Supporting information: Histogram-free reweighting with grand canonical Monte Carlo: Post-simulation optimization of non-bonded potentials for phase equilibria

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SI.I Bonded parameters

Table SI.I: Equilibrium (fixed) bond lengths (r_{eq}) . CH_x and CH_y represent CH_3 , $CH_2(sp^3)$, $CH(sp^3)$, or $C(sp^3)$ sites.

Bond sites	$r_{ m eq}$ (nm)				
	TraPPE	TraPPE MiPPE NEI			
$CH_x\text{-}CH_y$	0.154	0.154	0.154		
$C(sp)$ - CH_x	_	0.146	_		
CH≡CH	_	0.121	_		
C≡CH	_	0.121	_		

Table SI.II: Equilibrium bond angles (θ_{eq}) and force constants $(k_{\theta}/k_{\rm B})$, where $k_{\rm B}$ is the Boltzmann constant.

Bending sites	$\theta_{ m eq}$ (degrees)			$k_{ heta}/k_{ m B}$ (K/rad ²)
	TraPPE	MiPPE	NERD	
CH_x - CH_2 - CH_y	114.0	114.0	114.0	62500
CH_x - CH - CH_y	112.0	112.0	109.5	62500
$CH_x\text{-}C\text{-}CH_y$	109.5	109.5	109.5	62500
CH_x - CH_2 - $C(sp)$	_	112	<u> </u>	62500
CH_x - $C(sp)\equiv CH$	_	180	_	30800
CH_x - $C(sp)\equiv C$	_	180	_	30800

Table SI.III: Fourier constants $(c_n/k_{\rm B})$ in units of K.

Torsion sites	$c_0/k_{ m B}$	$c_1/k_{ m B}$	$c_2/k_{ m B}$	$c_3/k_{\rm B}$
CH_x - CH_2 - CH_y	0.0	355.03	-68.19	791.32
CH_x - CH_2 - CH - CH_y	-251.06	428.73	-111.85	441.27
$CH_x ext{-}CH_2 ext{-}C-CH_y$	0.0	0.0	0.0	461.29
$CH_x ext{-}CH ext{-}CH ext{-}CH_y$	-251.06	428.73	-111.85	441.27
CH_x - CH_2 - $C(sp)$	94.88	162.00	-205.40	980.40
CH_x - CH_2 - $C(sp)\equiv C(sp)$	0	0	0	0
CH_x - CH_2 - $C(sp)\equiv CH(sp)$	0	0	0	0
CH_x - $C(sp)\equiv C(sp)$ - CH_y	0	0	0	0

SI.II Compiler and hardware

With the exception of the 20 replicates performed for MiPPE cyclohexane, all simulations are run on a Linux 4.4.0-112-generic x86_64 on an Intel(R) Xeon(R) CPU E5-2699 v4 @ 2.20GHz machine. On this machine, GOMC was erroneously compiled using the suboptimal GNU compiler collection (GCC) instead of the preferred Intel compiler. GOMC compiled with the Intel compiler typically runs approximately twice as fast as GOMC compiled with the GCC compiler.

The 20 replicate simulations for MiPPE cyclohexane utilize several different machine hardware architectures, listed in Table SI.IV. GOMC was compiled with the Intel compiler on each of these machines.

Table SI.IV: Machine hardware for 20 replicate simulations of MiPPE cyclohexane

Intel(R) Core(TM) i7-4790K CPU @ 4.00GHz Intel(R) Core(TM) i5-3570 CPU @ 3.40GHz Intel(R) Core(TM) i5-2500K CPU @ 3.30GHz Intel(R) Xeon(R) CPU X5450 @ 3.00GHz Intel(R) Xeon(R) CPU X5355 @ 2.66GHz Intel(R) Xeon(R) CPU E5-2640 v3 @ 2.60GHz Intel(R) Core(TM)2 Quad CPU Q6600 @ 2.40GHz

SI.III ϵ -scaling

SI.III.1 Tabulated ψ values

Table SI.V: Optimal ϵ -scaling parameter (ψ) values and corresponding scoring function. Abbreviations correspond to those in Figure 2.

