1	3D-printed microcell for protein NMR at high ionic strengths
2	and small sample volumes
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23 OD samples are needed to study samples in conductive, radiofrequency absorbing solvents 24 such as water. We demonstrate an effective and inexpensive alternative for reducing the 25 active sample volume to 0.13 mL by 3D printing of ellipsoidal shaped cells that are inserted 26 into 5-mm OD NMR tubes. Static magnetic susceptibility, χ , of printer resin was measured 27 using a simple slice-selection pulse sequence. We found that the χ of water increases 28 linearly with NaCl concentration, from -9.05 ppm to -8.65 ppm for 0 to 2 M NaCl. The χ 29 of D₂O was measured to be -9.01 ppm. The susceptibility difference between the resin ($\chi =$ -9.40 ppm) and water can be minimized by paramagnetic doping of the resin. Such doping 30 was found unnecessary for obtaining high quality protein NMR spectra when using 31 32 ellipsoidal shaped cells that are insensitive to susceptibility mismatching. The microcells 33 offer outstanding RF and good B_0 homogeneities. Integrated 600-MHz HSQC signal 34 intensities for the microcell sample in PBS buffer were 6.5±4% lower than for 0.5 mL of 35 the same protein solution in a regular 5-mm sample tube. The cell is demonstrated for N-36 acetylated α -synuclein in PBS buffer, and for observing tetramerization of melittin at 2 M 37 NaCl.

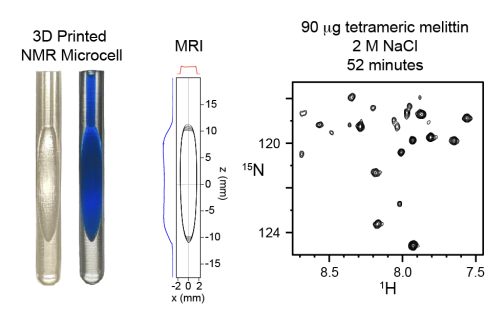
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41 **1 Introduction**

42 Nearly all solution NMR measurements are carried out using standard 5-mm outer diameter 43 (OD) sample tubes that require *ca* 0.5 mL of solvent to minimize "end effects" on magnetic 44 field homogeneity in the active volume of the sample. Use of a Shigemi cell, which consist 45 of glass with a magnetic susceptibility that is close to that of the selected solvent, can reduce the amount of sample required and thereby limit the cost of expensive protein 46 47 preparations. However, the tubes are expensive and need to be matched to the magnetic 48 susceptibility of the NMR solvent. The latter can be challenging considering that the 49 susceptibility of water is somewhat temperature-dependent (Schenck, 1996) and increases 50 substantially (becomes less negative) upon the addition of NaCl, whereas it increases by 51 ca 0.04 ppm in D₂O versus H₂O (see below). The sample volume also can be reduced by 52 inserting plugs, of a magnetic susceptibility close to that of the solvent, above and below 53 the active volume of the sample (Barbara, 2009).

Magnetic susceptibility measurements of ionic solutions are often carried out using a magnetic field that oscillates at frequencies ranging from ~50 Hz to low MHz (Tsukada et al., 2006;Gutiérrez-Mejía and Ruiz-Suárez, 2012). However, for magnetic resonance purposes it is the magnetic susceptibility measured in a static magnetic field that is relevant to distortions in field homogeneity (Sangal et al., 2023) and to our efforts to develop a small volume sample cell for protein NMR.

60 We describe a simple method for measuring magnetic susceptibility of solid material in a 61 high-resolution NMR spectrometer, and the development of a 3D printed microcell that 62 can be inserted into a regular 5-mm NMR tube. The resin that we used for 3D printing at high resolution (25 μ m) has a reported magnetic susceptibility of $\chi = -9.34$ ppm (Sangal et 63 64 al., 2023), which is well below that of water. However, by printing the cell with a spherical 65 (Hizawa et al., 2017) or ellipsoidal geometry, magnetic field homogeneity within the cell becomes insensitive to the susceptibility mismatch between solvent and the printer resin 66 67 (Schenck, 1996; VanderHart, 1996). The shape of our cell is perturbed by a narrow 68 diameter, cylindrical access port that is needed to fill it with the NMR sample. Magnetic 69 susceptibility mismatching effects, resulting from the deviation of a perfect ellipsoidal 70 shape caused by this access port and by the finite resolution of the printer, can be minimized 71 by paramagnetic doping of the printer resin with organic paramagnetic salt (Evans, 1959) but is found unnecessary for routine applications. The cells are reusable, but the cost ofprinting such cells is minimal and recycling them therefore may not be necessary.

74 We note that susceptibility mismatching for non-ellipsoidal shapes can also be obtained by 75 introducing nearby suitably shaped small "compensation structures" of different magnetic 76 susceptibility, that effectively shim the sample to homogeneity for microfluidic 77 applications (Ryan et al., 2014). Alternatively, the magnetic susceptibility of an aqueous 78 solution can be increased by addition of Eu³⁺-complexed diethyl-triamine pentaacetate (DTPA), which due to the short Eu^{3+} electron T₁ has minimal broadening effects on other 79 80 solutes (Hale et al., 2018) but is restricted to cases where the solvent susceptibility is more 81 negative than the surrounding material, which does not apply for 3D printer resin.

82 The microcell introduced by us for biological NMR spectroscopy purposes is particularly 83 useful when the available sample quantity is limited, or when high ionic strength is 84 required. High ionic strength lowers the quality factor (Q) of radiofrequency (RF) coils, 85 and thereby negatively impacts NMR sensitivity, an effect that scales steeply with 86 frequency (Ugurbil, 2018). RF penetration of water has been studied extensively for 87 applications to magnetic resonance imaging, where it impedes the observation of tissue far 88 from the body surface (Roschmann, 1987). RF absorption at high and ultrahigh magnetic 89 fields also challenges solution ¹H NMR spectroscopy, where the use of pulses that are 90 compensated for both offset and RF inhomogeneity (Freeman et al., 1980;Xia et al., 2017) 91 becomes essential, in particular for the vast majority of advanced experiments that include 92 multiple 180° pulses (Manu et al., 2023). Use of the 3D printed microcell greatly reduces 93 problems with probe detuning, lowering of Q, and RF absorption. Consequently, use of 94 the microcell results in short ¹Hpulse widths with superior RF homogeneity, even at 95 elevated ionic strength.

