

SUPPORTING INFORMATION

**Polarisation Effects on the Solvation Properties of Alcohols**

Stefan Henkel, Maria Cristina Misuraca, Pavle Troselj, Jonathan Davidson, Christopher A. Hunter\*

Department of Chemistry, University of Cambridge, Lensfield Road, Cambridge CB2 1EW, UK

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BINDING ISOTHERMS for the formation of 1:1 complexes of phenols **1-5** with phosphine oxide **6** in *n*-octane at 298 K.

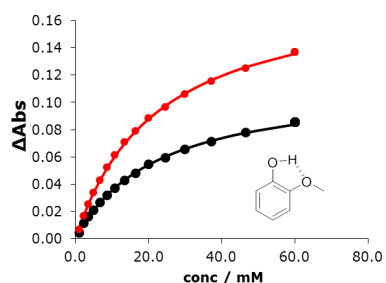


Figure S1. Fit of the experimental shifts of the UV-vis signals of 2-methoxyphenol upon titration with tri-*n*-butyl phosphine oxide.

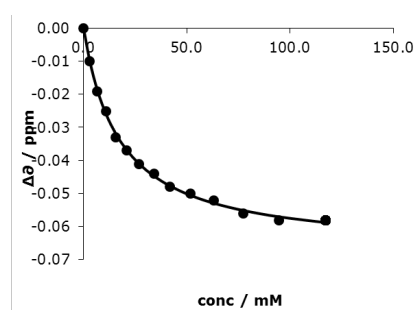


Figure S2. Fit of the experimental shifts of the  $^1\text{H}$  NMR signals of the methyl protons of 2-methoxyphenol upon titration with tri-*n*-butyl phosphine oxide.

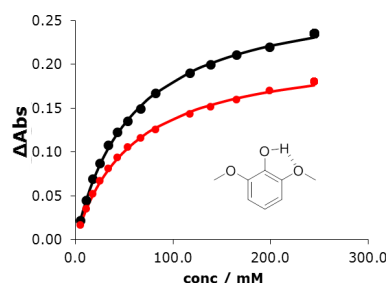


Figure S3. Fit of the experimental shifts of the UV-vis signals of 2,6-dimethoxyphenol upon titration with tri-*n*-butyl phosphine oxide.

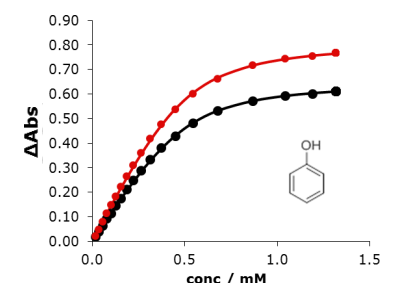


Figure S4. Fit of the experimental shifts of the UV-vis signals of phenol upon titration with tri-*n*-butyl phosphine oxide.

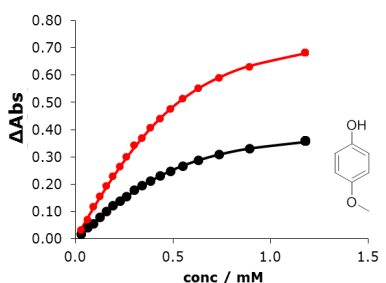


Figure S5. Fit of the experimental shifts of the UV-vis signals of 4-methoxyphenol upon titration with tri-*n*-butyl phosphine oxide.

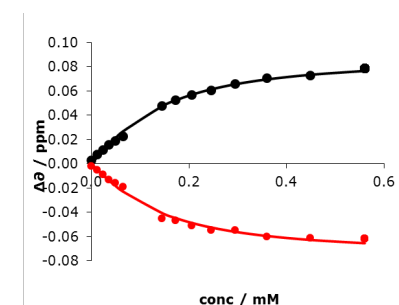


Figure S6. Fit of the experimental shifts of the  $^1\text{H}$  NMR signals of the aromatic protons of 4-methoxyphenol upon titration with tri-*n*-butyl phosphine oxide.