Molecular name	Abbreviation	Optimal ψ	Optimal score					
	Branched alkanes							
2-methylpropane	$2MC_3$	1.0015	0.3883					
2-methylbutane	$2MC_4$	1.0025	0.4281					
2-methylpentane	$2MC_5$	1.0020	0.4770					
3-methylpentane	$3MC_5$	1.0103	0.4050					
2,2-dimethylpropane	$22DMC_3$	1.0035	0.5132					
2,2-dimethylbutane	$22DMC_4$	0.9985	0.5445					
2,3-dimethylbutane	$23DMC_4$	1.0000	0.4724					
2,2,4-trimethylpentane	$234TMC_5$	1.0005	0.4367					
	Alkynes							
1-ethyne	C_2	1.0005	0.2931					
1-propyne	C_3	0.9965	0.3307					
1-butyne	$1C_4$	1.0063	1.143					
2-butyne	$2C_4$	1.0031	0.3191					
1-pentyne	$1C_5$	1.0087	1.8505					
2-pentyne	$2C_5$	1.0186	1.3801					
1-hexyne	$1C_6$	1.0063	1.908					
2-hexyne	$2C_6$	1.0228	1.0594					
1-heptyne	$1C_7$	1.0066	0.8415					
1-octyne	$1C_8$	1.0034	0.9777					
1-nonyne	1C ₉	1.0000	0.9128					

SI.III.2 Tabulated phase equilibria for optimal ψ

SI.III.2.1 Branched alkanes

Table SI.VI: GCMC-MBAR results for 2-methylpropane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
390	$390.5_{4.5}$	$85.7_{2.5}$	$2.718_{0.030}$	$9.90_{0.20}$	$0.569_{0.011}$
380	$418.0_{2.3}$	$65.5_{2.0}$	$2.272_{0.017}$	$11.81_{0.14}$	$0.639_{0.016}$
370	$440.86_{0.43}$	$51.3_{1.3}$	$1.8856_{8.8e-3}$	$13.31_{0.11}$	$0.695_{0.015}$
360	$460.16_{0.83}$	$40.72_{0.64}$	$1.5515_{4.8e-3}$	$14.524_{0.080}$	$0.740_{0.011}$
350	$477.25_{0.86}$	$32.43_{0.28}$	$1.2638_{3.8e-3}$	$15.559_{0.056}$	$0.78_{0.01}$
340	$492.86_{0.74}$	$25.78_{0.16}$	$1.0177_{3.4e-3}$	$16.473_{0.048}$	$0.81_{0.01}$
330	$507.34_{0.80}$	$20.38_{0.14}$	$0.8090_{3.3e-3}$	$17.292_{0.053}$	$0.84_{0.01}$
320	$521.13_{0.61}$	$15.97_{0.11}$	$0.6336_{3.4e-3}$	$18.042_{0.053}$	$0.87_{0.01}$
310	$534.54_{0.29}$	$12.374_{0.074}$	$0.4881_{3.6e-3}$	$18.745_{0.042}$	$0.89_{0.01}$
300	$547.12_{0.22}$	$9.452_{0.060}$	$0.3690_{3.5e-3}$	$19.389_{0.035}$	$0.910_{0.010}$
290	$558.97_{0.19}$	$7.101_{0.076}$	$0.2732_{3.2e-3}$	$19.980_{0.039}$	$0.927_{0.012}$
280	$570.80_{0.34}$	$5.230_{0.094}$	$0.1975_{2.7e-3}$	$20.552_{0.048}$	$0.943_{0.019}$
270	$582.96_{0.45}$	$3.76_{0.10}$	$0.1389_{2.5e-3}$	$21.122_{0.071}$	$0.956_{0.034}$
260	$593.85_{0.25}$	$2.63_{0.10}$	$0.0946_{2.7e-3}$	$21.62_{0.12}$	$0.966_{0.060}$
250	$603.32_{0.94}$	$1.787_{0.089}$	$0.0623_{3.4e-3}$	$22.06_{0.24}$	$0.97_{0.10}$