96 Measurements of magnetic susceptibilities of materials with application to magnetic 97 resonance imaging (MRI) as well as microfluidic NMR technology commonly have 98 employed MRI scanner equipment, and elegant methods applied to a wide range of 99 materials were presented by Wapler et al. (Wapler et al., 2014). The same approach was 100 recently used to measure magnetic susceptibility of 3D printed materials (Sangal et al., 101 2023), we demonstrate simple methods for deriving magnetic susceptibility differences 102 between ionic solutions and printer material in a high-field solution NMR magnet. We also

103	demonstrate that while using only 130 μ L of solvent, the sensitivity in common multi-
104	dimensional NMR experiments, such as the gradient-enhanced HSQC (Kay et al., 1992),
105	is comparable to what is obtained on a regular 500- μ L sample. In another application, we
106	show that high-quality spectra can be obtained for 90 μ g of recombinantly expressed and
107	chemically amidated melittin at 2 M NaCl concentration. Milligram-scale expression and
108	purification of this uniformly ¹⁵ N-enriched peptide in its post-translationally modified
109	state, which tetramerizes in a salt-dependent manner, is expensive and very labor-intensive
110	(Gelenter and Bax, 2023).
111	
112	2 Results and Discussion
113	
114	2.1 Measurement of magnetic susceptibility
115	For all materials pertinent to high-resolution solution NMR, the induced magnetization \mathbf{M}
116	depends linearly on the applied magnetic field H ₀ :
117	$\mathbf{M} = \chi \mathbf{H}_{0} \tag{1}$
118	where χ is the volume magnetic susceptibility, often expressed in units of parts per million
119	(ppm), i.e. χ is negative for diamagnetic media and positive for paramagnetic substances.
120	The magnetic flux density, \mathbf{B}_0 , is directly proportional to \mathbf{H}_0 :
121	$\mathbf{B}_{0} = \boldsymbol{\mu} \mathbf{H}_{0} \tag{2}$
122	where the magnetic permeability μ corresponds to $\mu = (1+\chi)\mu_0$, and $\mu_0 = 4\pi \ 10^{-7} \text{ H/m}$ is
123	the permeability of vacuum. In an NMR sample, the magnetic flux density corresponds to
124	the sum of the applied magnetic field \mathbf{H}_0 and the integral of the magnetic field contributions
125	from the induced magnetization over all sample volume elements at locations r relative to
126	the point of interest (Barbara, 1994). For a cylindrical sample of infinite length, with its
127	axis parallel to a homogeneous applied magnetic field, the integral over all space within
128	the cylinder is uniform across all locations, resulting in
100	
129	$\mathbf{B}_{0} = \mu_0(1+\chi) \mathbf{H}_{0} $ (3)

131 magnetic susceptibilities χ_1 and χ_2 . For the example of a Shigemi sample cell, where the

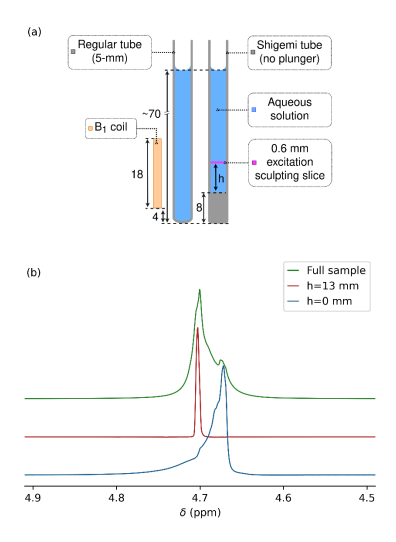
bottom segment of the cylindrical tube consists of glass with susceptibility χ_g , and χ_s is the solvent susceptibility above it (Figure 1a), the flux density at the interface is given by

134 $\mathbf{B}_{o} = \mu_{0} [1 + (\chi_{g} + \chi_{s})/2] \mathbf{H}_{o}$

Away from the interface, the integrated contributions to B_0 from volume elements below 135 136 and above the interface depend on the height above the interface as well as the transverse location, with values converging to $\mathbf{B}_0 = \mu_0(1 + \chi_s)\mathbf{H}_0$ at distances above the interface that 137 138 are large relative to the tube inner diameter, D. Therefore, when selectively observing 139 solvent signal with a chemical shift of δ_0 from a very thin slice perpendicular to the sample 140 axis at height h above the interface (Figure 1a), a narrow signal at a frequency $\delta(h) = [1 + 1]$ $(\chi_{g} + \chi_{s})/2]\delta_{o}$ is observed for $h \ll D$, with $\delta(h)$ approaching $(1 + \chi_{s})\delta_{o}$ for $h \gg D$, and a 141 142 complex line shape for slices taken at intermediate values (Barbara, 1994).

143 Hence, for a solution above the solid glass of a Shigemi cell with a total solvent height that 144 is large relative to both D and the height of the receiver coil, the line shape corresponds to $\delta(h)$ contributions ranging from $[1 + (\chi_g + \chi_s)/2]\delta_o$ to $(1 + \chi_s)\delta_o$. In other words, the total 145 width of the line shape at its base corresponds to half the difference between χ_g and χ_s in 146 147 units of ppm. The above analysis does not take into account the B_0 gradient inside the NMR tube that is caused by the large magnetic susceptibility mismatch at the bottom of the tube. 148 However, when assuming that the magnetic susceptibility of the Shigemi glass is much 149 150 closer to that of water than air, shimming on a water-filled sample tube inserted to the same 151 depth into the probe can be used to remove this B₀ gradient, to first order.

(4)





153 Figure 1. Measurement of magnetic susceptibility in a Shigemi sample tube. (a) Prior to 154 measurements on this sample, the magnetic field homogeneity was optimized for a regular 5-mm 155 NMR sample tube, filled to the same height of 70 mm above the bottom of the tube, and inserted 156 to the same depth into the probehead. Dimensions in mm. A thin slice (pink) through the sample 157 at height h above its bottom was selected by excitation sculpting. (b) HDO resonances for the 158 Shigemi sample tube containing 0.3% H₂O/99.7% D₂O (no plunger). The glass/liquid interface of 159 the Shigemi cell was positioned 5 mm below the center of the receiver coil, using the shim settings 160 of a regular 5-mm sample tube filled to the same total height with the same solution, inserted to 161 the same depth into the NMR probe (see left half of panel a). Overlayed are spectra recorded for 162 the entire sample (green) and for 0.6-mm thickness horizontal slices through the Shigemi cell, 163 centered at h = 0 mm (blue), and at h = 13 mm (red) by using excitation sculpting while applying z 164 gradients. Spectra were scaled to show the same intensity.

166 2.2 Measurement of magnetic susceptibility in a high-resolution NMR magnet

167 Careful measurements of solvent susceptibility in a high-resolution magnet that used the 168 gas phase of TMS as an internal reference have been reported by Hoffman (Hoffman, 169 2020;Hoffman, 2022). Here, we describe a different approach that also permits 170 susceptibility measurements of solid objects. First, we demonstrate the method for 171 measurement of the susceptibility of the glass bottom section of a Shigemi cell.

172 Bruker's topshim program was used to minimize magnetic field inhomogeneity of a regular 173 5-mm sample that contained a 1-mL solution of a variable NaCl concentration in 97% 174 $D_2O/3\%$ H₂O, that was inserted into the probe to have its bottom ~13 mm below the center 175 of the receiver coil. Then, without changing the shim settings, the HDO line shape on the 176 same solvent composition was observed for a Shigemi sample tube, inserted to the same 177 depth into the probehead. Because the Shigemi tube had an 8-mm bottom segment of solid 178 glass, the flat solvent-glass interface then is located 5 mm below the center of the receiver coil (Figure 1a). If χ_g were identical to χ_s , the same perfect line shape would be expected, 179 180 but with approximately 15% lower intensity because the bottom ~15% of the receiver coil 181 now was filled with solid glass from the Shigemi tube.