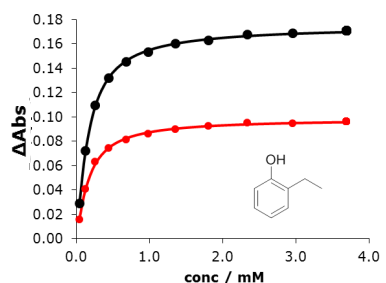
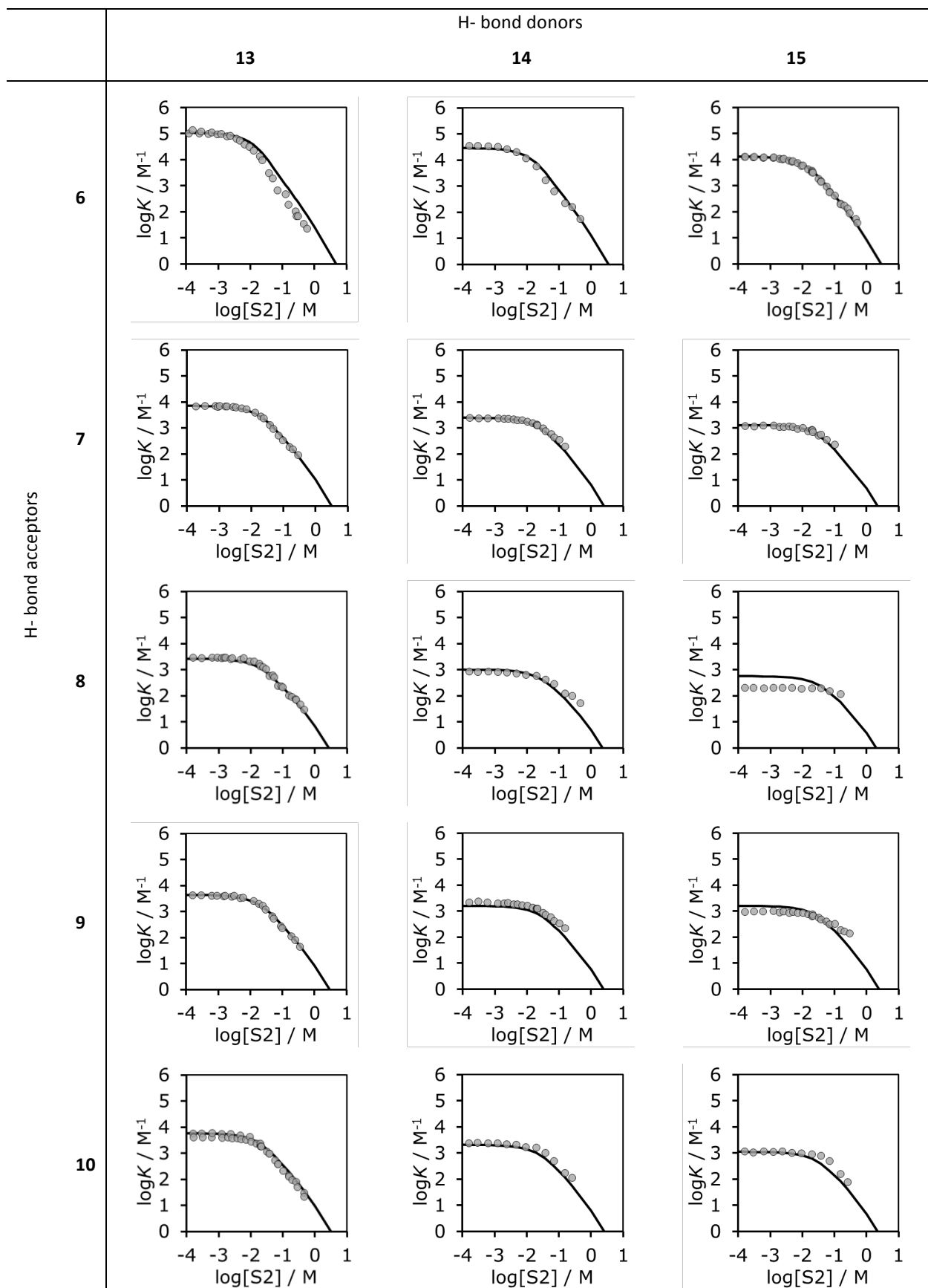
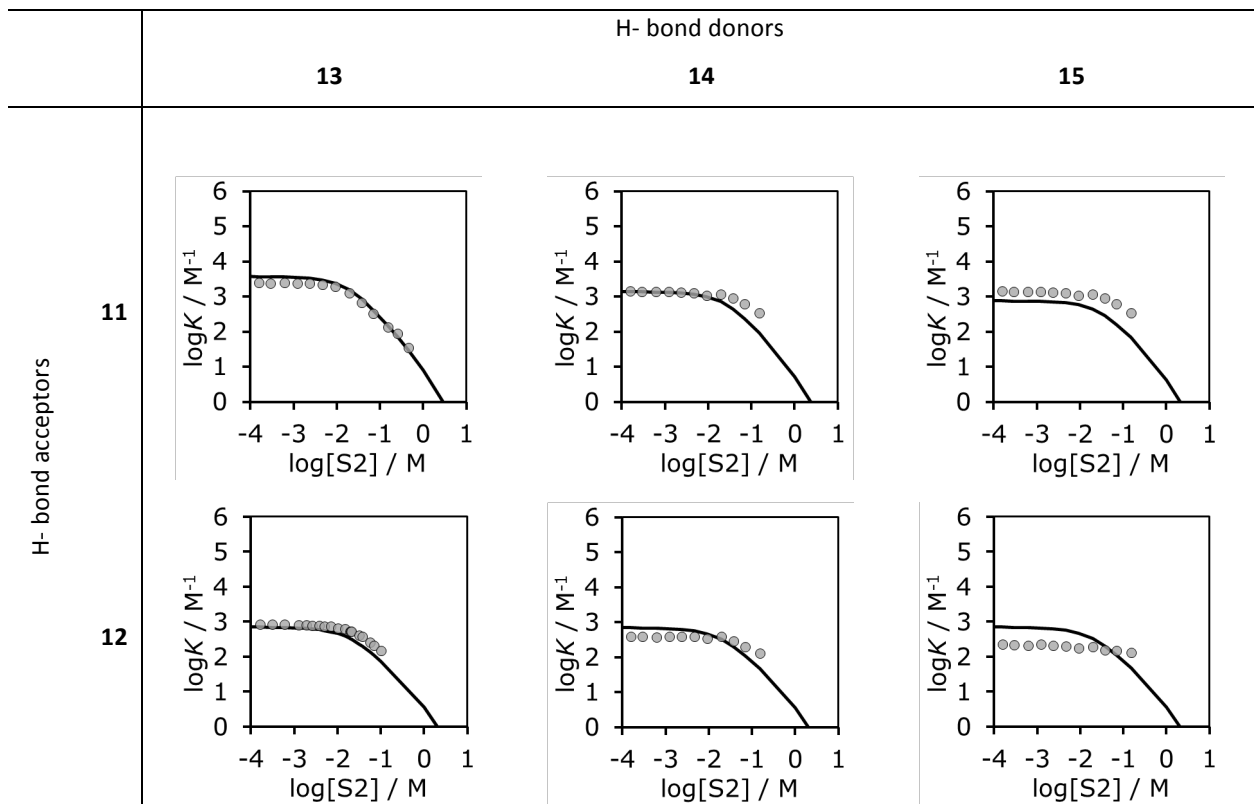


Figure S7. Fit of the experimental shifts of the UV-vis signals of 2-ethylphenol upon titration with tri-*n*-butyl phosphine oxide.

EXPERIMENTAL AND PREDICTED ASSOCIATION CONSTANTS of donors **12-14** and acceptors **5-11** the as a function of concentration of 1-octanol (S2) in *n*-octane (S1) at 298 K.





SPECIATION OF ALCOHOL AGGREGATES for 1-octanol **A1** and 3-ethyl-3-pentanol **A9** in tetrachloromethane based on literature data.