Table SI.VII: GCMC-MBAR results for 2-methylbutane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
440	$403.0_{2.4}$	$84.5_{1.4}$	$2.490_{0.015}$	$11.80_{0.11}$	$0.58_{0.01}$
430	$429.5_{1.4}$	$67.3_{1.1}$	$2.1160_{0.0099}$	$13.658_{0.094}$	$0.63_{0.01}$
420	$451.56_{0.83}$	$53.86_{0.75}$	$1.7868_{0.0058}$	$15.259_{0.071}$	$0.69_{0.01}$
410	$470.40_{0.84}$	$43.55_{0.44}$	$1.4985_{0.0031}$	$16.608_{0.052}$	$0.73_{0.01}$
400	$487.2_{1.1}$	$35.38_{0.25}$	$1.2466_{0.0017}$	$17.769_{0.044}$	$0.76_{0.01}$
390	$502.5_{1.3}$	$28.74_{0.17}$	$1.0277_{0.0016}$	$18.796_{0.044}$	$0.80_{0.01}$
380	$516.7_{1.4}$	$23.25_{0.13}$	$0.8385_{0.0025}$	$19.723_{0.041}$	$0.82_{0.01}$
370	$530.1_{1.2}$	$18.699_{0.091}$	$0.6765_{0.0033}$	$20.574_{0.033}$	$0.85_{0.01}$
360	$542.77_{0.83}$	$14.917_{0.061}$	$0.5388_{0.0038}$	$21.360_{0.027}$	$0.87_{0.01}$
350	$554.94_{0.49}$	$11.782_{0.058}$	$0.4234_{0.0039}$	$22.094_{0.027}$	$0.89_{0.01}$
340	$566.86_{0.43}$	$9.196_{0.079}$	$0.3274_{0.0037}$	$22.793_{0.033}$	$0.91_{0.01}$
330	$578.69_{0.60}$	$7.08_{0.10}$	$0.2489_{0.0032}$	$23.467_{0.052}$	$0.925_{0.011}$
320	$589.78_{0.55}$	$5.36_{0.12}$	$0.1856_{0.0027}$	$24.087_{0.073}$	$0.939_{0.019}$
310	$599.68_{0.39}$	$3.99_{0.12}$	$0.1354_{0.0024}$	$24.634_{0.10}$	$0.951_{0.031}$
300	$609.37_{0.41}$	$2.90_{0.11}$	$0.0965_{0.0025}$	$25.16_{0.14}$	$0.961_{0.050}$
290	$619.67_{0.47}$	$2.065_{0.092}$	$0.0670_{0.0030}$	$25.71_{0.20}$	$0.971_{0.076}$

Table SI.VIII: GCMC-MBAR results for 2-methylpentane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	$424.43_{0.72}$	$75.06_{0.98}$	$2.0415_{0.0063}$	$14.355_{0.063}$	$0.60_{0.01}$
460	$446.63_{0.44}$	$59.92_{0.85}$	$1.7383_{0.0032}$	$16.259_{0.079}$	$0.65_{0.01}$
450	$466.02_{0.34}$	$48.40_{0.58}$	$1.4722_{0.0024}$	$17.883_{0.082}$	$0.70_{0.01}$
440	$483.12_{0.37}$	$39.46_{0.30}$	$1.2388_{0.0030}$	$19.257_{0.075}$	$0.74_{0.01}$
430	$498.62_{0.46}$	$32.30_{0.13}$	$1.0347_{0.0032}$	$20.456_{0.064}$	$0.77_{0.01}$
420	$513.07_{0.46}$	$26.404_{0.097}$	$0.8567_{0.0028}$	$21.538_{0.049}$	$0.80_{0.01}$
410	$526.79_{0.34}$	$21.50_{0.12}$	$0.7026_{0.0021}$	$22.536_{0.031}$	$0.83_{0.01}$
400	$539.84_{0.33}$	$17.39_{0.13}$	$0.5701_{0.0016}$	$23.462_{0.027}$	$0.85_{0.01}$
390	$552.17_{0.47}$	$13.96_{0.12}$	$0.4573_{0.0015}$	$24.320_{0.043}$	$0.87_{0.01}$
380	$563.75_{0.44}$	$11.10_{0.10}$	$0.3622_{0.0018}$	$25.113_{0.052}$	$0.890_{0.011}$
370	$574.77_{0.27}$	$8.738_{0.085}$	$0.2829_{0.0022}$	$25.853_{0.051}$	$0.907_{0.015}$
360	$585.55_{0.30}$	$6.794_{0.076}$	$0.2177_{0.0027}$	$26.562_{0.058}$	$0.922_{0.021}$
350	$596.34_{0.30}$	$5.208_{0.079}$	$0.1647_{0.0031}$	$27.257_{0.083}$	$0.936_{0.030}$
340	$606.99_{0.31}$	$3.928_{0.084}$	$0.1222_{0.0035}$	$27.93_{0.12}$	$0.948_{0.045}$
330	$617.01_{0.30}$	$2.909_{0.083}$	$0.0888_{0.0039}$	$28.55_{0.18}$	$0.959_{0.067}$
320	$626.41_{0.21}$	$2.111_{0.076}$	$0.0631_{0.0043}$	$29.13_{0.26}$	$0.968_{0.099}$