182 For the 97% D₂O/3% H₂O Shigemi tube sample, a pronounced upfield shoulder was 183 observed (Figure 1b, green), indicative of susceptibility mismatching. Excitation sculpting 184 (Stott et al., 1995) (Appendix A1) while applying a pulsed z-gradient was then used to 185 select a slice of 0.6-mm thickness, centered at the glass-solvent interface, i.e. selecting a 186 ~ 0.3 -mm solvent layer just above the interface. A resonance for this layer was observed 187 that was 0.032 ppm upfield from the most intense segment of the solvent obtained with a 188 30° pulse, without slice selection (Figure 1b). When selecting a slice at a height of 13 mm 189 above the interface, the signal (Figure 1b, red) coincided with the maximum of the 190 resonance obtained without slice selection. Therefore, the difference in frequency between the red and blue resonances provides a good measure for the difference between $(\chi_g + \chi_s)/2$ 191 192 and χ_s . The precision at which the frequency at the glass-water interface can be measured 193 is impacted by the slice thickness of the selected solvent layer, which is subject to lateral 194 gradients that increase with slice layer thickness and towards the edges of the slice. By 195 varying the position of the center of the slice from ~ 0.2 mm below the interface to ~ 0.2 196 mm above the interface, increased total intensity with a strong downfield shoulder is

197 observed. The upfield edge of this line shape (Figure 1b, blue) remains invariant to the 198 precise position of the selected slice and represents the true $(\chi_g + \chi_s)/2$ value, which can be 199 determined at a precision of ~0.02 ppm.

200 Repeating the same measurement but using 97% H₂O/3% D₂O and strongly mistuning the probehead to reduce radiation damping, showed a shoulder that was 0.0183 ppm closer to 201 202 the frequency observed 8 mm above the center of the coil (i.e. h = 13 mm; Appendix A2). 203 Accounting for the solutions not being fully deuterated or protonated, their difference in 204 static magnetic susceptibility then equals $\chi_{H2O}-\chi_{D2O}=2\times0.0183\times(100/94)=0.04$ ppm. The widely used literature value for χ_{H2O} is -9.05 ppm (Sangal et al., 2023), yielding $\chi_{D2O} = -$ 205 206 9.01 ppm, in fair agreement with Hoffman's measurements (Hoffman, 2022). Using γ_{D20} = -9.01 ppm as a reference, the susceptibility of the Shigemi glass used in our 207 measurements is $\chi_g = -9.08$ ppm. 208

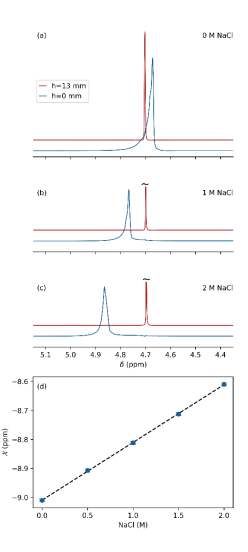


Figure 2. Effect of salt on magnetic susceptibility of 97% D₂O. Resonances shown correspond to a ca 0.6-mm thick slice through a Shigemi tube, centered at h = 0 (blue) and h = 13 mm (red) above the glass bottom of the Shigemi sample cell, using the same protocol as for Figure 1b at (a) 0 mM NaCl; (b) 1.0 M NaCl; (c) 2.0 M NaCl. (d) Magnetic susceptibility of D₂O as a function of NaCl concentration, using $\chi_{D2O} = -9.01$ ppm as a reference. The intensities of the slices at h = 0 are upscaled *ca* 16-fold to account for the 50% smaller aqueous volume and the strong magnetic field inhomogeneity in these lower slices.

217 2.3 Salt dependence of water magnetic susceptibility

The effect of dissolved NaCl on magnetic susceptibility of water is important to protein NMR. We therefore repeated the above measurements of the 97% D₂O/3% H₂O Shigemi tube sample after addition of 0.5, 1, 1.5, and 2 M analytical grade NaCl (Sigma-Aldrich) to the solvent (Figure 2a-c), showing a linear increase in solvent susceptibility with salt 222 concentration over this range: $\chi_{D2O+NaCl} = ([NaCl] \times 0.2 - 9.01)$ ppm, where [NaCl] denotes 223 the concentration molarity in units of moles per liter (Figure 2d).

224 2.4 Susceptibility and paramagnetic doping of 3D printer resin

225 Clear V4 resin was used because it enabled the highest precision (25-um resolution) of 226 printing on a Formlabs Form3+ 3D printer available in our laboratory, and its optical 227 transparency facilitated sample handling for 3D printed microcells. The susceptibility of 228 the printed Clear V4 resin was measured in the same manner as described above for the 229 Shigemi tube. A cylindrical plug of 2 cm length was printed and pushed to the bottom of a standard 5-mm NMR sample tube that was prefilled with 0.7 mL 1% H₂O/99% D₂O, 230 231 such that the top of the plug was again 5 mm below the center of the receiver coil once 232 inserted into the magnet. Shimming of the magnetic field was carried out on a sample 233 without the plug, filled to the same level by using $\sim 1 \text{ mL}$ of the same solvent, inserted into 234 the probehead at the same depth, i.e. with the bottom of the sample tube at 25 mm below 235 the center of the RF coil.

236 The frequency difference observed at the D_2O /resin interface versus the top of the coil was 237 -0.195 ppm (Figure 3a), corresponding $\gamma_{ClearV4} = -9.40$ ppm, which is close to the value 238 of -9.33 ppm measured for this resin by magnetic resonance imaging (Sangal et al., 2023). 239 Considering that a mismatch in magnetic susceptibility between the sample cell and the 240 solution impacts the achievable B₀ homogeneity, and that $\chi_{\text{ClearV4}} < \chi_{\text{H2O}}, \chi_{\text{D2O}}$ increasing 241 the value of χ_{ClearV4} by paramagnetic doping of the resin in principle allows elimination of 242 this difference. Finding a paramagnetic doping substance that is miscible with the printer 243 resin and does not impact the performance of the 3D printer proved challenging. For 244 example, the use of concentrated CuCl₂ in methanol strongly impacted the polymerization 245 kinetics The same problem was encountered for a range of strong paramagnetic chelated 246 substances, such as gadodiamide (Omniscan), which is commonly used in protein 247 paramagnetic relaxation enhancement measurements and as a contrast agent in magnetic 248 resonance imaging. However, the hydrophobic cobalt(II) complex, cobalt(II) bis(2-249 ethylhexanoate), available from Sigma-Aldrich as a 65 wt. % solution in mineral spirits

- 250 (product number: 444545), proved miscible with the Clear V4 printer resin without major
- adverse impact on print quality.

