



### *MS analysis*

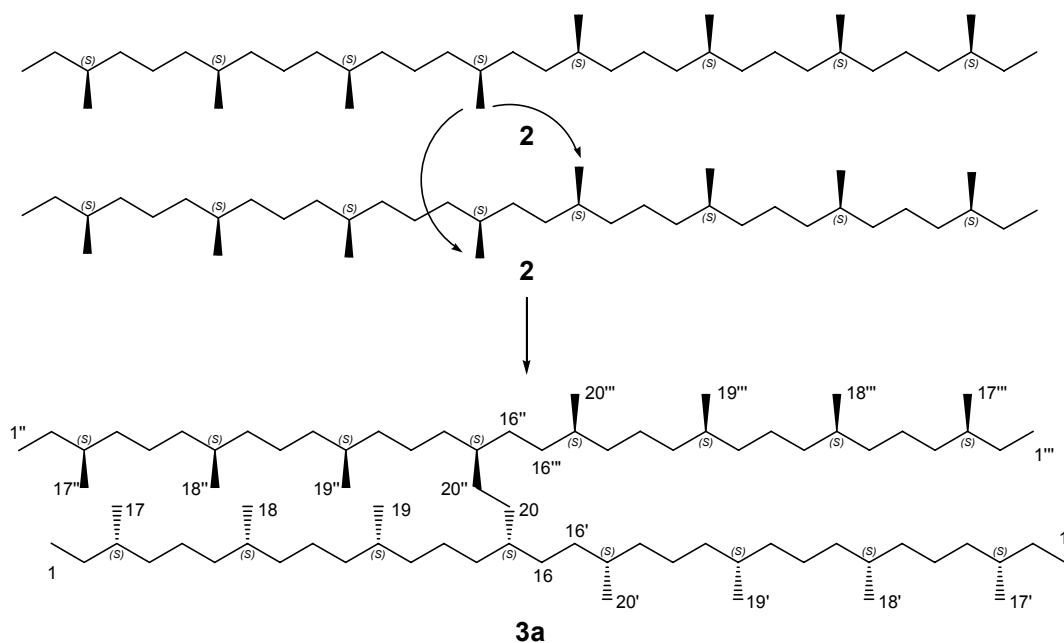
The positive mode electrospray ionisation mass spectrum (Fig. 13 below) of the permethylated 20-bis-16,16'-biphytanyl tetraacids was recorded on a Micromass QTOF 2W mass spectrometer. Positive mode APCI MS/MS spectra were also recorded, but structural fragmentation was not obtained with collision energies of 15, 20, 25 or 30 V. Accurate mass measurement (positive mode APCI) of the main component with 6 rings, gave  $m/z$  1288.1370 (calc. 1288.1403 for  $C_{84}H_{150}O_8$ ).

### *Derivatisation of the tetraacid 1 to the permethyl ester*

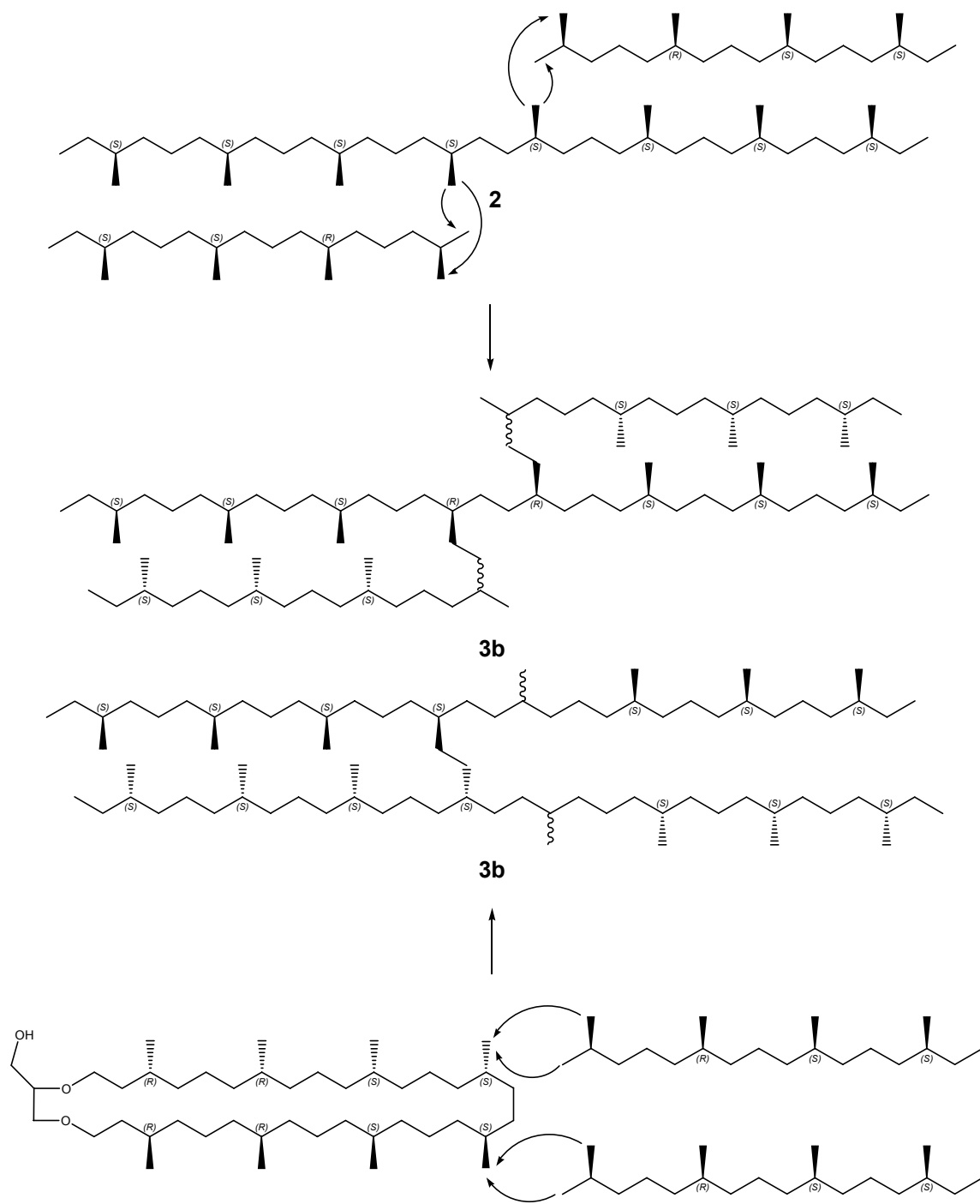
An aliquote of the tetraacid **1** (ca. 20 mg) which had been purified by extraction into toluene as described above was dissolved in THF (5 ml). MeOH (5 ml) and  $BF_3 \cdot OMe_2$  (1 ml) were added, and the solution refluxed for 5 min. Diethyl ether (15 ml) was added, and the solution washed with water (3x 10 ml) and saturated NaCl solution (2x 5 ml). The organic phase was dried over  $MgSO_4$  and taken to dryness under reduced pressure. NMR (400 MHz) showed that 95% of the acid moieties were methylated. This sample was submitted to electrospray MS analysis.