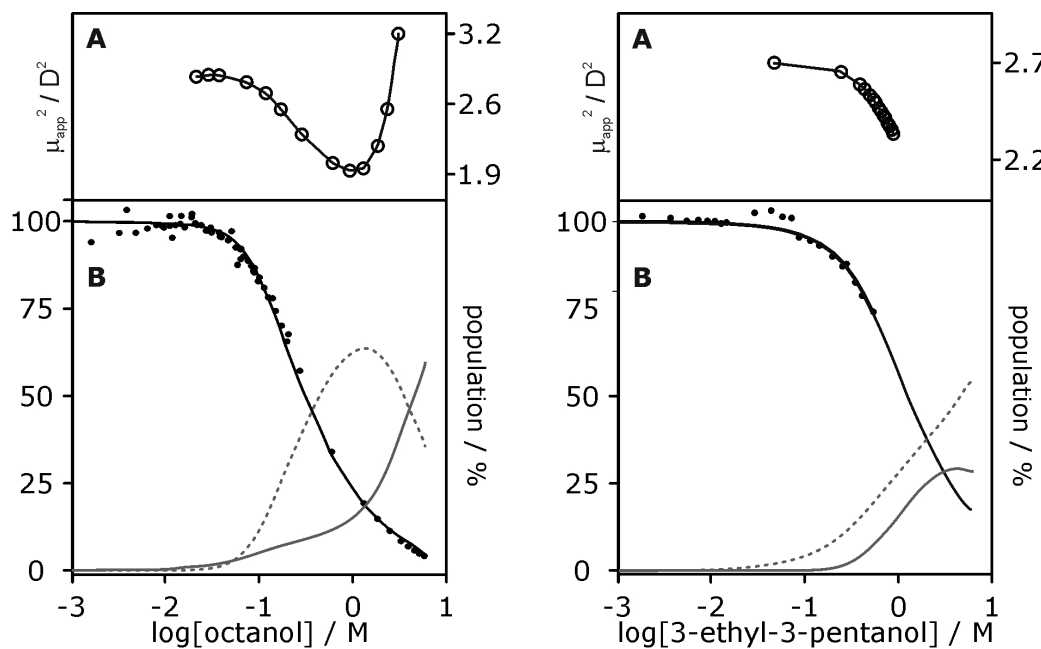


Figure S8. Left: **A:** Apparent dipole moment of 1-octanol in tetrachloromethane.<sup>[1]</sup> **B:** Population of monomeric 1-octanol in tetrachloromethane measured by IR spectroscopy.<sup>[2-4]</sup> Right: **A:** Apparent dipole moment of 3-ethyl-3-pentanol in tetrachloromethane.<sup>[5]</sup> **B:** Population of monomeric 3-ethyl-3-pentanol in tetrachloromethane measured by IR spectroscopy.<sup>[4]</sup> Black lines correspond to fits of the experimental data with  $K_n = 3 \text{ M}^{-1}$ ,  $\alpha_c = 13$  and  $K_4 = 75 \text{ M}^{-3}$  for 1-octanol and  $K_n = 0.7 \text{ M}^{-1}$ ,  $\alpha_c = 3.4$  and  $K_4 = 0.4 \text{ M}^{-3}$  for 3-ethyl-3-pentanol. The grey lines show the populations of linear aggregates (solid line) and cyclic tetramer (dashed line).

#### References

- [1] Campbell, C.; Brink, G.; Glasser, L., *J. Phys. Chem.* **1975**, *79*, 660.
- [2] Fletcher, A. N., *J. Phys. Chem.* **1969**, *73*, 2217.
- [3] Brink, G.; Drought, Z.; Glasser, L., *S. Afr. J. Chem.* **1986**, *39*, 163-168.
- [4] Nodland, E., *Appl. Spectrosc.* **2000**, *54*, 1339.
- [5] Kunst, M.; Vanduijn, D.; Bordewijk, P., *Ber. Bunsen Phys. Chem.* **1976**, *80*, 839.

H-BOND PARAMETERS FOR ALCOHOL MONOMERS AND DIMERS calculated from the molecular electrostatic potential.

Table S1. H-bond parameters for alcohols **A1-A10** calculated for the surface site interaction points at the B3LYP/6-31G(d) level of theory.

		Monomer		Dimer	
		$\alpha$	$\beta$	$\alpha$	$\beta$
1-Octanol	<b>A1</b>	2.8	5.3	3.5	7.4
1-Decanol	<b>A2</b>	2.8	5.3	3.5	7.4
2-Heptanol	<b>A3</b>	2.7	5.3	3.3	6.9
5-Nonanol	<b>A4</b>	2.7	5.1	3.3	5.8
2,4-Dimethyl-4-heptanol	<b>A5</b>	2.7	4.9	3.2	5.3
Menthol	<b>A6</b>	2.7	5.1	3.5	5.9
4-Penty-cyclohexanol	<b>A7</b>	2.7	5.3	3.3	6.5
2-Methyl-2-hexanol	<b>A8</b>	2.6	5.3	3.2	6.4
3-Ethyl-3-pentanol	<b>A9</b>	2.7	5.1	3.3	6.7
1,1,1-Trifluoro-2-octanol	<b>A10</b>	3.2	3.5	4.0	5.7

OPTIMIZED GEOMETRIES of monomer and dimers of alcohols **A1-A10** and methanol. Geometry optimizations, frequency calculations and calculations of the electrostatic potential were carried out using the B3LYP function and the 6-31G\* basis set as implemented in Gaussian 03.

(Gaussian 03, Revision E.01; Frisch, M. J. T., G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Montgomery, Jr., J. A.; Vreven, T.; Kudin, K. N.; Burant, J. C.; Millam, J. M.; Iyengar, S. S.; Tomasi, J.; Barone, V.; Mennucci, B.; Cossi, M.; Scalmani, G.; Rega, N.; Petersson, G. A.; Nakatsuji, H.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Klene, M.; Li, X.; Knox, J. E.; Hratchian, H. P.; Cross, J. B.; Bakken, V.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Ayala, P. Y.; Morokuma, K.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Zakrzewski, V. G.; Dapprich, S.; Daniels, A. D.; Strain, M. C.; Farkas, O.; Malick, D. K.; Rabuck, A. D.; Raghavachari, K.; Foresman, J. B.; Ortiz, J. V.; Cui, Q.; Baboul, A. G.; Clifford, S.; Cioslowski, J.; Stefanov, B. B.; Liu, G.; Liashenko, A.; Piskorz, P.; Komaromi, I.; Martin, R. L.; Fox, D. J.; Keith, T.; Al-Laham, M. A.; Peng, C. Y.; Nanayakkara, A.; Challacombe, M.; Gill, P. M. W.; Johnson, B.; Chen, W.; Wong, M. W.; Gonzalez, C.; Pople, J. A. Gaussian, Inc.: Pittsburgh PA, 2003.)

Methanol							
<b>Monomer</b>				<b>Dimer</b>			
C	-0.511	0.316	0.000	H	0.372	2.995	-0.092
H	-0.154	-0.693	0.000	O	1.123	2.648	-0.612
H	-0.154	0.820	-0.874	H	-1.817	3.861	0.944
H	-1.581	0.316	0.000	O	-1.316	3.061	0.724
O	-0.034	0.990	1.168	C	2.000	1.971	0.292
H	-0.352	1.896	1.167	H	2.835	1.578	-0.248
<b>Trimer</b>				H	2.347	2.659	1.035
H	0.372	2.995	-0.092	H	1.473	1.169	0.767
O	1.123	2.648	-0.612	C	-2.123	2.242	-0.127
C	2.000	1.971	0.292	H	-2.360	2.782	-1.020
H	2.835	1.578	-0.248	H	-1.585	1.352	-0.380
H	2.347	2.659	1.035	H	-3.026	1.980	0.382
<b>Cyclic tetramer</b>				C	0.121	2.783	1.138
H	-1.826	3.874	0.889	H	-0.764	3.418	1.183
O	-1.340	3.038	0.744	H	0.995	3.436	1.059
H	-2.745	6.240	1.302	H	0.190	2.235	2.083
O	-2.393	5.642	0.625	O	-0.002	1.928	0.024
C	-2.146	2.209	-0.097	H	0.777	1.322	-0.008
H	-1.640	1.284	-0.276	C	-2.791	0.123	-1.135
H	-3.083	2.017	0.383	H	-3.425	-0.762	-1.181
H	-2.317	2.707	-1.029	H	-2.250	0.199	-2.084
C	-1.256	6.264	0.020	H	-3.445	0.996	-1.046
H	-0.512	6.448	0.767	O	-1.928	-0.005	-0.028
H	-0.854	5.617	-0.732	H	-1.322	0.773	0.005
H	-1.551	7.190	-0.426	C	2.788	-0.122	-1.136
				H	3.423	0.763	-1.182
				H	2.242	-0.192	-2.082
				H	3.442	-0.995	-1.056
				O	1.932	0.002	-0.024
				H	1.326	-0.776	0.008
				C	-0.125	-2.780	1.140
				H	0.760	-3.413	1.197
				H	-0.997	-3.435	1.057
				H	-0.203	-2.227	2.081
				O	0.006	-1.931	0.022
				H	-0.772	-1.325	-0.018