Table SI.IX: GCMC-MBAR results for 3-methylpentane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(K)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
480	$428.2_{1.5}$	75_{12}	$2.171_{0.060}$	$14.81_{0.93}$	$0.625_{0.090}$
470	$449.9_{1.6}$	$61.1_{8.5}$	$1.863_{0.021}$	$16.56_{0.84}$	$0.672_{0.089}$
460	$468.1_{1.3}$	$50.3_{4.0}$	$1.5910_{0.0071}$	$18.01_{0.53}$	$0.712_{0.060}$
450	$484.5_{1.2}$	$41.6_{1.1}$	$1.350_{0.014}$	$19.28_{0.23}$	$0.747_{0.027}$
440	$500.3_{1.1}$	$34.45_{0.21}$	$1.137_{0.015}$	$20.464_{0.074}$	$0.78_{0.01}$
430	$515.5_{1.0}$	$28.41_{0.46}$	$0.950_{0.013}$	$21.563_{0.068}$	$0.81_{0.01}$
420	$529.38_{0.62}$	$23.34_{0.40}$	$0.786_{0.010}$	$22.556_{0.054}$	$0.83_{0.01}$
410	$541.89_{0.69}$	$19.07_{0.30}$	$0.6449_{0.0079}$	$23.443_{0.043}$	$0.85_{0.01}$
400	$553.62_{0.90}$	$15.49_{0.24}$	$0.5233_{0.0062}$	$24.264_{0.049}$	$0.88_{0.01}$
390	$565.07_{0.63}$	$12.48_{0.20}$	$0.4198_{0.0048}$	$25.051_{0.039}$	$0.89_{0.01}$
380	$575.97_{0.45}$	$9.95_{0.17}$	$0.3326_{0.0039}$	$25.789_{0.043}$	$0.912_{0.014}$
370	586_{95}	$7.8_{1.6}$	$0.260_{0.066}$	$26.5_{4.1}$	$0.93_{0.18}$
360	$597.06_{0.27}$	$6.10_{0.13}$	$0.2001_{0.0035}$	$27.178_{0.094}$	$0.945_{0.029}$
350	$607.29_{0.23}$	$4.67_{0.12}$	$0.1514_{0.0039}$	$27.83_{0.14}$	$0.959_{0.043}$

Table SI.X: GCMC-MBAR results for 2,2-dimethylpropane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
410	$413.2_{2.6}$	$75.9_{2.6}$	$2.173_{0.029}$	$11.65_{0.11}$	$0.606_{0.013}$
400	$437.1_{2.3}$	$59.3_{1.8}$	$1.831_{0.018}$	$13.410_{0.083}$	$0.669_{0.014}$
390	$457.4_{1.5}$	$47.4_{1.1}$	$1.532_{0.010}$	$14.814_{0.051}$	$0.719_{0.013}$
380	$474.6_{1.1}$	$38.29_{0.72}$	$1.2725_{0.0056}$	$15.944_{0.045}$	$0.759_{0.011}$
370	$489.78_{0.98}$	$31.00_{0.48}$	$1.0468_{0.0035}$	$16.909_{0.045}$	$0.792_{0.011}$
360	$504.12_{0.89}$	$25.02_{0.33}$	$0.8520_{0.0036}$	$17.785_{0.044}$	$0.821_{0.012}$
350	$518.05_{0.86}$	$20.05_{0.21}$	$0.6853_{0.0043}$	$18.604_{0.046}$	$0.847_{0.011}$
340	$531.32_{0.93}$	$15.93_{0.18}$	$0.5437_{0.0045}$	$19.363_{0.057}$	$0.871_{0.011}$
330	$543.65_{0.96}$	$12.52_{0.23}$	$0.4252_{0.0041}$	$20.056_{0.078}$	$0.893_{0.016}$
320	$555.0_{1.0}$	$9.71_{0.28}$	$0.3271_{0.0037}$	$20.68_{0.11}$	$0.913_{0.029}$
310	$565.5_{1.4}$	$7.43_{0.32}$	$0.2473_{0.0044}$	$21.25_{0.18}$	$0.931_{0.052}$
300	$576.4_{1.8}$	$5.60_{0.33}$	$0.1832_{0.0061}$	$21.82_{0.28}$	$0.947_{0.087}$
290	$588.4_{1.3}$	$4.13_{0.32}$	$0.1327_{0.0084}$	$22.41_{0.38}$	$0.96_{0.14}$
280	$600.2_{1.1}$	$2.98_{0.29}$	$0.093_{0.011}$	$22.98_{0.55}$	$0.97_{0.22}$