### *MM2 Calculations*

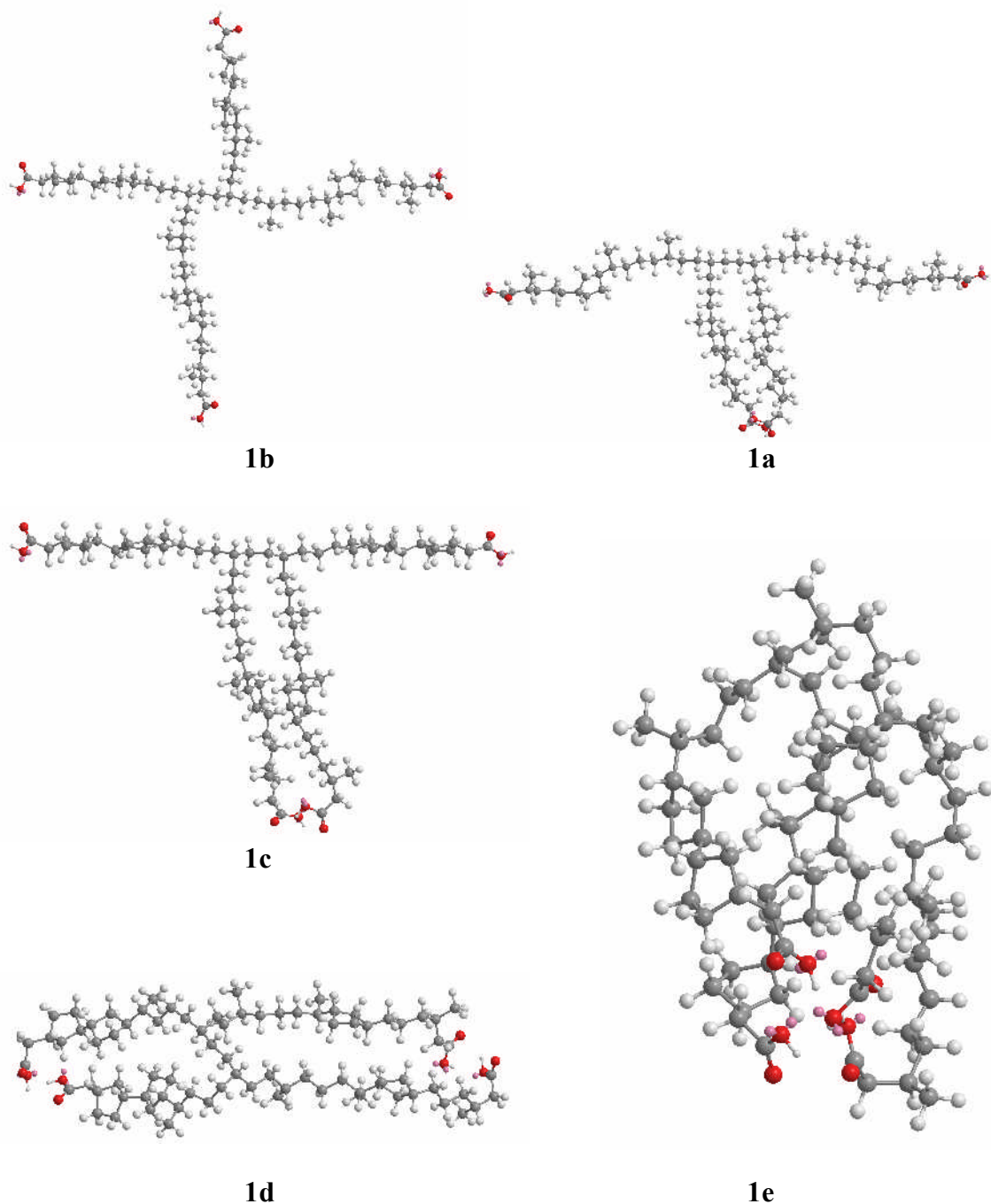
The MM2 program used here is incorporated in the ChemOffice2005 (Chem3D Ultra ver. 9.0.1) program package from CambridgeSoft, Cambridge, MA, USA. Structures for the tetraacids (**1**) calculated by MM2 methods are given in Fig. 5 and Fig 3 here in Chem3D format.