1-Octanol A1			
Monomer			Dimer
C	-3.764	0.450	0.000
H	-3.796	1.104	0.913
H	-3.796	1.104	-0.913
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-2.521	-1.074	0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
H	-0.011	-1.053	0.902
C	1.226	0.477	0.000
H	1.221	1.144	-0.903
C	2.482	-0.367	0.000
H	1.221	1.144	0.902
H	2.488	-1.034	-0.903
H	2.488	-1.034	0.902
C	3.727	0.496	0.000
H	3.719	1.162	-0.903
C	4.980	-0.341	0.000
H	3.718	1.163	0.902
H	5.018	-0.995	-0.904
H	5.017	-0.995	0.905
H	5.884	0.314	0.000
O	-4.875	-0.433	0.000
H	-5.665	0.119	-0.001
			H 0.372741 2.996967 -0.091911
			O 1.122719 2.648454 -0.611509
			C 1.992181 1.976722 0.284767
			C 3.123377 1.342321 -0.516714
			H 1.457628 1.194302 0.851784
			H 2.416195 2.673288 1.030060
			H -1.817467 3.860282 0.944951
			O -1.315777 3.060691 0.723950
			C -2.125856 2.238958 -0.130491
			H -1.442903 1.471132 -0.504812
			H -2.473665 2.816095 -1.000691
			C -3.306019 1.614610 0.609439
			H 2.682285 0.667406 -1.262828
			C 4.127960 0.580246 0.355024
			H 3.635526 2.134396 -1.079552
			H 3.601082 -0.202201 0.921866
			H 4.554049 1.262389 1.105982
			C 5.268597 -0.061339 -0.445994
			H 4.844284 -0.740527 -1.200216
			C 6.269353 -0.831805 0.424527
			H 5.801224 0.719884 -1.008230
			H 5.734908 -1.612726 0.986166
			H 6.691811 -0.152321 1.179898
			C 7.412074 -1.474556 -0.372026
			H 6.991363 -2.153443 -1.128948
			C 8.409202 -2.247405 0.500700
			H 7.949745 -0.694430 -0.931639
			H 7.872130 -3.027995 1.058403
			H 8.828746 -1.569122 1.257517
			C 9.549216 -2.885045 -0.300107
			H 9.164026 -3.594807 -1.042700
			H 10.242132 -3.429261 0.351998
			H 10.127732 -2.125155 -0.840007
			C -4.133949 0.675664 -0.278322
			H -3.952939 2.414834 1.001004
			H -2.921225 1.069993 1.481479
			H -4.499927 1.228086 -1.156382
			C -5.324653 0.041525 0.452647
			H -3.482852 -0.119601 -0.669488
			H -5.978418 0.836813 0.840373
			H -4.958626 -0.506694 1.333072
			C -6.147343 -0.906284 -0.429605
			H -6.511162 -0.357924 -1.311284
			C -7.338862 -1.541238 0.298691
			H -5.492407 -1.701153 -0.816223
			H -7.995835 -0.746855 0.683560
			H -6.975961 -2.088157 1.181749
			C -8.159680 -2.492240 -0.581962
			H -8.521056 -1.946193 -1.465031
			C -9.349014 -3.120328 0.151817
			H -7.503885 -3.287325 -0.964444
			H -10.043190 -2.351428 0.512958
			H -9.016693 -3.699634 1.022118
			H -9.911922 -3.795868 -0.502384

1-Decanol A2			
Monomer			
C	-3.764	0.450	0.000
H	-3.796	1.104	0.913
H	-3.796	1.104	-0.913
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-2.521	-1.074	0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
H	-0.011	-1.053	0.902
C	1.226	0.477	0.000
H	1.221	1.144	-0.903
C	2.482	-0.367	0.000
H	1.221	1.144	0.902
H	2.488	-1.034	-0.903
H	2.488	-1.034	0.902
C	3.727	0.496	0.000
H	3.719	1.162	-0.903
C	4.980	-0.341	0.000
H	3.718	1.163	0.902
H	5.018	-0.995	-0.904
H	5.017	-0.995	0.905
O	-4.875	-0.433	0.000
H	-5.665	0.119	-0.001
C	6.227	0.563	0.000
H	6.220	1.180	0.874
H	6.220	1.180	-0.874
C	7.495	-0.312	0.000
H	7.502	-0.930	-0.873
H	8.361	0.316	0.000
H	7.502	-0.930	0.874
Dimer			
H	0.372067	3.023060	-0.088831
O	1.122230	2.674763	-0.608316
C	1.990854	2.001517	0.287648
C	3.119731	1.363830	-0.514504
H	1.454905	1.220753	0.855604
H	2.417212	2.697526	1.032128
H	-1.821145	3.882311	0.945853
O	-1.317317	3.083938	0.725337
C	-2.124822	2.259957	-0.129421
H	-1.439586	1.494081	-0.503552
H	-2.473959	2.836160	-0.999702
C	-3.303460	1.632247	0.610073
H	2.676171	0.689569	-1.259743
C	4.123269	0.599668	0.356627
H	3.633318	2.154242	-1.078372
H	3.594797	-0.180542	0.925073
H	4.552587	1.281262	1.106235
C	5.260632	-0.046438	-0.445452
H	4.832854	-0.724536	-1.198692
C	6.259610	-0.820124	0.424255
H	5.795338	0.732509	-1.008878
H	5.722765	-1.598085	0.987666
H	6.686311	-0.141715	1.178155
C	7.398091	-1.468879	-0.373898
H	6.971952	-2.145262	-1.129834
C	8.391766	-2.246458	0.498650
H	7.938133	-0.691224	-0.934397
H	7.850843	-3.024144	1.058471
H	8.816565	-1.570053	1.255412
C	9.532379	-2.896032	-0.295414
H	9.109338	-3.572607	-1.053105
H	10.075809	-2.119137	-0.853995
C	-4.127798	0.690193	-0.277774
H	-3.953166	2.430584	1.000880
H	-2.917465	1.089251	1.482590
H	-4.494764	1.240901	-1.156498
C	-5.317112	0.052752	0.452574
H	-3.473903	-0.103295	-0.667896
H	-5.974090	0.846190	0.838668
H	-4.950149	-0.493168	1.334050
C	-6.135373	-0.899052	-0.429546
H	-6.500156	-0.353129	-1.312305
C	-7.325407	-1.537443	0.298497
H	-5.477116	-1.691981	-0.814443
H	-7.985310	-0.744467	0.680860
H	-6.960626	-2.080456	1.183040
C	-8.140583	-2.493592	-0.581965
H	-8.503835	-1.951059	-1.467523
C	-9.331814	-3.131241	0.144398
H	-7.480502	-3.287087	-0.962898
H	-9.993789	-2.338260	0.523614
H	-8.969707	-3.672707	1.031197
C	10.523144	-3.674445	0.579611
H	10.944985	-2.998372	1.337112
H	9.980409	-4.451774	1.136377
C	11.661116	-4.318839	-0.218679
H	12.349459	-4.866852	0.435062
H	11.273414	-5.026596	-0.961871
H	12.245183	-3.562461	-0.757522
C	-10.145317	-4.089804	0.734852
H	-10.505866	-3.549335	-1.621708
H	-9.484732	-4.883769	-1.111493
C	-11.334639	-4.719885	-0.002733
H	-11.892288	-5.400918	-0.655813
H	-11.003001	-5.293681	0.871503
H	-12.033441	-3.952361	0.352405