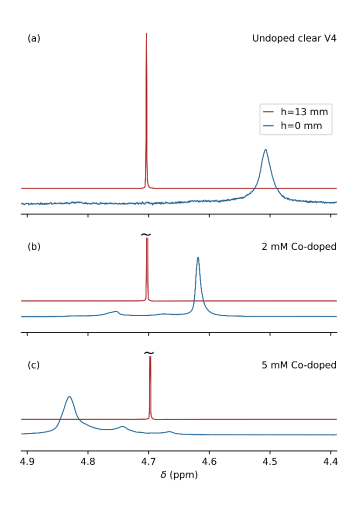




Figure 3. Effect of cobalt(II)2-ethylhexanoate doping on the magnetic susceptibility of printed Formlabs Clear V4 resin. A solid plug of printed resin that is 2 cm in length and has a 4.0-mm outer diameter (OD) was inserted into a regular 5mm OD NMR sample tube (Wilmad-507; ID 4.2 mm), prefilled with 0.7 mL 99%D₂O/1%H₂O. Overlaid spectra are shown from 0.6-mm thickness slices centered at the top of a Clear V4 plug (blue), and 13 mm above the plug (red). (a) No doping; (b) 2 mM cobalt(II) doping; (c) 5 mM cobalt(II) doping.

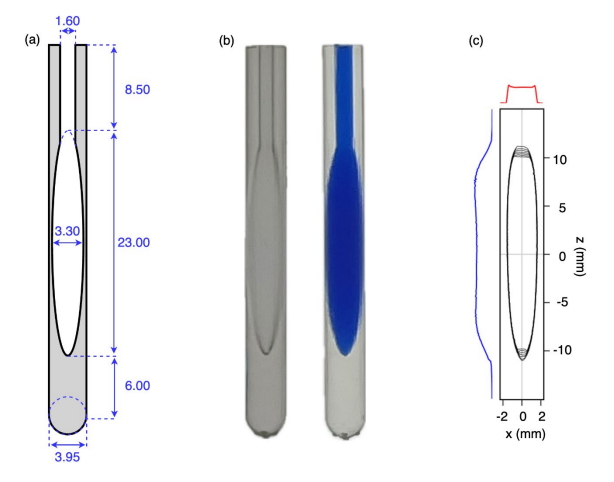
Comparison of the difference in HDO resonance frequencies obtained for slices at 13 mm above the interface between solvent and printed plug and at the interface for three different levels of the cobalt(II) doping, 0 mM (Figure 3a); 2 mM (Figure 3b) and 5 mM (Figure 3c), shows a doping-dependent decrease from +117 Hz to -80 Hz for the NaCl-free 99% D₂O sample. This result indicates that it is possible to match the susceptibility of the solvent to that of the printed resin. However, that would require a large number of printed cells to 265 cover the ionic strength range from 0 to 2 M salt, while also accounting for the difference
266 between D₂O and H₂O samples.

267 In practice, printing sample cells with different levels of cobalt(II) doping is labor-intensive 268 because it requires thorough cleaning of the printing vat used by the Formlabs 3D laser 269 printer. Printing with doped resin requires mixing of the viscous resin at the molecular level 270 with the doping agent and keeping it homogenous during printing. The latter also required 271 some modification of the printer to prevent refreshing of the printer resin with undoped 272 resin from a sealed cassette during printing. We therefore resorted to printing the sample 273 cells with an ellipsoidal shape, that to a good approximation were insensitive to the 274 susceptibility mismatch between solvent and printed resin (Schenck, 1996; VanderHart, 275 1996).

276 2.4 Performance of a Clear V4 ellipsoidal microcell

277 For applications to proteins, we settled on an ellipsoidal microcell design with a volume of 130 µL (Figure 4a). A printed access channel of 1.6 mm diameter (measured 1.3 mm) and 278 279 8.5-mm length and a volume of $\sim 11 \mu$ L was used for cleaning of the sample after the initial 280 print and prior to further hardening of the resin in a Formlabs light chamber (see Methods). 281 A subsequent overnight rinse with Milli-Q H₂O at 60 °C was used to remove small water-282 soluble contaminants. Comparison of the NMR spectrum of these impurities with those in 283 a sample obtained by briefly vortexing a mixture of unpolymerized resin and water followed by entering the aqueous phase into an NMR sample tube (Figure A3) suggests 284 285 that these impurities consist of residual, methacrylate-based small oligomeric species that 286 apparently can diffuse from the cell walls into the aqueous contents of the cell. Leaving 287 the sample cells filled with H₂O for a week prior to a final rinse reduces the impurity levels 288 further, but in our experience is not necessary considering we have not observed any 289 interaction between these very low impurity concentrations and isotopically enriched 290 proteins. For applications to samples in D_2O , leaving the microcell filled with D_2O for 24 291 h, prior to using it, reduces the intensity of a weak, very broad ($\sim 1 \text{ kHz}$) signal at $\sim 3.8 \text{ ppm}$, 292 that results from H₂O diffusing into the resin.

Because the digital printer increases the size of printed parts by a small amount due to partial polymerization adjacent to the laser-selected spots, the printed walls of the chamber are actually slightly thicker than designed, and the total volume of the cell including its access channel was measured gravimetrically to be 130 μ L (Figure 4b).

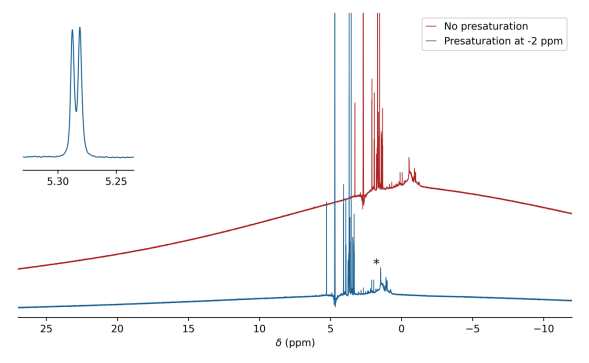




299 Figure 4. Images of the printed microcell. (a) Technical drawing; note that the actual dimensions 300 of printed material are slightly larger due to polymerization of a thin (~0.15 mm) sticky surface 301 layer that remains on the printed cell prior to subsequent UV curing. (b) Photographs of the 302 printed cells (left) prior to and (right) after filling with a blue dye solution; (c) Contour plot of a 303 sagittal xz cross-section through a 3D image of a 97% D₂O, 3% H₂O sample containing 1.8 mM 304 CuCl₂ to shorten the ¹HT₁ value to ~0.7 s. The image was recorded in absorption mode on an 800-305 MHz NMR spectrometer equipped with a 3-axis pulsed field gradient probehead. The total 306 measurement time was 9 minutes.

A sagittal (xz) cross-section through the absorption mode 3D image of the cell, recorded on a Bruker Neo 800-MHz instrument equipped with a 3-axis pulsed field gradient probehead, yielded a shape that matched the ellipsoidal design (Figure 4c) but that did not 310 include the access channel because the solvent in that channel falls outside the RF coil. 311 Small distortions near the bottom of the image correspond to the drop off in RF coil 312 receptivity. Distortions at the top of the sample originate from the access channel which 313 also distorts the ellipse. However, cross-sections taken through the 3D image along the x 314 and z axis through the center of the sample, plotted along the sides of the 3D image, show 315 the expected nearly rectangular shape, indicative of linear imaging gradients.