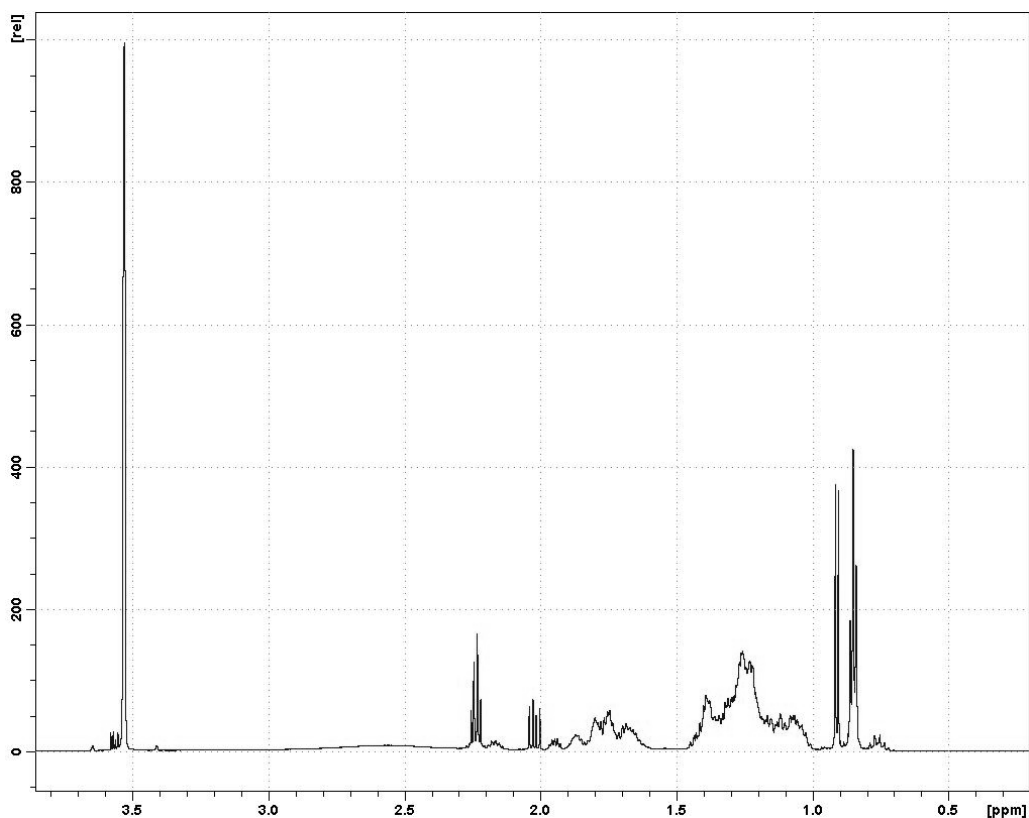
**Fig. 1** Head-to-head coupling of two 16,16'-biphytane (**2**) moieties provides one stereoisomer of 20-bis-16,16'-biphytane (**3a**) regardless of whether the coupling, indicated by arrows, occurs between Me-20 and Me-20'' or Me-20' and Me-20''.