2-Heptanol <b>A3</b>			
Monomer			
C	-3.764	0.450	0.000
H	-3.796	1.104	0.913
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-2.521	-1.074	0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
H	-0.011	-1.053	0.902
O	-4.875	-0.433	0.000
H	-5.665	0.119	-0.001
C	1.248	0.492	0.000
H	1.254	1.109	-0.875
H	1.252	1.112	0.873
C	2.499	-0.406	0.003
H	2.495	-1.026	-0.870
H	3.377	0.205	0.002
H	2.493	-1.022	0.878
C	-3.808	1.346	-1.251
H	-3.179	2.199	-1.102
H	-3.462	0.793	-2.099
H	-4.813	1.670	-1.423
Dimer			
H	0.373	2.997	-0.092
O	1.123	2.648	-0.612
C	1.992	1.977	0.285
C	3.123	1.342	-0.517
H	1.458	1.194	0.852
H	-1.817	3.860	0.945
O	-1.316	3.061	0.724
C	-2.126	2.239	-0.130
H	-1.443	1.471	-0.505
C	-3.306	1.615	0.609
H	2.682	0.667	-1.263
C	4.128	0.580	0.355
H	3.636	2.134	-1.080
H	3.601	-0.202	0.922
H	4.554	1.262	1.106
C	5.269	-0.061	-0.446
H	4.844	-0.741	-1.200
C	6.269	-0.832	0.425
H	5.801	0.720	-1.008
H	5.735	-1.613	0.986
H	6.692	-0.152	1.180
C	-4.134	0.676	-0.278
H	-3.953	2.415	1.001
H	-2.921	1.070	1.481
H	-4.500	1.228	-1.156
C	-5.325	0.042	0.453
H	-3.483	-0.120	-0.669
H	-5.978	0.837	0.840
H	-4.959	-0.507	1.333
C	-6.147	-0.906	-0.430
H	-6.511	-0.358	-1.311
H	-5.492	-1.701	-0.816
C	-7.343	-1.544	0.302
H	-8.002	-0.774	0.646
H	-7.869	-2.189	-0.370
H	-6.990	-2.111	1.137
C	7.416	-1.477	-0.375
H	8.071	-1.994	0.295
H	7.012	-2.169	-1.084
H	7.963	-0.716	-0.892
C	2.583	2.948	1.324
H	2.950	2.393	2.163
H	3.388	3.497	0.881
H	1.824	3.627	1.650
C	-2.613	3.046	-1.348
H	-2.827	2.378	-2.156
H	-3.499	3.586	-1.087
H	-1.850	3.734	-1.647

5-Nonanol A4			
Monomer			
C	-3.764	0.450	0.000
H	-3.796	1.104	0.913
H	-3.796	1.104	-0.913
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-2.521	-1.074	0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
C	1.226	0.477	0.000
H	1.221	1.144	-0.903
C	2.482	-0.367	0.000
H	1.221	1.144	0.902
H	2.488	-1.034	-0.903
H	2.488	-1.034	0.902
C	3.727	0.496	0.000
H	3.719	1.162	-0.903
C	4.980	-0.341	0.000
H	3.718	1.163	0.902
H	5.018	-0.995	-0.904
H	5.017	-0.995	0.905
H	5.884	0.314	0.000
O	-0.010	-1.236	1.150
H	0.002	-2.155	0.871
C	-4.970	-0.508	0.000
H	-5.876	0.059	-0.039
H	-4.914	-1.151	-0.853
H	-4.956	-1.098	0.893
Dimer			
H	0.309234	2.434278	0.106119
O	1.124640	2.441793	-0.435009
C	2.121692	1.666554	0.226875
C	1.606806	0.241264	0.497179
H	2.369222	2.132354	1.199384
H	-1.801004	3.325769	0.615759
O	-1.484754	2.412108	0.717416
C	-2.280785	1.581095	-0.161512
H	-1.833113	0.586208	-0.045588
C	-3.733190	1.538468	0.326333
H	0.642204	0.331195	1.018916
C	2.534307	-0.647469	1.337095
H	1.393961	-0.239965	-0.468290
H	2.809764	-0.117636	2.261673
H	3.474460	-0.828221	0.798277
C	1.905176	-1.998959	1.702856
H	0.970562	-1.824659	2.254823
H	1.619641	-2.526262	0.781398
C	-3.891612	1.019404	1.760937
H	-4.318500	0.909492	-0.357366
H	-4.160618	2.551605	0.245937
H	-3.486126	-0.001768	1.820053
C	-5.345833	1.014531	2.248722
H	-3.276414	1.629020	2.434325
H	-5.957577	0.405540	1.567628
H	-5.752115	2.034687	2.189996
C	3.371058	1.706957	-0.656244
H	4.152077	1.082101	-0.202971
H	3.118256	1.248825	-1.623640
C	-2.106629	2.038078	-1.616381
H	-2.548695	3.041389	-1.730449
H	-1.029302	2.143774	-1.793667
C	3.919177	3.121186	-0.883774
H	3.127526	3.742833	-1.318281
H	4.168428	3.573910	0.088258
C	5.159062	3.151747	-1.786191
H	5.945911	2.519242	-1.349645
H	4.911230	2.699382	-2.757199
C	5.707648	4.564989	-2.008594
H	5.997117	5.031329	-1.058503
H	6.590859	4.556858	-2.658067
H	4.955125	5.211737	-2.476454
C	2.831268	-2.891239	2.535520
H	3.108940	-2.403397	3.478106
H	2.352618	-3.845749	2.783202
H	3.758960	-3.114619	1.994187
C	-2.704372	1.094871	-2.670364
H	-2.311371	0.079232	-2.512803
H	-3.793929	1.024185	-2.547729
C	-2.393090	1.535814	-4.107304
H	-2.778878	2.553103	-4.264716
H	-1.304023	1.600552	-4.236401
C	-2.980890	0.597073	-5.165571
H	-2.741175	0.938174	-6.179070
H	-2.587026	-0.420818	-5.055250
H	-4.073428	0.539304	-5.083533
C	-5.497424	0.487960	3.679385
H	-6.544987	0.499166	4.001275
H	-5.134255	-0.543807	3.762536
H	4.923817	1.097433	4.388396