Table SI.XI: GCMC-MBAR results for 2,2-dimethylbutane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	410.4 _{5.2}	86.7 _{8.5}	$2.34_{0.11}$	$12.61_{0.20}$	$0.595_{0.027}$
460	$436.5_{4.9}$	$70.3_{8.7}$	$2.014_{0.071}$	$14.48_{0.34}$	$0.645_{0.049}$
450	$457.9_{3.1}$	$57.7_{7.2}$	$1.724_{0.036}$	$16.04_{0.45}$	$0.689_{0.061}$
440	$476.4_{1.5}$	$47.7_{4.6}$	$1.467_{0.011}$	$17.37_{0.41}$	$0.725_{0.057}$
430	$493.01_{0.95}$	$39.4_{2.1}$	$1.2386_{0.0045}$	$18.53_{0.25}$	$0.757_{0.039}$
420	$508.03_{0.71}$	$32.56_{0.58}$	$1.0375_{0.0094}$	$19.573_{0.098}$	$0.786_{0.020}$
410	$521.74_{0.38}$	$26.79_{0.14}$	$0.862_{0.010}$	$20.513_{0.035}$	$0.81_{0.01}$
400	$534.75_{0.29}$	$21.94_{0.16}$	$0.7088_{0.0094}$	$21.383_{0.026}$	$0.84_{0.01}$
390	$547.49_{0.34}$	$17.86_{0.11}$	$0.5771_{0.0086}$	$22.210_{0.027}$	$0.859_{0.010}$
380	$559.93_{0.37}$	$14.416_{0.067}$	$0.4643_{0.0081}$	$22.998_{0.038}$	$0.878_{0.013}$
370	$571.83_{0.42}$	$11.528_{0.058}$	$0.3687_{0.0077}$	$23.738_{0.053}$	$0.896_{0.017}$
360	$582.92_{0.45}$	$9.115_{0.074}$	$0.2885_{0.0074}$	$24.416_{0.062}$	$0.911_{0.022}$
350	$593.10_{0.52}$	$7.115_{0.099}$	$0.2224_{0.0070}$	$25.030_{0.081}$	$0.925_{0.029}$
340	$602.70_{0.55}$	$5.47_{0.12}$	$0.1685_{0.0067}$	$25.60_{0.11}$	$0.938_{0.041}$
330	$612.20_{0.41}$	$4.14_{0.14}$	$0.1253_{0.0065}$	$26.15_{0.16}$	$0.949_{0.061}$
320	$621.65_{0.33}$	$3.08_{0.13}$	$0.0912_{0.0065}$	$26.69_{0.23}$	$0.959_{0.091}$
310	$631.20_{0.30}$	$2.24_{0.12}$	$0.0649_{0.0068}$	$27.22_{0.34}$	$0.97_{0.14}$

Table SI.XII: GCMC-MBAR results for 2,3-dimethylbutane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
480	$411.7_{1.1}$	$88.4_{2.2}$	$2.369_{0.027}$	$12.87_{0.14}$	$0.58_{0.01}$
470	$437.1_{1.0}$	$71.6_{2.1}$	$2.037_{0.019}$	$14.80_{0.17}$	$0.627_{0.013}$
460	$459.23_{0.86}$	$58.1_{1.6}$	$1.742_{0.013}$	$16.53_{0.18}$	$0.676_{0.015}$
450	$478.07_{0.79}$	$47.5_{1.1}$	$1.4813_{0.0074}$	$18.00_{0.16}$	$0.718_{0.014}$
440	$494.54_{0.81}$	$39.07_{0.72}$	$1.2513_{0.0041}$	$19.26_{0.13}$	$0.754_{0.012}$
430	$509.51_{0.77}$	$32.20_{0.43}$	$1.0493_{0.0024}$	$20.37_{0.10}$	$0.79_{0.01}$
420	$523.51_{0.64}$	$26.49_{0.25}$	$0.8726_{0.0019}$	$21.382_{0.079}$	$0.81_{0.01}$
410	$536.78_{0.54}$	$21.70_{0.15}$	$0.7189_{0.0020}$	$22.310_{0.063}$	$0.84_{0.01}$
400	$549.39_{0.59}$	$17.68_{0.10}$	$0.5863_{0.0021}$	$23.171_{0.060}$	$0.86_{0.01}$
390	$561.38_{0.73}$	$14.296_{0.082}$	$0.4727_{0.0023}$	$23.971_{0.062}$	$0.88_{0.01}$
380	$572.70_{0.68}$	$11.455_{0.085}$	$0.3764_{0.0025}$	$24.712_{0.056}$	$0.896_{0.011}$
370	$583.42_{0.40}$	$9.083_{0.092}$	$0.2957_{0.0028}$	$25.402_{0.050}$	$0.912_{0.016}$
360	$593.75_{0.21}$	$7.115_{0.095}$	$0.2289_{0.0033}$	$26.054_{0.064}$	$0.926_{0.024}$
350	$603.97_{0.20}$	$5.497_{0.091}$	$0.1742_{0.0037}$	$26.686_{0.091}$	$0.938_{0.034}$
340	$614.11_{0.22}$	$4.179_{0.082}$	$0.1302_{0.0042}$	$27.30_{0.13}$	$0.950_{0.047}$
330	$623.72_{0.26}$	$3.121_{0.069}$	$0.0953_{0.0047}$	$27.87_{0.18}$	$0.959_{0.066}$
320	$632.83_{0.34}$	$2.284_{0.057}$	$0.0682_{0.0050}$	$28.41_{0.25}$	$0.968_{0.092}$