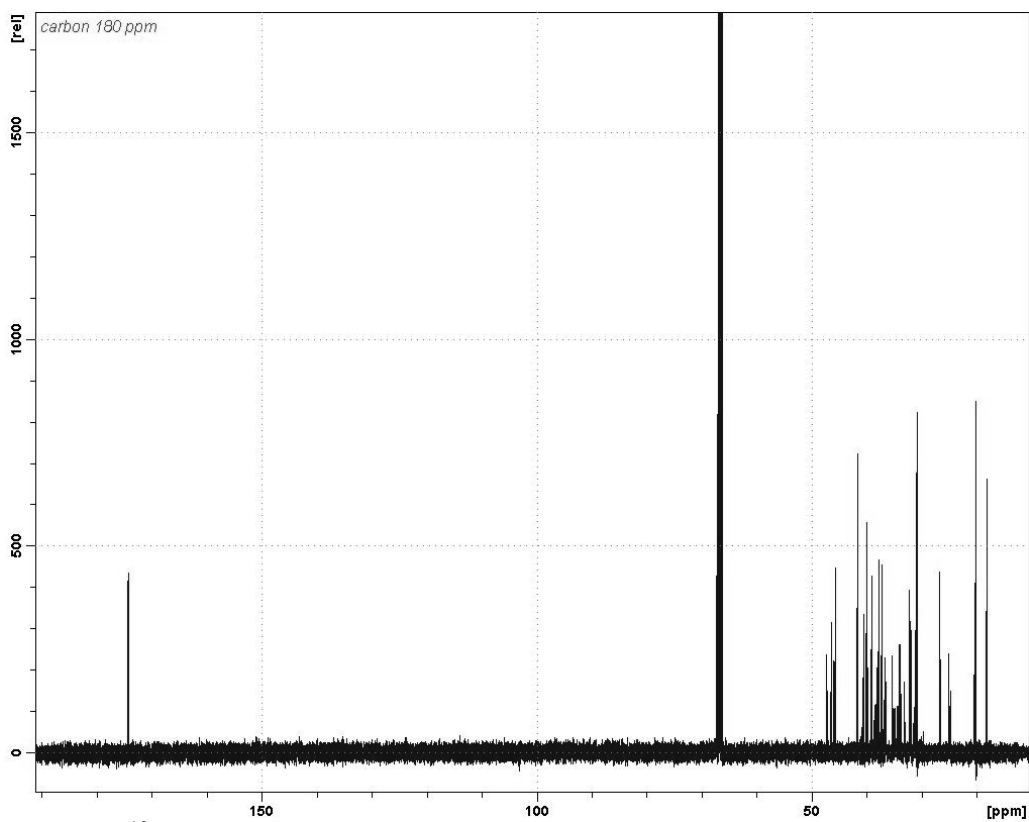
**Fig. 2**



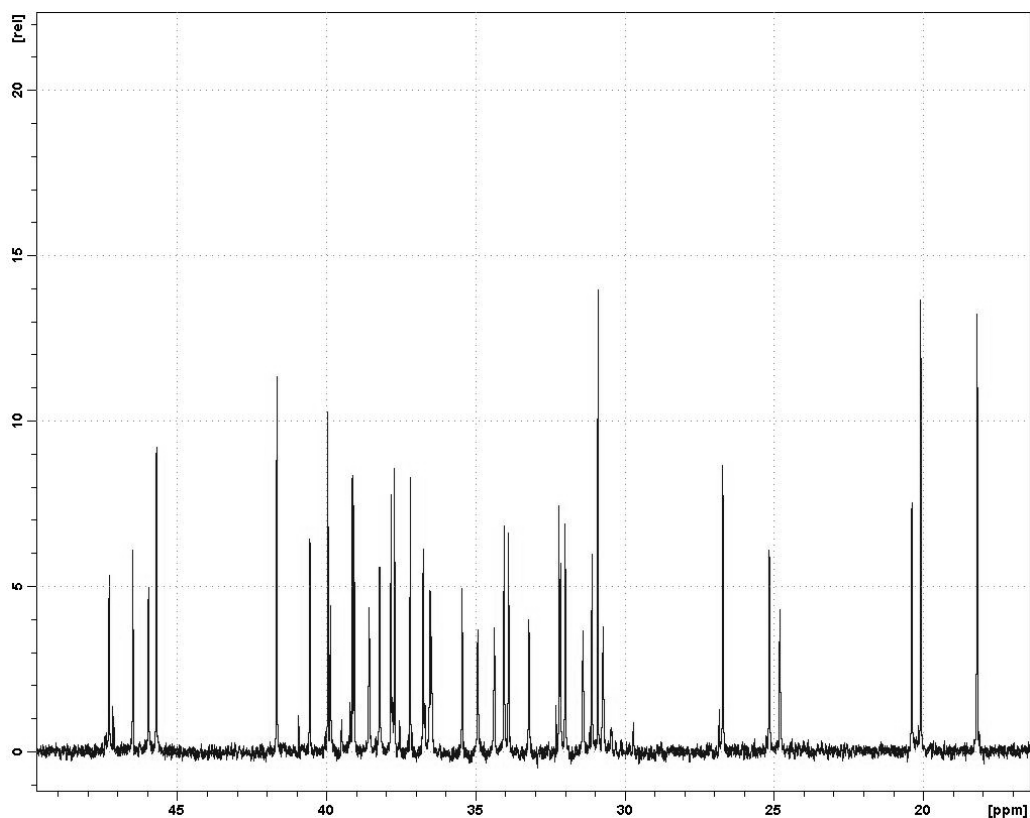
**Fig. 3** Conformations calculated for (6:17,10:18,10':18',6'':17'',10'':18'',10''':18''')-hexacyclo-20-bis-16,16'-biphytane-1,1',1'',1'''-tetracarboxylic acid (**1**), with stereochemistry preserved from 20-bis-16,16'-biphytane (**3**). Free energies of the conformations are given in Table 2 below. Structures can be viewed in 3D by Cambridgesoft Chem3D software.