2,4-Dimethyl-4-heptanol A5						
Monomer				Dimer		
C	-2.512	-0.409	0.000	C	2.267690	2.208619
C	-1.272	0.458	0.000	H	2.338816	2.640858
H	-2.521	-1.074	0.902	C	3.501955	2.625373
H	-1.277	1.125	-0.902	H	4.402503	2.268674
C	-0.016	-0.387	0.000	C	3.639874	4.138477
H	-1.277	1.124	0.903	H	3.469029	2.101924
H	-0.012	-1.053	-0.903	H	2.736499	4.457222
C	1.226	0.477	0.000	O	1.128704	2.755563
H	1.221	1.144	-0.903	H	0.331810	2.578073
C	2.482	-0.367	0.000	C	4.853312	4.410116
H	1.221	1.144	0.902	H	4.937001	5.476357
H	2.488	-1.034	0.902	H	5.788121	4.108862
O	-0.010	-1.236	1.150	H	4.782515	3.856045
H	0.002	-2.155	0.871	C	3.732596	4.957223
C	-3.782	0.462	0.000	H	2.827008	4.856244
H	-3.790	1.080	-0.874	H	4.588398	4.639560
H	-4.647	-0.168	0.000	H	3.864159	6.023358
H	-3.791	1.080	0.874	C	-1.834043	4.026894
C	3.748	0.510	0.000	C	-1.988329	2.521517
H	4.615	-0.116	0.000	H	-1.116194	4.443167
H	3.753	1.128	-0.874	H	-2.732100	2.094145
H	3.753	1.128	0.874	C	-2.399361	2.159866
C	2.490	-1.282	-1.239	H	-1.031390	2.027375
H	1.953	-2.181	-1.021	H	-3.378995	2.602472
H	2.024	-0.777	-2.058	C	-2.467816	0.646116
H	3.500	-1.524	-1.497	H	-3.168872	0.233845
C	-2.524	-1.323	-1.240	C	-2.879036	0.177315
H	-2.066	-0.815	-2.062	H	-1.480255	0.229187
H	-1.981	-2.220	-1.027	H	-2.185808	0.639236
H	-3.535	-1.569	-1.491	O	-1.443031	2.667863
				H	-1.507954	3.635878
				C	-1.215695	4.223595
				H	-1.859312	3.796929
				H	-1.079887	5.287972
				H	-0.236605	3.736474
				C	-2.732826	-1.347783
				H	-2.980031	-1.697968
				H	-3.403765	-1.860377
				H	-1.708632	-1.669320
				C	-4.304788	0.610385
				H	-4.413873	1.700405
				H	-5.038847	0.210737
				H	-4.575317	0.239339
				C	-3.157804	4.797776
				H	-3.900697	4.413072
				H	-3.596632	4.726426
				H	-3.012708	5.862561
				C	3.226258	-0.095701
				H	4.175956	0.086688
				C	2.947739	-1.605341
				H	2.007549	-1.854478
				H	2.864186	-1.949675
				H	3.748472	-2.179459
				C	3.397181	0.352122
				H	4.169835	-0.241867
				H	3.687313	1.405471
				H	2.461609	0.221409
				C	2.112842	0.680573
				H	2.000860	0.282582
				H	1.158810	0.482030
						1.167561

Menthol A6			
Monomer		Dimer	
C	-3.323	-1.241	0.000
C	-1.808	-1.241	0.000
C	-1.256	0.170	0.000
C	-1.806	0.975	1.161
C	-3.321	0.975	1.160
C	-3.874	-0.435	1.159
H	-0.137	0.136	0.063
H	-1.433	-1.791	-0.902
H	-3.696	-0.808	-0.965
H	-3.699	-2.295	0.064
H	-1.430	2.029	1.099
H	-3.696	1.525	2.062
H	-3.608	-0.941	2.124
H	-4.992	-0.400	1.094
H	-1.525	0.676	-0.965
C	-1.297	-1.990	1.245
H	-1.667	-2.994	1.234
H	-0.227	-2.004	1.238
H	-1.641	-1.492	2.127
C	-3.831	1.726	-0.084
H	-3.493	1.224	-0.967
C	-3.289	3.167	-0.073
H	-2.361	3.203	-0.605
H	-3.995	3.818	-0.544
H	-3.133	3.482	0.938
C	-5.371	1.757	-0.069
H	-5.716	1.846	0.940
H	-5.716	2.593	-0.640
H	-5.750	0.852	-0.497
O	-1.331	0.420	2.390
H	-0.765	1.058	2.832
C	4.103936	1.743444	-1.996220
C	2.973847	0.751234	-2.351339
C	2.176599	0.378587	-1.079725
C	1.734242	1.567120	-0.212511
C	2.917794	2.509692	0.143185
C	3.600204	2.941000	-1.174587
H	1.284149	-0.200205	-1.348838
H	3.449246	-0.172763	-2.712755
H	4.883759	1.212732	-1.430908
H	4.585664	2.099780	-2.916849
H	1.296394	1.161392	0.710468
H	2.474887	3.411379	0.598422
H	2.860112	3.496166	-1.761608
H	4.421963	3.638291	-0.978121
H	2.797975	-0.277845	-0.454881
C	2.081083	1.264385	-3.494186
H	2.681359	1.463033	-4.391423
H	1.325423	0.514199	-3.757597
H	1.548285	2.176698	-3.217229
C	3.881028	1.915266	1.212873
H	4.219341	0.925745	0.874276
C	3.179976	1.724305	2.570996
H	2.326613	1.041637	2.517002
H	3.878254	1.316790	3.311771
H	2.813381	2.684912	2.957218
C	5.136487	2.781346	1.417557
H	4.865491	3.814039	1.675510
H	5.745546	2.385651	2.239125
H	5.771061	2.815518	0.526812
O	0.720200	2.283340	-0.921926
H	0.263241	2.869357	-0.285063
C	-4.057536	2.254883	1.891188
C	-3.085257	1.529659	0.934343
C	-2.672596	2.467662	-0.224993
C	-2.250047	3.882777	0.188571
C	-3.261804	4.570634	1.137454
C	-3.527834	3.628184	2.333373
H	-1.857363	2.020995	-0.805448
H	-3.632237	0.691572	0.479096
H	-5.029050	2.380882	1.392705
H	-4.245707	1.628358	2.772905
H	-2.119179	4.479218	-0.723059
H	-2.770509	5.479578	1.528128
H	-2.584338	3.495024	2.875045
H	-4.220334	4.092810	3.042567
H	-3.522802	2.577812	-0.911532
C	-1.883224	0.919137	1.675013
H	-2.221775	0.189434	2.421223
H	-1.222471	0.398573	0.972208
H	-1.284129	1.678794	2.184049
C	-4.543172	5.066086	0.402186
H	-5.007528	4.211915	-0.109655
C	-4.221764	6.128391	-0.666149
H	-3.582581	5.748213	-1.468695
H	-5.144061	6.492246	-1.133427
H	-3.717054	6.994619	-0.217602
C	-5.585294	5.637787	1.379391
H	-5.155886	6.446507	1.985528
H	-6.435397	6.054813	0.827440
H	-5.979761	4.879822	2.062536
O	-0.954436	3.783397	0.832873
H	-0.677413	4.681995	1.074289