Table SI.XIII: GCMC-MBAR results for 2,2,4-trimethylpentane with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
530	$401.6_{2.0}$	$97.3_{2.8}$	$2.113_{0.013}$	$13.65_{0.15}$	$0.563_{0.014}$
520	$428.08_{0.71}$	$79.0_{1.7}$	$1.8383_{0.0085}$	$15.95_{0.17}$	$0.615_{0.012}$
510	$449.27_{0.69}$	$64.82_{0.80}$	$1.5924_{0.0067}$	$17.85_{0.12}$	$0.66_{0.01}$
500	$467.26_{0.83}$	$53.68_{0.34}$	$1.3733_{0.0059}$	$19.449_{0.068}$	$0.70_{0.01}$
490	$484.19_{0.91}$	$44.67_{0.32}$	$1.1784_{0.0050}$	$20.898_{0.058}$	$0.74_{0.01}$
480	$500.20_{0.83}$	$37.25_{0.39}$	$1.0051_{0.0038}$	$22.224_{0.066}$	$0.77_{0.01}$
470	$514.73_{0.48}$	$31.08_{0.39}$	$0.8515_{0.0027}$	$23.414_{0.060}$	$0.80_{0.01}$
460	$527.91_{0.42}$	$25.89_{0.30}$	$0.7160_{0.0022}$	$24.486_{0.050}$	$0.83_{0.01}$
450	$540.02_{0.69}$	$21.49_{0.18}$	$0.5973_{0.0026}$	$25.458_{0.051}$	$0.85_{0.01}$
440	$551.36_{0.72}$	$17.746_{0.095}$	$0.4939_{0.0030}$	$26.355_{0.050}$	$0.87_{0.01}$
430	$562.92_{0.74}$	$14.553_{0.071}$	$0.4046_{0.0031}$	$27.239_{0.050}$	$0.89_{0.01}$
420	$574.83_{0.92}$	$11.836_{0.082}$	$0.3278_{0.0033}$	$28.116_{0.049}$	$0.906_{0.013}$
410	$585.23_{0.62}$	$9.54_{0.11}$	$0.2626_{0.0037}$	$28.885_{0.042}$	$0.922_{0.022}$
400	$594.42_{0.22}$	$7.62_{0.14}$	$0.2078_{0.0042}$	$29.567_{0.098}$	$0.937_{0.036}$
390	$604.54_{0.21}$	$6.02_{0.18}$	$0.1623_{0.0051}$	$30.29_{0.17}$	$0.949_{0.058}$
380	$615.63_{0.16}$	$4.70_{0.20}$	$0.1247_{0.0061}$	$31.04_{0.27}$	$0.959_{0.089}$
370	$625.43_{0.14}$	$3.62_{0.20}$	$0.0942_{0.0072}$	$31.71_{0.40}$	$0.97_{0.13}$

