities obtained from CP, rampCP, COMPOZER-CP, and the supercycled COMPOZER-CP experiments, respectively. The signal intensities were normalized by the maximum intensity of the CH₂ peak obtained in a CP. A contact time of 7.92 ms was used in all experiments. The solid, long-dashed, dashed, and dotted lines are for eye guidance.

4. Conclusions

Our experimental results suggest that the new pulse sequence, COMPOZER-CP, is capable of transferring Z-magnetization via heteronuclear dipolar couplings from high- γ to low- γ nuclei. The structure of the homonuclear dipolar coupling Hamiltonian is such that it can assist the spin diffusion under the COMPOZER spinlocks, which further enhances the uniformity of CP. Experimental results for adamantane under a static experimental condition confirm that the COMPOZER-CP is better in overcoming the HH mismatch than the rampCP. Our experimental results also infer that the contact time required for cross-polarization in COMPOZER-CP sequence is shorter than that of the rampCP sequence. We believe that the COMPOZER-CP sequence and its ramped version will be useful to study static samples, particularly semi-solids like lipid bilayers and liquid crystalline materials. Such application is in progress and will be reported elsewhere.

Acknowledgments

M. Fukuchi was supported by research fellowship of Global COE program, International Center for Integrated Research and Advanced Education in Material Science, Kyoto-University, Japan. We thank the Chemistry exchange program between Japan and USA, and the NIH, for the funding support to A.R.

References

- K. Schmidt-Rohr, H.W. Spiess, Multidimensional Solid-State NMR and Polymers, Academic Press, New York, 1994.
- [2] S.R. Hartmann, E.L. Hahn, Nuclear double resonance in the rotating frame, Phys. Rev. 128 (1962) 2042–2053.
- [3] A. Pines, M.G. Gibby, J.S. Waugh, Proton-enhanced NMR of dilute spins in solids, J. Chem. Phys. 59 (1973) 569–590.
- [4] (a) J. Schaefer, E.O. Stejskal, Carbon-13 nuclear magnetic resonance of polymers spinning at the magic angle, J. Am. Chem. Soc. 98 (1976) 1031–1032;
 (b) E.O. Stejskal, J. Schaefer, J.S. Waugh, Magic-angle spinning and polarization transfer in proton-enhanced NMR, J. Magn. Reson. 28 (1977) 105–112.
- [5] M.H. Levitt, D. Suter, R.R. Ernst, Spin dynamics and thermodynamics in solidstate NMR cross polarization, J. Chem. Phys. 84 (1986) 4243–4255.
- [6] (a) O.B. Peersen, X.L. Wu, I. Kustanovich, S.O. Smith, Variable-amplitude cross-polarization MAS NMR, J. Magn. Reson. A104 (1993) 334–339;
 (b) O.B. Peersen, X.L. Wu, S.O. Smith, Enhancement of CP-MAS signal by variable-amplitude cross-polarization. Compensation for inhomogeneous B₁ fields, J. Magn. Reson. A106 (1994) 127–131.
- [7] G. Metz, X. Xiaoling, S. Smith, Ramped-amplitude cross polarization in magicangle-spinning NMR, J. Magn. Reson. A110 (1994) 219–227.
- [8] (a) S. Hediger, B.H. Meier, N.D. Kurur, G. Bodenhausen, R.R. Ernst, NMR cross polarization by adiabatic passage through the Hartmann-Hahn condition (APHH), Chem. Phys. Lett. 223 (1994) 283–288;
 (b) S. Hediger, B.H. Maier, P.R. Frist, Adiabatic passage Hartmann, Hahn cross

(b) S. Hediger, B.H. Meier, R.R. Ernst, Adiabatic passage Hartmann-Hahn cross polarization in NMR under magic angle sample spinning, Chem. Phys. Lett. 240

(1995) 449-456:

(c) M. Baldus, D.G. Geurts, S. Hediger, B.H. Meier, Efficient ¹⁵N-¹³C polarization transfer by adiabatic-passage Hartmann-Hahn cross polarization, J. Magn. Reson. A118 (1996) 140-144;

(d) S. Hediger, P. Signer, M. Tomaselli, R.R. Ernst, B.H. Meier, A combination of slow and fast RF field modulation for improved cross polarization in solid-state MAS NMR, J. Magn. Reson. 125 (1997) 291–301.

- [9] (a) W.K. Peng, K. Takeda, M. Kitagawa, A new technique for cross polarization in solid-state NMR compatible with high spinning frequencies and high magnetic fields, Chem. Phys. Lett. 417 (2006) 58–62;
 (b) W.K. Peng, K. Takeda, Efficient cross polarization with simultaneous adiabatic frequency sweep on the source and target channels, J. Magn. Reson. 188 (2007) 267–274;
 (c) W.K. Peng, A. Samoson, M. Kitagawa, Simultaneous adiabatic spin-locking cross polarization in solid-state NMR of paramagnetic complexes, Chem. Phys. Lett. 460 (2008) 531–535.
- [10] A. Ramamoorthy, C.H. Wu, S.J. Opella, Experimental aspects of multidimensional solid-state NMR correlation spectroscopy, J. Magn. Reson. 140 (1999) 131–140.
- [11] (a) A. Bax, B.L. Hawkins, G.E. Maciel, Off-resonance cross-polarization: a technique to reduce rf power requirements for magnetization transfer experiments in solids, J. Magn. Reson. 59 (1984) 530–535;
 (b) S.C. Shekar, D.-K. Lee, A. Ramamoorthy, Chemical shift anisotropy and offset effects in cross polarization solid-state NMR spectroscopy, J. Magn. Reson. 157 (1996) 223–234;
 (c) S.C. Shekar, D.-K. Lee, A. Ramamoorthy, An experimental strategy to dramatically reduce the RF power used in cross polarization solid-state NMR spectroscopy, J. Am. Chem. Soc. 123 (2001) 7467–7468.
- [12] K. Takegoshi, C.A. McDowell, Cross polarization using a time averaged precession frequency. A simple technique to reduce radiofrequency power requirements for magnetization transfer experiments in solids, J. Magn. Reson. 67 (1986) 356–361.
- [13] S.V. Dvinskikh, K. Yamamoto, A. Ramamoorthy, Separated local field NMR spectroscopy by windowless isotropic mixing, Chem. Phys. Lett. 419 (2006) 168–173.
- [14] K. Yamamoto, S.V. Dvinskikh, A. Ramamoorthy, Measurement of heteronuclear dipolar couplings using a rotating frame solid-state NMR experiment, Chem. Phys. Lett. 419 (2006) 533–536.
- [15] H. Geen, J.J. Titman, H.W. Spiess, Heteronuclear magnetization transfer in rapidly spinning solids, Chem. Phys. Lett. 213 (1993) 145–152.
- [16] A.E. Bennett, C.M. Rienstra, M. Auger, K.V. Lakshmi, R.G. Griffin, Heteronuclear decoupling in rotating solids, J. Chem. Phys. 103 (1995) 6951-6958.
- [17] (a) U. Haeberlen, J.S. Waugh, Coherent averaging effects in magnetic resonance, Phys. Rev. 175 (1968) 453–467;
 (b) M. Mehring, Principles of High Resolution NMR in Solids, second ed., Springer, Verlag, Berlin, 1983.