4-Pentylcyclohexanol A7			
Monomer		Dimer	
H	0.372	2.995	-0.092
O	1.123	2.648	-0.612
C	2.000	1.971	0.293
C	2.107	0.512	-0.100
C	3.358	2.644	0.283
H	1.552	2.050	1.318
C	3.092	-0.227	0.783
H	2.437	0.437	-1.169
H	1.099	0.028	-0.024
C	4.342	1.906	1.167
H	3.750	2.675	-0.767
H	3.255	3.702	0.637
C	4.449	0.446	0.776
H	3.195	-1.285	0.428
H	2.698	-0.260	1.833
H	5.351	2.390	1.091
H	4.012	1.982	2.236
H	4.899	0.366	-0.249
C	5.391	-0.282	1.753
H	5.482	-1.308	1.463
H	4.991	-0.225	2.743
C	6.779	0.385	1.720
H	6.775	1.248	2.353
H	7.010	0.679	0.718
C	9.207	0.090	2.304
H	9.220	0.747	3.148
H	9.373	0.655	1.410
C	10.319	-0.965	2.457
H	11.105	-0.758	1.761
H	10.707	-0.932	3.454
H	9.917	-1.937	2.263
C	7.840	-0.613	2.220
H	7.860	-1.431	1.531
H	7.606	-0.977	3.199
			H 0.767409 1.674088 -0.418660
			O 1.445803 1.083554 -0.805621
			H -0.672729 3.548067 -0.377708
			O -0.790585 2.657738 -0.008417
			C 2.245893 0.564979 0.249913
			C 3.241373 -0.419753 -0.361619
			C 2.986168 1.673534 1.011040
			H 1.612781 0.014991 0.970905
			C 4.197073 -0.999822 0.691890
			H 3.812548 0.112458 -1.135475
			H 2.692005 -1.220774 -0.871171
			C 3.937581 1.096805 2.070475
			H 3.550886 2.268335 0.279236
			H 2.258156 2.350460 1.479252
			C 4.947701 0.097882 1.472322
			H 4.906911 -1.676940 0.202193
			H 3.627138 -1.613780 1.407058
			H 4.473040 1.910575 2.577371
			H 3.350366 0.584016 2.848113
			H 5.573440 0.650310 0.751619
			C -1.882103 2.031315 -0.713813
			C -1.666522 2.052175 -2.229460
			C -1.992186 0.598613 -0.199683
			H -2.810686 2.574174 -0.470683
			C -2.782758 1.292497 -2.965542
			H -0.692478 1.592433 -2.442791
			H -1.619041 3.091398 -2.587631
			C -3.106533 -0.164600 -0.931008
			H -1.024838 0.106687 -0.365350
			H -2.171985 0.610064 0.882078
			C -2.924084 -0.157718 -2.461756
			H -2.579960 1.309078 -4.042375
			H -3.741584 1.815170 -2.821824
			H -3.144457 -1.198741 -0.565586
			H -4.081608 0.284445 -0.685755
			H -1.979784 -0.680310 -2.685266
			C 5.873199 -0.467635 2.565888
			H 6.288404 0.374026 3.140319
			H 5.269088 -1.049061 3.279509
			C 7.035715 -1.335636 2.063440
			H 6.646748 -2.217450 1.536413
			H 7.618295 -0.769255 1.321240
			C 7.968129 -1.801197 3.189794
			H 7.383907 -2.358529 3.937455
			H 8.371078 -0.922507 3.715606
			C 9.131050 -2.676202 2.704311
			H 8.728312 -3.554359 2.179607
			H 9.716122 -2.119490 1.958352
			C 10.054549 -3.137849 3.836397
			H 10.873901 -3.759643 3.457595
			H 10.500017 -2.282010 4.358627
			H 9.505031 -3.727981 4.580241
			C -4.065689 -0.926793 -3.151839
			H -4.159566 -1.910942 -2.669240
			H -5.016918 -0.406343 -2.960401
			C -3.898605 -1.138457 -4.663383
			H -3.884512 -0.169131 -5.180417
			H -2.920509 -1.602910 -4.859160
			C -5.003456 -2.011405 -5.273541
			H -5.010777 -2.990880 -4.772206
			H -5.983751 -1.557412 -5.063970
			C -4.857721 -2.220068 -6.786321
			H -4.858841 -1.242062 -7.288475
			H -3.876203 -2.668140 -6.997140
			C -5.959679 -3.101150 -7.383699
			H -6.951613 -2.662813 -7.217732
			H -5.829062 -3.228962 -8.464403
			H -5.959992 -4.099156 -6.928140

2-methyl-2-hexanol <b>A8</b>			
Monomer			Dimer
C	-3.764	0.450	0.000
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-2.521	-1.074	0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
H	-0.011	-1.053	0.902
O	-4.875	-0.433	0.000
H	-5.665	0.119	-0.001
C	-3.808	1.346	-1.251
H	-3.179	2.199	-1.102
H	-3.462	0.793	-2.099
H	-4.813	1.670	-1.423
C	1.248	0.492	0.000
H	1.230	1.144	0.849
H	2.116	-0.131	0.048
H	1.277	1.075	-0.897
C	-3.808	1.346	1.251
H	-4.801	1.722	1.387
H	-3.523	0.775	2.110
H	-3.131	2.166	1.127
			H 0.372 2.995 -0.092
			O 1.123 2.648 -0.612
			H -1.817 3.861 0.944
			O -1.316 3.061 0.724
			C 2.000 1.971 0.292
			C -2.123 2.242 -0.127
			C 3.203 1.405 -0.486
			H 2.855 0.717 -1.228
			H 3.729 2.206 -0.961
			C -2.464 3.019 -1.412
			H -2.999 2.380 -2.084
			H -3.070 3.866 -1.167
			H -1.560 3.351 -1.879
			C 2.499 2.961 1.361
			H 1.668 3.316 1.934
			H 3.193 2.467 2.008
			H 2.983 3.787 0.884
			C -1.349 0.961 -0.491
			H -1.104 0.426 0.403
			H -1.956 0.345 -1.121
			H -0.449 1.223 -1.008
			C -3.424 1.865 0.606
			H -3.185 1.349 1.513
			H -3.975 2.753 0.835
			C 1.242 0.818 0.975
			H 0.868 0.146 0.231
			H 1.907 0.292 1.628
			H 0.425 1.214 1.541
			C 4.147 0.676 0.488
			H 4.496 1.364 1.229
			H 3.619 -0.124 0.965
			C 5.347 0.106 -0.290
			H 4.998 -0.583 -1.031
			H 5.875 0.905 -0.768
			C 6.292 -0.623 0.684
			H 7.126 -1.019 0.143
			H 6.641 0.066 1.424
			H 5.764 -1.422 1.161
			C -4.275 0.951 -0.295
			H -3.768 0.019 -0.438
			H -4.425 1.424 -1.243
			C -5.639 0.697 0.373
			H -5.488 0.248 1.333
			H -6.157 1.626 0.491
			C -6.475 -0.249 -0.509
			H -5.954 -1.176 -0.631
			H -7.420 -0.429 -0.043
			H -6.630 0.201 -1.468