316 ¹H non-spinning linewidths of \sim 1 Hz at half height (600 MHz) obtained with the cell 317 (expanded anomeric doublet in Fig.5) were slightly larger than those obtained for a regular 318 5-mm NMR sample in the same probehead. With a width of only ~13 Hz at 0.55% of the 319 HDO peak height the lineshape is also very good, which is most important for protein NMR 320 studies, where protein ¹H line widths at half height commonly exceed 10-20 Hz due to fast 321 transverse relaxation and the absence of a "hump" in the water line shape is critical for 322 good solvent suppression.





323 324 Figure 5. ¹H background of printed microcell. The regular ¹H NMR spectrum of a 1 mg/mL solution 325 of sucrose in 99% D_2O , recorded without presaturation (red) and with ¹H presaturation at -2 ppm (blue), using a 100 Hz RF field strength to suppress the ¹H background of the solid resin. 326 327 Resonances from small impurities released from the printed cell when it has only briefly (~ca 2 h) 328 been rinsed with H_2O are marked with an asterisk. However, a subset of these resonances very 329 slowly reappear over a period of multiple days, at concentrations much lower than seen in the

figure (see Figure A3). For display purposes, the not-presaturated spectrum has been shifted upfield by 2 ppm and offset vertically. The inset shows the splitting for the sucrose anomeric doublet at 5.39 ppm.

333

We note that sometimes microscopic air bubbles can form inside the cell after filling it with the NMR sample solution. When this happens, it has a very strong adverse impact on both lineshape and linewidth. To eliminate the potential presence of such air bubbles, we briefly $(\sim 20 \text{ sec})$ spin the filled sample cell in a speedvac, operating at a pressure of *ca* 150 Pa, and replenish any lost volume by adding a few μ L of the protein solution to the access channel.

340 For observation of simple one-dimensional ¹H spectra, without echo delays, the protons of 341 the resin yield a strong background signal that is broad due to the rapid transverse 342 relaxation of this solid material. This background signal is readily suppressed by spin echo delays prior to the start of signal acquisition, as are already present in nearly all protein 343 344 NMR experiments. It can also be effectively reduced by presaturating this broad background with a weak RF field outside the spectral region of interest. Saturation with a 345 346 100-Hz RF field, applied at -2 ppm in the ¹H spectrum, attenuates the background signal 347 by about six-fold (Figure 5).

348 The ¹H RF field homogeneity was compared for the microcell and for a Shigemi sample (straight wall) containing 280 µL of phosphate buffered saline (PBS) solution in 99% D₂O. 349 350 Due to the substantial detuning of the 600 MHz cryoprobe used for this work by the ionic 351 solution in the Shigemi sample, the RF power for this sample was increased by 2.7 dB over the power used for the microcell to yield the same 90° pulse width of ~17 μ s. Comparison 352 of the decay of the signal when the excitation pulse is increased from 0.2 to 497.7 µs using 353 354 the Bruker "paropt" module (Figure 6) shows slightly better RF field homogeneity for the 355 smaller 3D printed microcell than for the Shigemi sample. Notably, the intensity of the 356 signal after a 90° pulse was only ~10% lower for the printed microcell than for the Shigemi 357 tube that contained more than double the volume of the same solution.

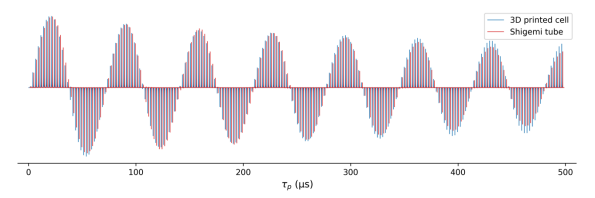


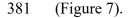


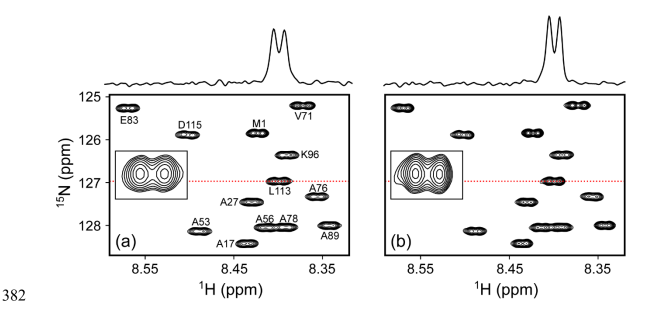
Figure 6. Comparison of ¹H RF field homogeneity in a 280- μ L Shigemi sample cell (red) and in the 360 3D printed 130- μ L 23×3.3-mm ellipsoidal microcell (blue). Signal intensity is shown as a function 361 of ¹H pulse duration, ranging from 0.2 to 497.7 μ s, generated by the Bruker macro 'paropt'. Both 362 samples contained the same solution of PBS buffer in 99% D₂O. The RF power for the microcell 363 sample was adjusted 2.7 dB lower than for the Shigemi sample to equalize the 90° pulse lengths.

365 2.6 Observation of high-resolution protein NMR spectra

366 The high resolution and sensitivity obtained with the microcell are illustrated for two proteins, N-acetylated α-synuclein and native C-terminal amidated melittin in its 367 monomeric and tetrameric forms. Complete or nearly complete N-acetylation of α -368 synuclein is invariably present in mammalian cells (Bartels et al., 2011) and strongly 369 370 impacts its interaction with phospholipids (Kang et al., 2012; Maltsev et al., 2012). By 371 simultaneously including a plasmid for expressing the NatB complex, needed for 372 acetylation of α -synuclein, together with a plasmid for α -synuclein, fully N-acetylated 373 protein can also be obtained from bacterial expression systems (Johnson et al., 2010). 374 Although this combined expression reduced protein yields in our hands, it enabled the pivotal incorporation of stable isotopes, such as ¹⁵N, in the biologically relevant state of 375 376 the protein.

377 As can be seen, in *ca* 20 min, a high-quality gradient-enhanced ¹H-¹⁵N HSQC spectrum 378 was obtained for 130 μ g of N-acetylated α -synuclein (14.5 kDa) in PBS buffer, pH 6.5, 379 when using the microcell, which approaches the sensitivity and resolution that was 380 obtained for 500 µg protein in a regular 5-mm sample cell, using a 0.5 mL sample volume

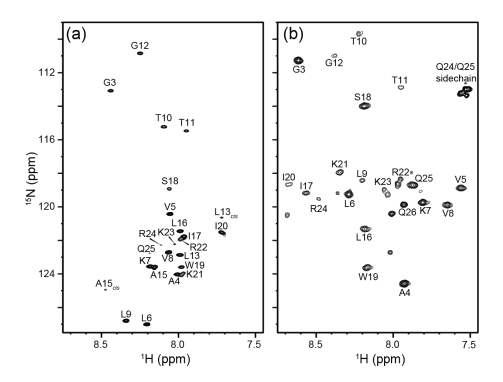




383 **Figure 7.** Comparison of small regions of the 600-MHz 1 H- 15 N HSQC spectra of 70 μ M N-acetylated 384 α -synuclein at 20 °C in PBS-buffer, pH 6.5. Each spectrum results from 175* × 4000* data points 385 with two transients per FID, for total measuring times of 21 min each. (a) 130 μ L in the 3D printed 386 microcell; (b) 500 µL in a standard 5-mm NMR tube. Lowest contours are drawn at the same level 387 above the respective RMS noise. Insets show the expansion of the L113 cross peak, with cross-388 sections through L113 above the panels at locations marked by the red dotted lines. The 389 comparison shows slightly lower resolution of the ${}^{1}H^{N-1}H^{\alpha}$ doublets for the microcell, reflected in 390 12% lower S/N versus 6.5% lower peak integrals relative to RMS noise.