SI.III.2.2 Alkynes

Table SI.XIV: GCMC-MBAR results for ethyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
290	$421.8_{1.7}$	$69.4_{1.8}$	$4.071_{0.027}$	$8.835_{0.094}$	$0.634_{0.013}$
280	$451.1_{1.2}$	$50.48_{0.65}$	$3.187_{0.018}$	$10.290_{0.067}$	$0.71_{0.01}$
270	$475.2_{2.5}$	$37.51_{0.33}$	$2.457_{0.013}$	$11.409_{0.072}$	$0.76_{0.01}$
260	$497.6_{1.7}$	$27.84_{0.34}$	$1.8584_{0.0078}$	$12.379_{0.043}$	$0.80_{0.01}$
250	$518.16_{0.83}$	$20.45_{0.22}$	$1.3737_{0.0065}$	$13.231_{0.027}$	$0.84_{0.01}$
240	$537.09_{0.92}$	$14.759_{0.088}$	$0.9884_{0.0067}$	$13.991_{0.021}$	$0.87_{0.01}$
230	$554.32_{0.93}$	$10.40_{0.10}$	$0.6899_{0.0056}$	$14.664_{0.024}$	$0.90_{0.01}$
220	$571.08_{0.74}$	$7.12_{0.12}$	$0.4649_{0.0046}$	$15.298_{0.035}$	$0.929_{0.019}$

Table SI.XV: GCMC-MBAR results for propyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
380	$436.3_{5.1}$	$85.3_{1.7}$	$3.923_{0.045}$	$10.64_{0.20}$	$0.58_{0.01}$
370	$468.9_{3.2}$	$64.9_{1.9}$	$3.234_{0.036}$	$12.56_{0.18}$	$0.649_{0.012}$
360	$495.3_{1.2}$	$50.2_{1.4}$	$2.643_{0.025}$	$14.12_{0.13}$	$0.705_{0.014}$
350	$517.6_{1.1}$	$39.26_{0.77}$	$2.137_{0.018}$	$15.39_{0.10}$	$0.750_{0.011}$
340	$537.2_{1.3}$	$30.77_{0.40}$	$1.708_{0.013}$	$16.469_{0.084}$	$0.79_{0.01}$
330	$554.7_{1.7}$	$24.00_{0.28}$	$1.3456_{0.0097}$	$17.416_{0.089}$	$0.82_{0.01}$
320	$570.4_{2.2}$	$18.56_{0.23}$	$1.0444_{0.0070}$	$18.26_{0.10}$	$0.85_{0.01}$
310	$585.8_{2.1}$	$14.18_{0.18}$	$0.7969_{0.0050}$	$19.048_{0.098}$	$0.87_{0.01}$
300	$601.7_{1.6}$	$10.67_{0.12}$	$0.5964_{0.0038}$	$19.823_{0.073}$	$0.90_{0.01}$
290	$616.5_{2.2}$	$7.899_{0.078}$	$0.4365_{0.0038}$	$20.532_{0.077}$	$0.918_{0.012}$

Table SI.XVI: GCMC-MBAR results for 1-butyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
410	$452.8_{4.0}$	$70.9_{5.6}$	$2.840_{0.093}$	$13.41_{0.28}$	$0.636_{0.032}$
400	$478.29_{0.92}$	$55.9_{2.7}$	$2.366_{0.066}$	$15.10_{0.20}$	$0.689_{0.016}$
390	$500.1_{1.3}$	$44.5_{1.3}$	$1.955_{0.052}$	$16.52_{0.12}$	$0.73_{0.01}$
380	$520.1_{1.4}$	$35.6_{1.1}$	$1.599_{0.043}$	$17.771_{0.095}$	$0.77_{0.01}$
370	$538.8_{1.1}$	$28.3_{1.2}$	$1.293_{0.033}$	$18.90_{0.10}$	$0.802_{0.015}$
360	$555.7_{1.3}$	$22.5_{1.2}$	$1.032_{0.021}$	$19.92_{0.14}$	$0.831_{0.031}$
350	$571.1_{1.3}$	$17.64_{0.98}$	$0.8126_{0.0098}$	$20.83_{0.15}$	$0.856_{0.041}$
340	$585.1_{1.0}$	$13.72_{0.65}$	$0.6302_{0.0033}$	$21.64_{0.16}$	$0.879_{0.044}$
330	$598.8_{2.7}$	$10.54_{0.36}$	$0.4806_{0.0057}$	$22.41_{0.19}$	$0.899_{0.042}$
320	$613.6_{4.5}$	$7.97_{0.19}$	$0.3595_{0.0080}$	$23.21_{0.25}$	$0.917_{0.039}$
310	$627.0_{2.2}$	$5.92_{0.13}$	$0.2629_{0.0086}$	$23.93_{0.17}$	$0.932_{0.040}$