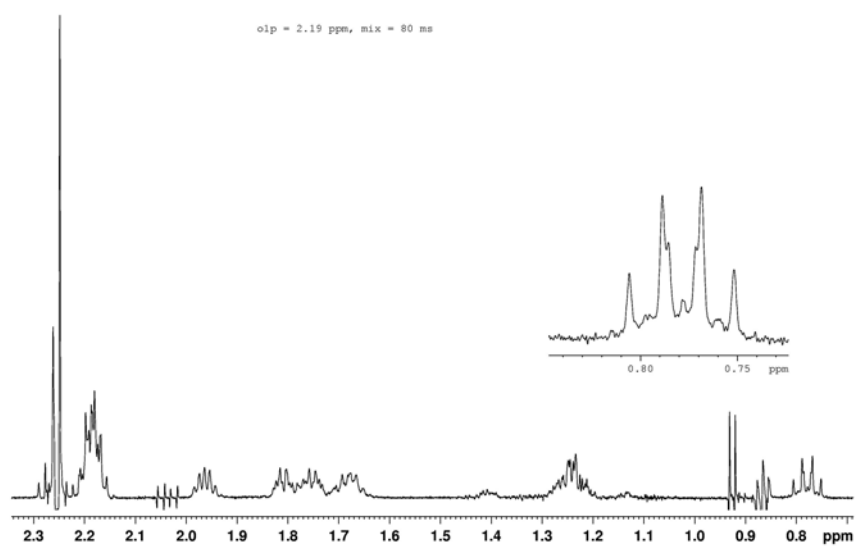
**Fig. 4** 600 MHz <sup>1</sup>H spectrum in dioxane-*d*<sub>8</sub> of the 20-bis-16,16'-biphtanyl tetraacids.



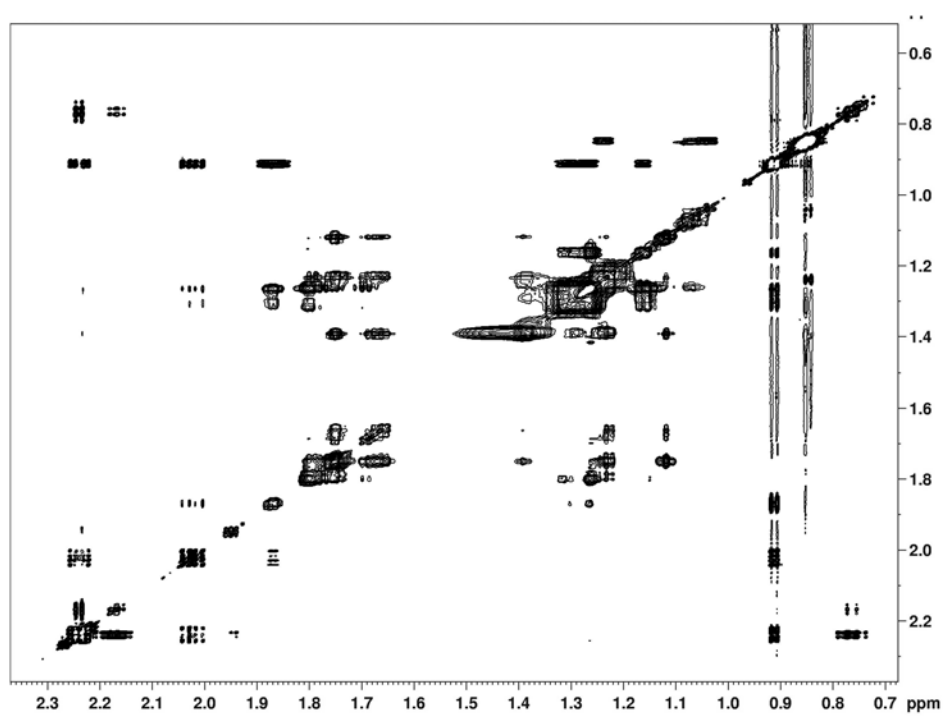
**Fig. 5** 100 MHz <sup>13</sup>C NMR spectrum of the 20-bis-16,16'-biphtanyl tetraacids.



