We present a new technique that combines the versatility of magic angle spinning (MAS) nuclear magnetic resonance (NMR) spectroscopy with the superior sensitivity provided by very small detection coils. This opens the way for NMR studies of solid samples with nanoliter volumes. Furthermore, the very strong radio frequency (rf) fields that can be generated by these microcoils facilitate a much broader excitation bandwidth and/or decoupling efficiency.

Although solid-state NMR is the method of choice for investigating local structure, alignment, and dynamics of nonosoluble and noncrystalline functional materials, its feasibility for studying chemically and biologically relevant systems is often hampered by sensitivity. For sensitivity reasons, microcoil technology was introduced in liquid-state NMR for mass-limited samples and high-sensitivity. For sensitivity reasons, microcoil technology was introduced in liquid-state NMR for mass-limited samples and high-sensitivity. Small-volume MRI experiments, based on the observation that the efficiency of a solenoid coil scales inversely with its diameter. By reciprocity, this also allows the generation of very high rf fields with limited amount of power, which has been exploited for static wide-line NMR studies in solids.

Two important pillars underpin the success of solid-state NMR as an analytical tool in materials science: first, sample rotation about the so-called "magic angle" needed to average anisotropic contributions to the resonance lines and second, polarization transfer from abundant nuclei (e.g., protons) to less-abundant nuclei with low gyromagnetic ratios such as $^{13}$C or $^{15}$N. The evolution of a variety of techniques for studying internuclear distances, bond angles, molecular orientation and dynamics, spin diffusion, and chemical exchange processes, is built on these foundations.

Here we describe a microMAS design, featuring microcoil-based resonators with either 400/300 μm outer/inner diameter or 335/235 μm outer/inner diameter coils and sample holders down to 170/125 μm outer/inner diameter (10 mL sample volume), mounted on a regular commercially available MAS unit (Figure 1). The microcoil resonator is a solenoid coil integrated into a capacitor similar to that developed for static experiments leaving a 220 μm opening (30 mL internal volume) for the rotor. The advantage of this design is its mechanical stability and minimization of signal losses. Compared to regular millimeter-sized coils with only a few windings, our microcoil design with over 10 windings allows for a better $B_0$ homogeneity over the sample volume, thus improving efficiency. Using microcoils, susceptibility broadening can be a serious issue that limits the resolution. In the present design, this is not the case, as these effects are averaged by MAS. The resolution of most solids spectra is, therefore, determined by the intrinsic spin interactions in the materials. Low-rf-power operation combined with the fact that only a small homogeneous $B_0$ volume is needed may help to fulfill the promise of tabletop NMR equipment. Low-power requirements and the absence of frictional heating in combination with the advantageous surface/volume ratio, which results in better cooling efficiency, can alleviate heating problems in, for example, hydroxyl biological samples with high salt concentrations such as membranes. In the present contribution, we focus on the feasibility of using microcoil circuits to study mass-limited (bio)organic compounds using CP-MAS.

A critical step in the probe assembly is the alignment of the microcoil circuit with respect to the rotor axis. Nevertheless, we found spinning to be very stable without any sizable excursions of the rotation axis; as a result, the attainable spinning speeds are those imposed by the supporting rotor, being 2–15 kHz for the 4 mm design used here. The spinning axis was set to the magic angle using KBr in the latest design the 4-mm support rotor is fitted with a separate circuit tuned for $^{87}$Br observation. In this way, one can quickly set the spinning axis to the magic angle and immediately proceed with the measurement of interest without the need of changing the sample. This approach makes simultaneous acquisition of spectra from samples in both the 4-mm rotor and the microrotor possible, thus allowing for efficient use of instrument time.

A cross-polarization (CP) experiment was performed at 14.1 T (600 MHz) using 6 mL of 25% $^{13}$C-enriched glycine in the microrotor spinning at 8.2 kHz. The line width of the C$_2$-resonance is 117 Hz, identical to the line width in a commercial probe. This demonstrates that MAS can spin out residual line broadenings due to susceptibility effects. A spectrum with a S/N = 61.5 ($I_{\text{spin}}/I_{\text{noise}}$) was obtained in 10 800 scans. Although further work is needed to assess the exact sensitivity of the probe head, this shows the feasibility of getting CP-MAS spectra for minute sample quantities in a reasonable amount of time. A first assessment of the sensitivity shows that the circuit is not performing at the theoretical optimum yet, leaving room for further improvements in the circuit design. The microcoil circuit allowed the generation of a proton decoupling field (expressed as nutation frequencies ($\gamma B_0/2\pi$) in Hz) of 200 kHz with as little as 1.35 W of rf power. As the circuits are designed to handle hundreds of watts of power, one can generate rf fields well beyond the capabilities of commercial CP-MAS probes. This is shown for a uniformly $^{13}$C-labeled trialanine sample in antiparallel $\beta$-sheet form. Its CP-MAS spectrum obtained in the microrotor,
proof of principle, we studied a silk rod prepared from the silkworm Bombyx mori silk gland 10 with 20% $^{13}$C enrichment of the carbonyl groups. In a systematic study of $^{13}$C CSA tensors in peptides, an isotropic shift variation of 7 ppm was observed for the carbonyl region of a $^{13}$C CP-MAS spectrum of uniformly $^{13}$C-labeled trialanine in the form of silk I, random coil, and silk I, antiparallel $\beta$-sheet form) as a function of proton CW decoupling fields (Figure 2). Increasing the decoupling rf field strength up to 600 kHz, using the 335/235 $^{13}$C spin systems, shows better line separation at high decoupling fields (Figure 3). As a result of the narrowing of the resonances, which levels off at 500 kHz decoupling (Figure 3). 

At 4 kHz spinning (348 164 scans), the spinning sidebands are more prominent. The arrows indicate the spinning sideband pattern from the carbonyl resonance at 171 ppm, accompanied by two spinning sidebands. At 8.2 kHz spinning (108 000 scans), the spectrum is dominated by a 1 kHz broad carbonyl resonance at 171 ppm, accompanied by two spinning sidebands. At 4 kHz spinning (348 164 scans), the spinning sidebands are more prominent. The arrows indicate the spinning sideband pattern from the Vespel rotor material.

In conclusion, we have shown that high-resolution solid-state NMR can be miniaturized to study samples in the nanoliter regime. We implemented a viable microdesign that incorporates cross-polarization and MAS, which is an essential step to optimize high-resolution solid-state NMR for mass-limited samples. Furthermore, this microdesign can either operate at extremely low rf power or generate extremely high rf fields, helping to excite broad spectra and achieve highly efficient proton decoupling, thus, substantially improving spectral resolution.

Acknowledgment. The authors are very grateful to Prof. T. Asakura and Dr. K. Yamauchi (Tokyo University of Agriculture and Technology) for supplying the silk rod and trialanine samples and for their interest in this work. We thank Jan van Os and Gerrit Janssen for their technical support and discussions. Edwin Sweers is acknowledged for his (micro)mechanical craftsmanship.

Supporting Information Available: Materials, experimental NMR procedures, and glycine CP-MAS spectra (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

References


JA061350+