391 The utility of the microcell becomes even more compelling for the study of the 392 tetramerization of native melittin, which requires C-terminal amidation. The latter involves 393 a large number of chemical derivatization and purification steps (Gelenter and Bax, 2023), 394 making it very challenging to generate adequate quantities of peptide for driving it to its 395 tetrameric state within a standard 5-mm NMR tube. Tetramerization is promoted by 396 increasing the NaCl concentration, but the tuning of the RF circuitry in cryoprobes often 397 limits salt concentrations to be below ca 0.5 M. Here, we demonstrate that the microcell 398 enables observation of the monomer-tetramer equilibrium at salt concentrations of 2 M, 399 and that indeed even at a moderate peptide concentration of 250 μ M the peptide is fully 400 tetrameric in the presence of 2 M NaCl (Figure 8). Interestingly, residues L9-R24, close 401 to the center of the peptide, are attenuated by an exchange process that is much less 402 prevalent at low ionic strength and high peptide concentration (Gelenter et al., 2024). We 403 speculate that the line-broadening associated with these weaker resonances arises from the 404 exchange process between the asymmetric arrangement of the two dimers seen in its X-ray 405 tetrameric structure (Terwilliger and Eisenberg, 1982) switching from fast exchange at 406 lower ionic strength to intermediate exchange at 2 M NaCl.

407 Notably, the same microcell was used for the two melittin spectra. After recording of the
408 low-ionic strength spectrum, contents of the microcell was removed and used to dissolve
409 15.2 mg NaCl, prior to insertion of the high ionic strength sample into the original cell,
410 thereby demonstrating the recyclability of the tube.



411

412 Figure 8. 800-MHz ¹H-¹⁵N HSQC spectra from 90 μg (250 μM) of ¹⁵N-labeled native melittin in a 413 130-µL microcell. Both spectra were collected at 288 K in 10 mM sodium phosphate buffer, pH 414 7.0, containing 3% D₂O. (a) Sample containing no NaCl, where melittin remains an intrinsically 415 disordered monomer. $L13_{cis}$ and $A15_{cis}$ correspond to residues in monomers with P14 in the 416 cis conformation. The data was collected with 2 transients per FID, using 200* points in the 417 indirect dimension corresponding to an evolution of 103 ms and a total measurement time of 27 418 min. (b) 15.2 mg of NaCl was added to the sample from (a) to reach a final concentration of 2 M 419 NaCl. Under these conditions melittin adopts an α -helical tetrameric conformation. The data was 420 collected with 8 transients, using 100^{*} points in the indirect dimension ($t_{1,max}$ = 51 ms) and a total 421 experimental time of 52 min.

422 **3 Concluding remarks**

3D printing enables the efficient and relatively inexpensive creation of complex, customized products with minimal waste. It is extensively used for prototyping new designs of items with intricate or complex geometries and has broad impact in science and engineering. 3D printing also enabled the design and construction of high performance solid-state NMR probes, offering similar or improved filling factors due to the coil being in close proximity to the sample, resulting in high RF transmit and receive efficiencies (Long et al., 2021;Pereira et al., 2023).

In solution NMR spectroscopy, 3D-printed bioreactor platforms were introduced that are compatible with low-field NMR spectrometers that accommodate bioengineered 3D cell models (Mangas-Florencio et al., 2025). That work consisted of a bioreactor made of biocompatible materials and included a microfluidic system for optimization of cell culture conditions during the actual NMR data collection process.

The application of 3D printing to high-field solution NMR spectroscopy has remained 435 436 rather limited, largely due to the requirements of high magnetic field homogeneity and 437 minimal background signals. Our study demonstrates that the homogeneity requirement 438 can be met by printing small sample cells with an ellipsoidal shape. The half-height line 439 widths achievable for our microcell is *ca* 1 Hz and remains limited by the precision at 440 which the cell's surface can be printed. For shimming purposes, we first used Bruker's 441 topshim program to adjust field homogeneity to starting values on a regular solution NMR 442 sample filled to the same height (~40 mm) as the length of the microcell. Subsequently, 443 after entering the microcell into the magnet, we used topshim followed by iterative tuning of z, z^2 , x, y, xz and yz gradients. The microcells can easily be recycled as they slide in 444 445 and out of standard high-quality Wilmad-507, New Era NE-HP5, or Norell Standard Series 446 5-mm NMR tubes. The access channel of the microcell is sufficiently small that surface 447 tension prevents the aqueous solution from leaving the microcell when the 5-mm NMR 448 tube is fully inverted while the microcell slides out of it.

Although inexpensive, sample cells are easily recycled as highlighted for the melittinsample, where 15.2 mg of salt was added to the initial sample by first removing the sample

451 solution, dissolving the NaCl, and re-injecting the solution into the original cell, all with 452 minimal losses. It is advisable to briefly spin and expose the sample to vacuum to remove 453 dissolved gasses, in particular for lengthy experiments. Formation of even microscopic air 454 bubbles deteriorates homogeneity for the microcell sample more than for the Shigemi or 455 larger conventional NMR sample tubes.

456 The sample cell appears unsuitable for the use of organic solvents which dissolve and 457 release resin components, resulting in strong narrow background signals. Even when using 458 water as the solvent, slowly increasing signals from micromolar quantities of resin-derived 459 small molecules appear in the ¹H spectrum over a period of days (marked by an asterisk in 460 Figure 5). However, the standard use of isotope-enriched multi-dimensional multinuclear 461 NMR experiments keeps these resonances well below the signal-to-noise threshold level, 462 and unless such contaminants have a strong interaction with the protein studied they have 463 no effect on the acquired spectra. We note that commercial 3D printing resins are 464 formulated in a way that tries to meet various performance specifications, such as material 465 strength, printing resolution, and printing speed. As such, they may not be optimal for high 466 precision work such as required for protein NMR. Future development of resins is needed 467 to allow high precision printing while minimizing release of small, incompletely 468 polymerized precursors into the aqueous phase.

469 While a small amount of solvent (<0.1%) can diffuse into cavities of the polymerized resin, 470 no detectable loss of solute signal was observed. For example, the intensity of the sucrose 471 NMR resonances remained unchanged, to within $\pm 0.2\%$ over a duration of more than one 472 week.

473 We cleaned the microcells by soaking them overnight at 60 °C in Milli-Q water and 474 subsequent removal of most of the solvent with a standard gel micropipette tip that didn't 475 quite reach the bottom of the cell, followed by upside down centrifugation after insertion 476 into an Eppendorf tube to remove the remainder of the solvent. An additional rinse with 477 130 μ L D₂O, followed by centrifugation and vacuum exposure can be used to remove any 478 residual solvent protons if the sample is intended for measurements in highly deuterated 479 D₂O. These are the most labor-intensive steps in preparing the sample cells, but limited 480 quantities of unrinsed sample cells are available upon request.