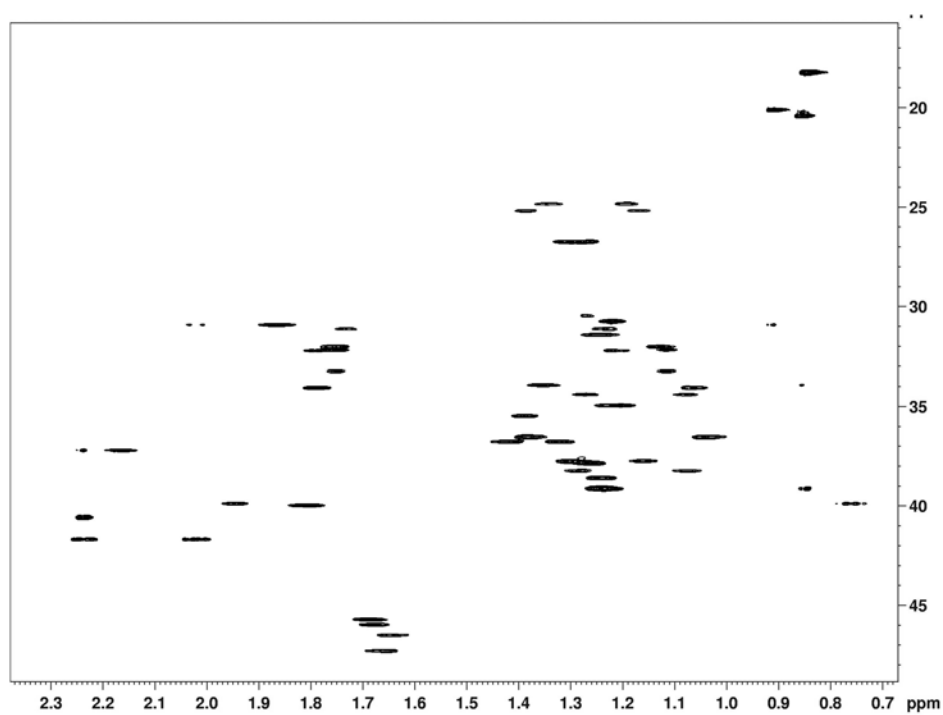
**Fig. 6** Aliphatic region of 150 MHz  $^{13}\text{C}$  spectrum (dioxane- $d_8$ ) of the 20-bis-16,16'-biphytanyl tetraacids.



**Fig. 7** 1D TOCSY spectrum of the 20-bis-16,16'-biphytanyl tetraacids, with irradiation of the  $\alpha$ -carbonyl protons at 2.16 ppm. Enlargement of the multiplet at 0.77 ppm inserted.

