

SUPPORTING INFORMATION

Direct Evaluation of Protein-Lipid Contacts Reveals Protein Membrane Immersion and Isotropic Bicelle Structure

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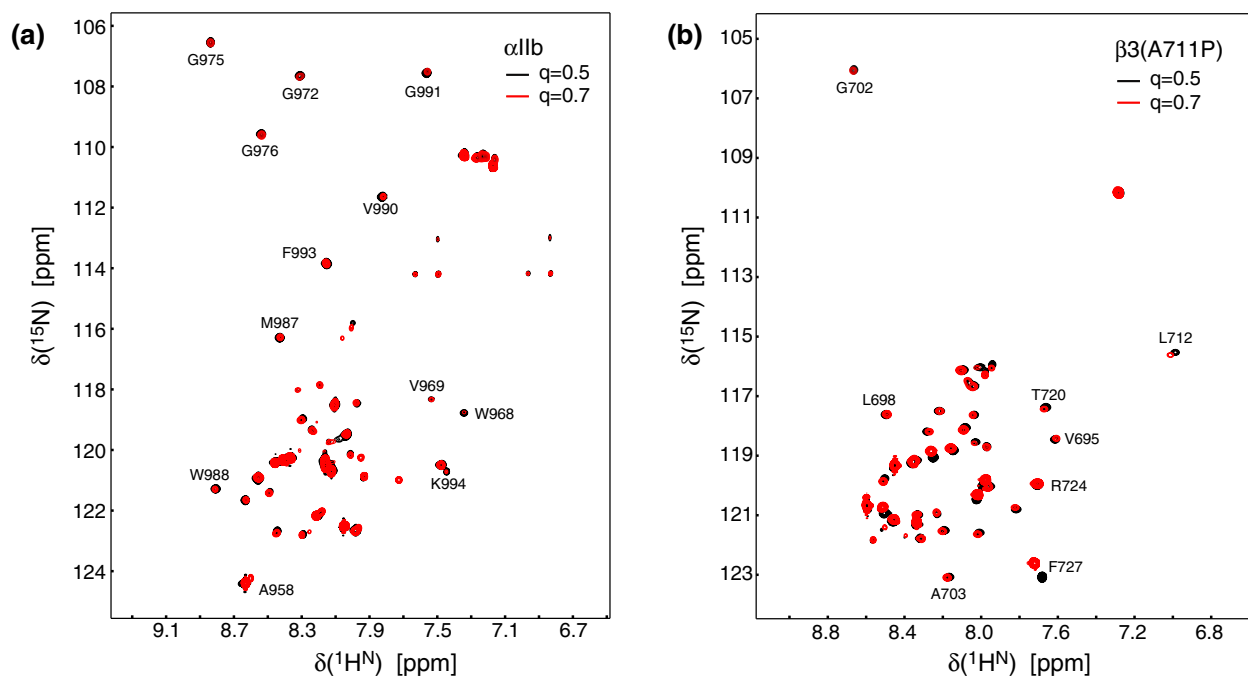


Figure S1. Illustration of spectral quality of bicelle-immersed integrin α IIb and β 3(A711P) TM peptides. TROSY-type H-N correlation spectra of 1 mM of the indicated integrin TM peptides in 350 mM DHPC/175 mM POPC ($q=0.5$ bicelles) and 400 mM DHPC/280 mM POPC ($q=0.7$ bicelles), respectively. The solutions additionally contained 5 mM tris(2-carboxyethyl)phosphine, 0.02% NaN_3 , 6% D_2O , and 25 mM $\text{NaH}_2\text{PO}_4/\text{Na}_2\text{HPO}_4$, pH 7.4. Spectra were recorded at 700 MHz and 35 °C. Selected backbone assignments are shown.

Table S1. Effect of presaturation on intensity of vicinal signals.

Presaturation ^{a,b}	Attenuation [%] DHPC(#4 CH ₂)	Attenuation [%] POPC(#3 CH ₂)	Attenuation [%] DHPC(#2 CH ₃)	Attenuation [%] POPC(#1 CH ₃)
POPC(#1 CH ₃)	0	2	18	80
DHPC(#2 CH ₃)	3	2	85	19
POPC(#3 CH ₂)	18	87	1	4
DHPC(#4 CH ₂)	74	25	4	1

^aPeak labels refer to the numbering of DHPC and POPC resonances depicted in Figure 1a.

^bPresaturation of 500 ms, B₀= 700 MHz and B₁= 11.2 Hz.