482 **4. Methods**

483 4.1 Magnetic susceptibility measurements

484 Selective excitation while applying a 11.5 G/cm (20% on the Bruker Neo-600 instrument) 485 was used for collecting the HDO solvent resonance measurements (Appendix A1) of 0.6-486 mm thickness slices at various heights, h, above the flat interface between a solid printed 487 plug and the aqueous solvent. The printed plug with outer diameter of 4-mm and a length 488 of 20-mm, including its hemispheric bottom, was inserted and pushed to the bottom of a 489 regular NMR tube, prefilled with 0.7 mL of 97% D₂O or 97% H₂O solution. A similar tube 490 without the plug was filled to the same height with the same solution and used for shimming 491 the magnetic field using topshim prior to inserting the sample with the plug at its bottom, 492 where the same shim settings of the sample without the plug were used. The thickness of 493 the aqueous fraction of the slice collected for h=0 was two-fold smaller than for slices at 494 h>0.3 mm, with a correspondingly lower integrated volume. The frequency of slices 495 collected at heights $>\sim 10$ mm above the plug became essentially independent of h. The 496 slice collected at *h*=0 shows extensive line broadening due to the large field gradient at the 497 solvent-plug interface (Figure 2a-c). The difference in ppm frequency was used as a 498 measure for $(\chi_{solvent} - \chi_{resin})/2$.

499 4.2 Printing of the microcell

500 Cells were printed in Clear V4 resin on a Form 3+ printer at 25-micron resolution to achieve 501 a smooth surface finish. The designs were created in OpenSCAD (2021.01) using standard 502 STL export settings, then prepared for printing in PreForm (3.43.2). Models were oriented 503 so that the bottom of each cell faced downward on the build platform, with 0.2 mm 504 touchpoint supports attached only at the bottom of the cells. The cells were washed with 505 15 mL isopropyl alcohol (IPA) at a rate of 5 mL/min through a syringe needle (0.58 mm 506 ID) with a blunt tip, connected to a Fast Protein Liquid Chromatography (FPLC) pump. 507 Cells were kept vertical during printing and subsequent UV-curing using Form Cure for 16 508 hours at 60 °C. Subsequently, cells were filled with Milli-Q water and immersed in a water509 filled falcon tube that was heated at 60 °C for 12 hours to remove water-soluble 510 components. After this cleaning, to further minimize the amounts of resin-derived 511 impurities from leaching from the cell wall into the solvent, which occurs at a rate that 512 decreases steadily with time, cells can be stored filled with and immersed in water for a 513 week at room temperature. After removal of the water from the cells by pipetting followed 514 by centrifugation upside down in Eppendorf tubes, they were briefly dried under vacuum. 515 Subsequent use of such a cell showed strongly reduced intensities of these impurity signals 516 (Figure A3).

517 The paramagnetically doped resin was prepared by mixing Formlabs Clear V4 Resin with

518 a cobalt(II) bis(2-ethylhexanoate) solution (65 wt.% in mineral spirits, Sigma-Aldrich,

519 444545) on a shaker at 37 °C and 200 rpm for 30 minutes. The plugs printed from doped

- 520 resin were washed with IPA and cured at 60 °C for 16 hours using Form Cure.
- 521

522 4.3 NMR sample preparation

523 The microcells were filled to the top with the sample solution (\sim 130 µL) using a gel-tip 524 pipette and degassed at *ca* 150 Pa pressure for 20-30 seconds using a SpeedVac (Savant, 525 SVC-100-H). After degassing, the cells were topped off with *ca* 2 µL of additional sample 526 solution and inserted into standard Wilmad-507, New Era NE-HP5, or Norell Standard 527 Series 5-mm NMR tubes.

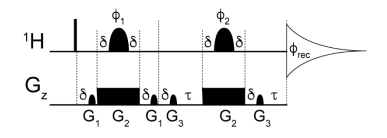
528 4.4 Imaging of the microcell

529 Although MRI normally uses absolute value mode displays, higher resolution absorption 530 mode spectra can also be obtained (Bretthorst, 2008). For generating images of the 531 microcell, we used a very simple one-pulse sequence with variable durations of the x, y, 532 and z gradients for encoding the three spatial dimensions (Appendix A4). The experiment 533 used Rance-Kay quadrature selection (Palmer et al., 1991;Kay et al., 1992) in both the x 534 and y dimensions, by collecting four scans per hypercomplex time domain data point 535 (ns=1). NMRPipe (Delaglio et al., 1995) processing of the 3D time domain matrix was 536 used to generate regular, amplitude-modulated quadrature (States et al., 1982) in both 537 indirect dimensions (see Appendix A5 for NMRPipe processing script). A total of $20^{*}(x)$ ×

- 538 $20^{*}(y) \times 512^{*}(z)$ data points were collected with a total acquisition time of *ca* 9 minutes
- 539 for a sample that contained 1.8 mM CuCl_2 in 99% D₂O.

541 Appendix A: Supplementary data and code

- 542
- 543 A1: Pulse diagram used for slice selection

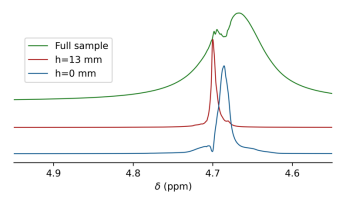


544

545 Figure A1. Diagram of the slice selection pulse sequence used to measure magnetic 546 susceptibility. The narrow bar represents the 90 ¹H excitation pulse. The shaped pulses ϕ_1 and 547 ϕ_2 have a profile of Reburp (Geen and Freeman, 1991) with a duration of 2 ms and a variable 548 offset frequency, ranging from ca + 23 to -30 kHz from the water signal resonating at 4.7 ppm. 549 The gradient pulses are either sine-bell shaped or rectangular, with durations of 0.5, 2.1 and 550 0.5 ms and a strength of 2, -11.5 and 3.5 Gauss/cm for G_1 , G_2 and G_3 , respectively. Delays: δ 551 = 50 µs and τ = 60 µs. The following phase cycling scheme was used: ϕ_1 = y, y, y, y, -x, -x, -x, -x, 552 -y, -y, -y, -y, x, x, x, x; $\phi_2 = x$, y, -x, -y; $\phi_{rec} = x$, -x, x, -x, -x, x, -x, x. The interscan delay was set to 4 553 s, number of scans to 4, spectral width to 8620 Hz and the acquisition time to 2 s.