3-Ethyl-3-pentanol A9			
Monomer		Dimer	
C	-0.012	-0.336	-0.023
C	0.008	0.947	-0.874
H	0.864	0.935	-1.516
H	-0.881	0.997	-1.466
C	-1.270	-0.343	0.866
H	-1.284	-1.234	1.457
H	-1.255	0.512	1.510
C	0.078	2.176	0.052
H	0.509	2.999	-0.479
H	-0.908	2.436	0.374
H	0.683	1.946	0.904
C	-2.527	-0.295	-0.022
H	-3.359	-0.700	0.516
H	-2.735	0.719	-0.291
H	-2.360	-0.872	-0.908
C	2.502	-0.380	-0.024
H	2.286	-0.873	-0.949
H	2.799	0.629	-0.219
H	3.295	-0.895	0.477
C	1.245	-0.384	0.865
H	1.158	-1.270	1.458
H	1.332	0.468	1.508
O	-0.031	-1.478	-0.883
H	-0.047	-2.279	-0.353
C	-2.524	-0.052	-0.322
C	-2.165	0.167	1.159
H	-1.391	-0.515	1.442
H	-1.825	1.171	1.299
C	-3.603	0.963	-0.745
H	-3.853	0.811	-1.773
H	-4.477	0.825	-0.142
C	-3.411	-0.079	2.031
H	-3.107	-0.348	3.021
H	-4.001	0.813	2.069
H	-3.991	-0.873	1.608
C	-3.067	2.394	-0.551
H	-3.601	3.064	-1.192
H	-3.201	2.690	0.468
H	-2.025	2.420	-0.796
C	-1.981	-2.496	-0.096
H	-1.011	-2.086	-0.290
H	-2.075	-2.708	0.948
H	-2.103	-3.400	-0.656
C	-3.060	-1.482	-0.516
H	-3.329	-1.563	-1.548
H	-3.917	-1.690	0.093
O	-1.357	0.132	-1.128
H	-1.581	-0.001	-2.052
C	2.320	0.074	-0.025
C	3.428	-0.088	1.032
H	3.362	0.709	1.742
H	3.308	-1.024	1.535
C	2.383	-1.101	-1.020
H	1.614	-0.988	-1.754
H	3.339	-1.107	-1.502
C	4.805	-0.049	0.343
H	5.548	0.263	1.047
H	5.048	-1.024	-0.024
H	4.775	0.642	-0.473
C	2.182	-2.427	-0.263
H	1.822	-3.173	-0.940
H	3.114	-2.744	0.156
H	1.469	-2.284	0.523
C	2.455	2.573	0.212
H	1.790	2.325	1.013
H	3.431	2.764	0.606
H	2.096	3.447	-0.291
C	2.521	1.399	-0.782
H	1.752	1.439	-1.524
H	3.479	1.465	-1.257
O	1.044	0.083	0.620
H	0.352	0.180	-0.039

1,1,1-Trifluoro-2-octanol A10			
Monomer		Dimer	
C	-3.764	0.450	0.000
C	-2.512	-0.409	0.000
C	-1.272	0.458	0.000
H	-2.521	-1.074	-0.902
H	-1.277	1.125	-0.902
C	-0.016	-0.387	0.000
H	-1.277	1.124	0.903
H	-0.012	-1.053	-0.903
H	-0.011	-1.053	0.902
C	1.226	0.477	0.000
H	1.221	1.144	-0.903
C	2.482	-0.367	0.000
H	1.221	1.144	0.902
H	2.488	-1.034	-0.903
H	2.488	-1.034	0.902
C	3.727	0.496	0.000
H	3.719	1.162	-0.903
C	4.980	-0.341	0.000
H	3.718	1.163	0.902
H	5.018	-0.995	-0.904
H	5.017	-0.995	0.905
H	5.884	0.314	0.000
O	-2.523	-1.258	1.151
H	-2.880	-2.116	0.912
F	-4.821	-0.390	0.000
F	-3.802	1.236	1.097
F	-3.802	1.236	-1.097
H	0.798794	1.497284	-1.177693
O	1.643826	1.980467	-1.267453
C	1.664179	3.023891	-0.318652
C	3.098249	3.559163	-0.257010
H	1.350122	2.674530	0.676380
H	-0.394634	-0.528250	-0.452998
O	-0.650351	0.324359	-0.843159
C	-1.933564	0.683497	-0.329573
H	-2.145660	1.671328	-0.744147
C	-3.001553	-0.330720	-0.760923
H	3.732460	2.677192	-0.106593
C	3.390494	4.598099	0.834087
H	3.354423	3.959212	-1.245502
H	3.066115	4.207710	1.809964
H	2.802626	5.506658	0.659879
C	4.878592	4.965591	0.905242
H	5.470784	4.056754	1.088879
C	5.197200	6.006088	1.986226
H	5.204826	5.347188	-0.073602
H	4.874963	5.623040	2.966161
H	4.600654	6.912589	1.804827
C	6.682187	6.384547	2.055259
H	7.279058	5.479037	2.235636
H	7.003071	6.769325	1.076741
C	-4.463953	0.109042	-0.589660
H	-2.829144	-1.278148	-0.230902
H	-2.803967	-0.524905	-1.822155
H	-4.697077	0.240884	0.471964
C	-5.448287	-0.893334	-1.207262
H	-4.603758	1.092636	-1.061008
H	-5.293153	-1.882711	-0.751855
H	-5.224428	-1.012539	-2.277547
C	-6.916130	-0.482025	-1.036868
H	-7.138359	-0.364792	0.034110
C	-7.906262	-1.478916	-1.652845
H	-7.071228	0.508947	-1.488845
H	-7.749128	-2.469728	-1.203181
H	-7.685768	-1.593002	-2.723733
C	0.641879	4.094139	-0.718355
C	-1.803090	0.849371	1.185864
C	6.991575	7.423008	3.138468
H	6.713387	7.052963	4.133105
H	8.058785	7.671739	3.162401
H	6.436516	8.353417	2.965925
C	-9.369914	-1.063188	-1.474142
H	-9.566053	-0.091042	-1.942965
H	-10.051087	-1.793634	-1.925139
H	-9.630560	-0.976028	-0.412167
F	-0.571790	3.530880	-0.970457
F	0.448143	5.002298	0.267042
F	0.997458	4.769248	-1.827209
F	-0.872940	1.769998	1.506847
F	-2.959043	1.226161	1.766077
F	-1.418487	-0.323317	1.754517