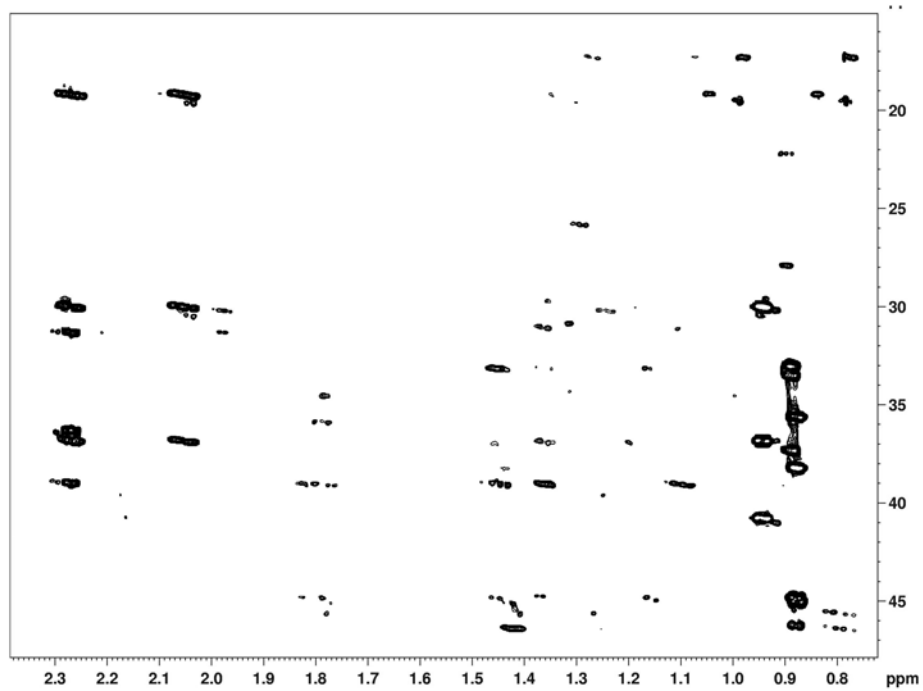


**Fig. 8** 2D ROESY spectrum (300 ms mix) of the 20-bis-16,16'-biphytanyl tetraacids.

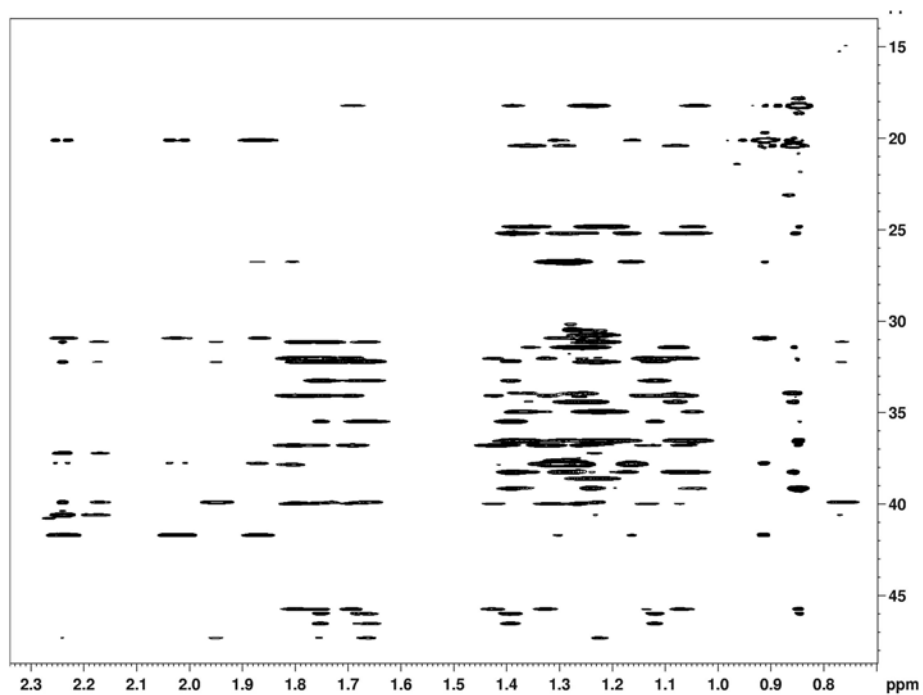


**Fig. 9**  $^1\text{H}$ - $^{13}\text{C}$  HSQC spectrum of the 20-bis-16,16'-biphytanyl tetraacids.

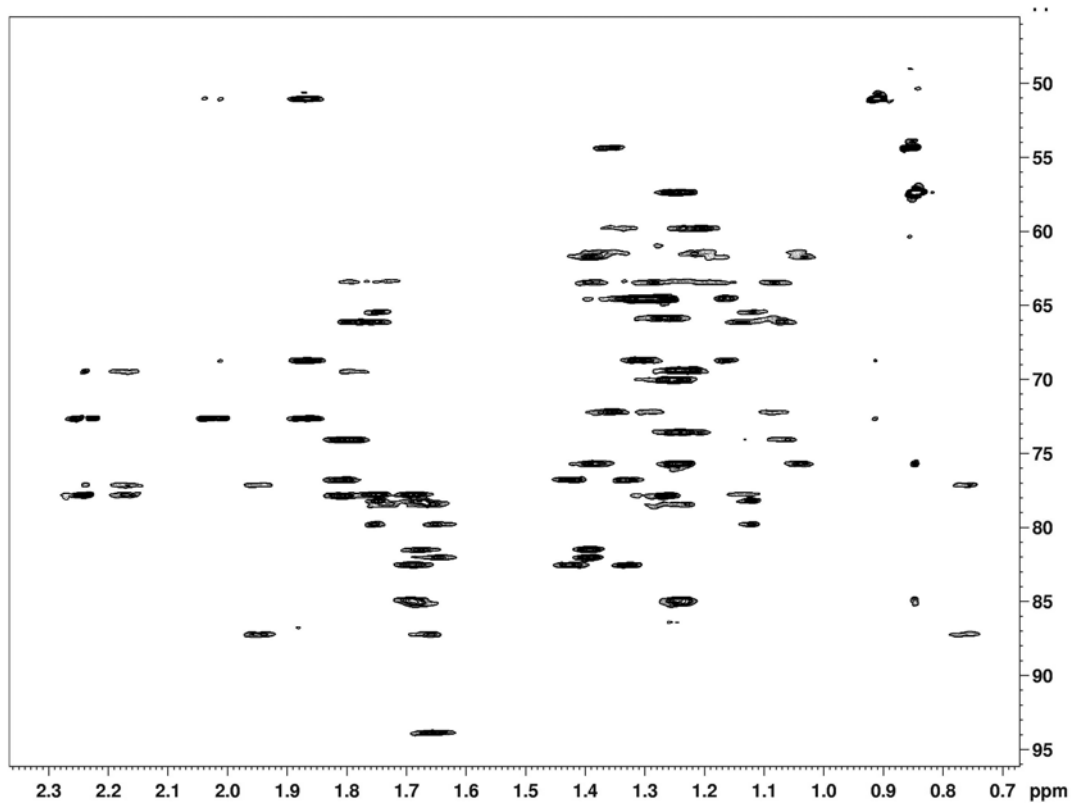




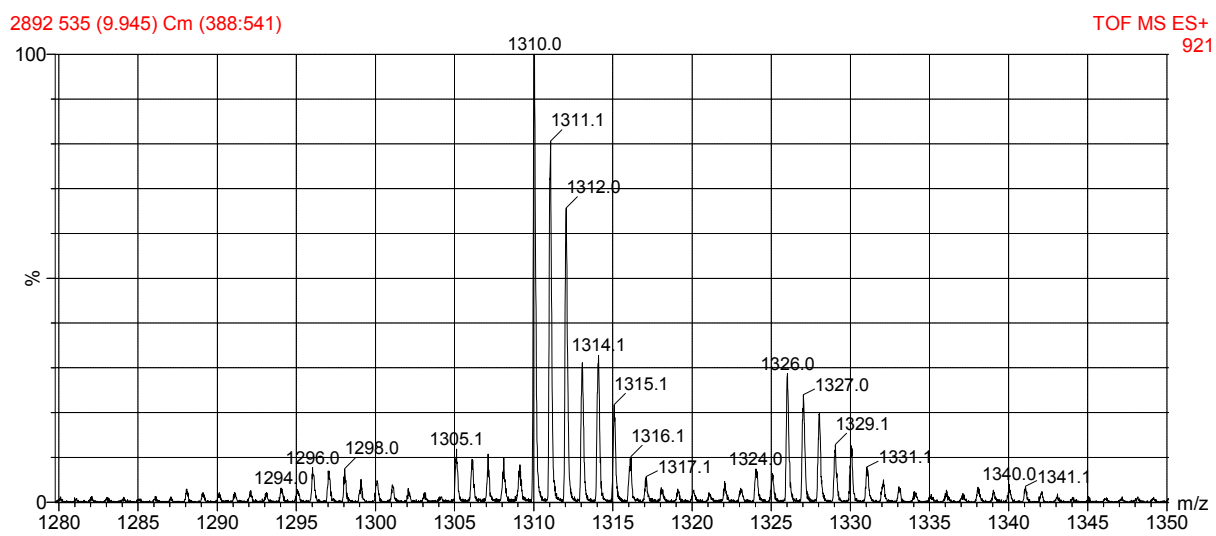
**Fig. 10** Long-range part of the  $^1\text{H}$ - $^{13}\text{C}$  HMSC spectrum of the 20-bis-16,16'-biphytanyl tetraacids.



**Fig. 11** 2D HSQC-TOCSY spectrum of the 20-bis-16,16'-biphytanyl tetraacids.



**Fig. 12** 1,1-ADEQUATE spectrum of the 20-bis-16,16'-biphytanyl tetraacids.



**Fig. 13** ESI (positive) mass spectrum of the permethylated 20-bis-16,16'-biphytanyl tetraacids.

**Table 1**  $^{13}\text{C}$ - and  $^1\text{H}$ -NMR data of **1**.

Carbon #	$\delta_{\text{C}}$ (ppm)				$\delta_{\text{H}}$ (ppm)
	$\text{CH}_3$	$\text{CH}_2$	$\text{CH}$	$\text{C}$	
1, 1''				174.37	
1', 1'''				174.36	
2, 2''		40.60			2.240
2', 2'''		41.70			2.236, 2.020
3, 3''			37.23		2.160
3', 3'''			30.94		1.870
4, 4''		32.25			1.794, 1.221
4', 4'''		37.76			1.300, 1.162
5, 5''		31.14			1.724, 1.233
5', 5'''		26.74			1.310, 1.270
6, 6''			47.31		1.657
6', 6'''		37.86			1.419, 1.324
7, 7''			46.53		1.648
7', 7'''			39.99		1.807
8, 8''		33.23			1.752, 1.126
8', 8'''		34.07			1.794, 1.064
9, 9''		32.18			1.751, 1.123