554

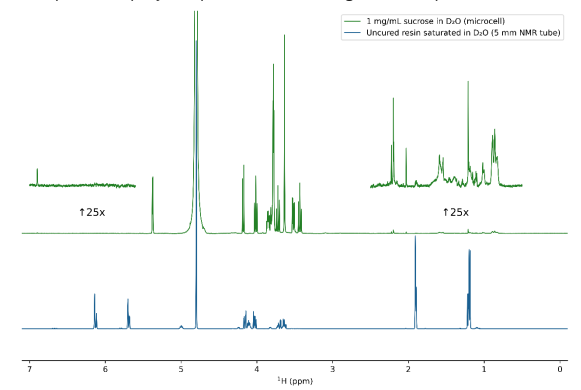
555 A2: Spectra used for measuring $\chi_{glass} - \chi_{H2O}$



556 557

Figure A2. Same as Figure 1b (main text), but for a solution containing 97% H₂O/3% D₂O. The

- resonance of the full Shigemi tube sample (green) is broadened by radiation damping and shown together with slices taken at h = 0 mm (blue), and at h=13 mm (red). The center of the
- 560 green resonance is shifted relative to its regular position due to radiation damping effect in
- the deliberately mistuned probe (Torchia, 2009).



562 A3: Spectra of polymer precursors and oligomeric impurities 563

Figure A3. ¹H NMR spectra (500 MHz) of liquid Formlabs Clear V4 resin, saturated in D₂O (5) 566 mm tube, 128 scans; blue trace), and 1 mg/mL sucrose in D₂O recorded in a 3D-printed 567 sample cell, 48 h after preparation (512 scans; green trace). The blue spectrum is scaled 568 down 10-fold relative to the green spectrum and corresponds to ca10 mg/mL precursor 569 concentration. The dissolved oligomer impurity concentration in the 3D-printed microcell 570 sample corresponds to *ca* 24 μ g mL⁻¹. Insets correspond to the olefinic and aliphatic 571 regions, upscaled 25-fold. According to the Formlabs Clear V4 datasheet, the monomeric 572 precursors in the blue spectrum correspond to methacrylic acid monoester with propane-573 1,2-diol and 7,7,9(or 7,9,9)-trimethyl-4,13-dioxo-3,14-dioxa-5,12-diazahexadecane-1,16-574 diylbismethacrylate (see https://formlabs-media.formlabs.com/datasheets/1801037-SDS-575 ENEU-0.pdf).

576 A4: Bruker pulse program used for imaging the microcell:

```
577
      #include<Avance.incl>
578
      #include<Grad.incl>
579
      #include<De.incl>
580
581
582
      "d11=30m"
583
584
      "in0=inf1".
                  ;125u
585
      "in10=inf2"
                   ;125u
586
587
      "d0=0"
588
      "d10=0"
589
590
      define list<gradient> EA2 = \{ 1.000 - 1.000 \}
591
592
      aqseq 321
593
594
595
     1 ze
596
       1m
597
     2 dll groff
598
       10u BLKGRAD
599
       10u pl9:f1
                                  ;set to 1000dB unless presat is needed
600
       10u fq=cnst2(bf ppm):f1 ;chemical shift for the resin to be
601
     presat'ed
602
      d1 cw:f1
603
       10u do:f1
604
       10u UNBLKGRAD
605
       p20:gp20. ;3m at 33% of z-grad
1m fq=0:f1 ;shift carrier back to be on-resonance with water
606
607
       1m pl1:f1
608
       (p1 ph1):f1 ;20 degree flip angle
lu gron1*EA ;x-grad 8%
       (p1 ph1):f1
609
610
       d0
611
       lu groff
612
       lu gron2*EA2 ;y-grad 8%
613
614
       d10
615
        lu groff
616
       600u
                      ;delay to dephase a very broad hump
617
       1u gron3
                     ;z-grad -1%
618
       go=2 ph31
619
       dll groff mc #0 to 2
620
       F1EA(calgrad(EA), caldel(d0)) ;TD1=40
621
       F2EA(calgrad(EA2), caldel(d10)) ;TD2=40
622
       d11 BLKGRAD
623
      exit
624
625
626
     ph1=0
627
     ph31=0
```

```
629
      #!/bin/csh
630
631
      bruk2pipe -verb -in ./ser \
632
        -bad 0.0 -ext -aswap -DMX -decim 1240 -dspfvs 20 -grpdly 68 -ws 8 -
633
      noi2f \
634
                           1024 -vN
                                                       40
                                                            -zN
                                                                                 40
        -xN
635
      \backslash
636
                                                       20
                                                                                 20
         -xT
                             512 -yT
                                                           -zT
637
      \backslash
638
        -xMODE
                             DQD -yMODE Echo-AntiEcho -zMODE Echo-AntiEcho
639
      \setminus
640
        -xSW
                      16129.032 -ySW
                                                8000.000 -zSW
                                                                          8000.000
641
      \setminus
642
                        800.134 -yOBS
                                                800.134 -zOBS
                                                                          800.134
        -xOBS
643
      \setminus
644
                          4.821 -yCAR
                                                    4.821 -zCAR
                                                                              4.821
        -xCAR
645
      \setminus
646
        -xLAB
                             Hz -yLAB
                                                       Hy -zLAB
                                                                                 Нx
647
      \setminus
648
                               3 -aq2D
                                                Complex
        -ndim
649
      \backslash
650
      | pipe2xyz -x -out ./fid/test%03d.fid -ov
651
652
653
      xyz2pipe -in fid/test%03d.fid -x
654
      | nmrPipe -fn SP -off 0.5 -end 0.98 -pow 2 -c 0.5
655
      | nmrPipe -fn ZF -zf 2 -auto
                                                                  \
656
      | nmrPipe -fn FT
                                                                  \backslash
      | nmrPipe -fn PS -p0 149 -p1 38 -di
657
                                                                  \backslash
658
     | nmrPipe -fn TP
                                                                  \backslash
659
                                                                  \setminus
     | nmrPipe -fn SP -off 0.5 -end 0.98 -pow 1 -c 0.5
660
     | nmrPipe -fn ZF -zf 2 -auto
                                                                  \
661
                                                                  \setminus
      | nmrPipe -fn FT
662
      | nmrPipe -fn PS -p0 135 -p1 0 -di
                                                                  \setminus
663
      | nmrPipe -fn ZTP
                                                                  \
664
      | nmrPipe -fn SP -off 0.5 -end 0.98 -pow 1 -c 0.5
                                                                  \
665
      | nmrPipe -fn ZF -zf 2 -auto
                                                                  \
666
                                                                  \
      | nmrPipe -fn FT
667
      | nmrPipe -fn PS -p0 90 -p1 0 -di
                                                                  \backslash
668
      | nmrPipe -fn TP
                                                                  669
      | nmrPipe -fn POLY -ord 4 -nw 300 -nl 301 1740
                                                                  \backslash
670
     | pipe2xyz -out ft/test%04d.ft3 -x -ov -verb
671
```

628 A5: NMRPipe conversion and processing scripts

672 Author contributions

TK: conceptualization (equal), 3D printing and cleaning, figures 1-6, review and editing
(equal). MDG: review and editing (equal), data collection (equal), figure 8. JY: review and
editing (equal), data collection (equal), figure 7. AB: conceptualization (equal), writing
original draft, review and editing (equal), recording data (equal).

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682 Competing interests

AB is a member of the editorial board of *Magnetic Resonance*. The peer-review process
was guided by an independent editor, and the authors also have no other competing
interests to declare.

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- 689
- 690

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