9', 9'''		32.02			1.751, 1.130
10, 10''			46.00		1.678
10', 10'''			45.74		1.691
11, 11''			39.10		1.236
11', 11'''			39.15		1.236
12, 12''		36.51			1.381, 1.038
12', 12'''		36.57			1.381, 1.038
13, 13''		24.86			1.342, 1.192
13', 13'''		25.20			1.380, 1.173
14, 14''		34.98			1.221
14', 14'''		38.26			1.290, 1.082
15, 15''			38.62		1.243
15', 15'''			33.94		1.236
16, 16''		31.46			1.249
16', 16'''		34.44			1.274, 1.083
17, 17''		39.89			1.942, 0.765
17', 17'''	20.12				0.910
18, 18''		35.48			1.396
18', 18'''		36.78			1.419, 1.324
19, 19''	18.22				0.844
19', 19'''	18.22				0.844
20, 20''		30.81			1.225
20', 20'''	20.41				0.854

**Table 2** MM2 calculated free energies (kcal/mol) for conformations **1a-e** of tetraacid **1**. For structures, see Fig. 3 above.

Conformation	<b>1b</b>	<b>1a</b>	<b>1c</b>	<b>1d</b>	<b>1e</b>
Stretch	8.24	8.34	8.26	8.33	8.38
Bend	31.97	32.91	32.71	34.76	37.50
Stretch-Bend	1.44	1.54	1.51	1.61	1.70
Torsion	37.31	38.18	40.07	43.11	50.62
Non-1,4 VDW*	-21.70	-33.55	-40.74	-50.99	-89.11
1,4 VDW*	51.05	51.29	51.38	52.38	54.62
Dipole/Dipole	6.29	2.99	4.66	3.32	7.30
Total	114.60	101.70	97.85	92.52	70.71

\*) van der Waals interaction

## References

1. H. Mediaas, K. V. Grande, B. M. Hustad, A. Rasch, H. G. Rueslåtten, J. E. Vindstad, *Proceedings - SPE 5<sup>th</sup> International Oilfield Scale Symposium*, Aberdeen, SPE 80404, **2003**.