Table SI.XVII: GCMC-MBAR results for 2-butyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
450	$437.2_{4.2}$	$90.5_{3.2}$	$3.554_{0.053}$	$12.28_{0.20}$	$0.568_{0.014}$
440	$470.0_{3.7}$	$71.8_{2.3}$	$3.012_{0.037}$	$14.30_{0.20}$	$0.620_{0.017}$
430	$494.6_{1.6}$	$57.4_{1.2}$	$2.535_{0.029}$	$15.96_{0.15}$	$0.668_{0.014}$
420	$514.58_{0.50}$	$46.27_{0.57}$	$2.118_{0.025}$	$17.336_{0.085}$	$0.71_{0.01}$
410	$532.82_{0.48}$	$37.39_{0.49}$	$1.757_{0.022}$	$18.554_{0.040}$	$0.75_{0.01}$
400	$550.44_{0.50}$	$30.16_{0.62}$	$1.444_{0.017}$	$19.684_{0.065}$	$0.78_{0.01}$
390	$567.00_{0.68}$	$24.23_{0.63}$	$1.175_{0.012}$	$20.72_{0.11}$	$0.809_{0.013}$
380	$582.34_{0.91}$	$19.36_{0.53}$	$0.9452_{0.0067}$	$21.66_{0.13}$	$0.836_{0.017}$
370	$596.57_{0.71}$	$15.35_{0.39}$	$0.7513_{0.0033}$	$22.51_{0.12}$	$0.860_{0.019}$
360	$610.15_{0.68}$	$12.07_{0.25}$	$0.5893_{0.0027}$	$23.32_{0.12}$	$0.882_{0.019}$
350	$623.0_{1.1}$	$9.39_{0.13}$	$0.4552_{0.0037}$	$24.06_{0.12}$	$0.901_{0.018}$
340	$635.13_{0.92}$	$7.213_{0.060}$	$0.3458_{0.0041}$	$24.75_{0.10}$	$0.917_{0.016}$
330	$647.43_{0.75}$	$5.454_{0.041}$	$0.2577_{0.0039}$	$25.410_{0.086}$	$0.931_{0.015}$

Table SI.XVIII: GCMC-MBAR results for 1-pentyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(K)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
450	$445.1_{1.3}$	$77.7_{1.0}$	$2.582_{0.012}$	$14.14_{0.12}$	$0.61_{0.01}$
440	$471.10_{0.57}$	$61.18_{0.83}$	$2.1759_{0.0091}$	$16.143_{0.079}$	$0.66_{0.01}$
430	$493.03_{0.73}$	$48.93_{0.57}$	$1.8221_{0.0082}$	$17.791_{0.054}$	$0.71_{0.01}$
420	$512.1_{1.0}$	$39.44_{0.30}$	$1.5143_{0.0083}$	$19.182_{0.025}$	$0.75_{0.01}$
410	$529.2_{1.4}$	$31.85_{0.22}$	$1.2476_{0.0079}$	$20.400_{0.045}$	$0.78_{0.01}$
400	$545.2_{1.3}$	$25.65_{0.31}$	$1.0180_{0.0064}$	$21.503_{0.073}$	$0.81_{0.01}$
390	$560.39_{0.90}$	$20.55_{0.33}$	$0.8216_{0.0042}$	$22.528_{0.075}$	$0.84_{0.01}$
380	$574.87_{0.92}$	$16.34_{0.27}$	$0.6551_{0.0022}$	$23.483_{0.088}$	$0.865_{0.012}$
370	$588.2_{1.7}$	$12.87_{0.20}$	$0.5155_{0.0014}$	$24.35_{0.13}$	$0.887_{0.014}$
360	$600.8_{2.1}$	$10.03_{0.16}$	$0.4000_{0.0020}$	$25.16_{0.15}$	$0.907_{0.018}$
350	$613.6_{1.5}$	$7.72_{0.15}$	$0.3052_{0.0029}$	$25.96_{0.14}$	$0.925_{0.026}$
340	$626.2_{1.2}$	$5.86_{0.16}$	$0.2288_{0.0041}$	$26.72_{0.17}$	$0.941_{0.042}$

Table SI.XIX: GCMC-MBAR results for 2-pentyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(K)$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	$470.2_{2.0}$	$67.8_{1.5}$	$2.443_{0.022}$	$16.285_{0.075}$	$0.628_{0.016}$
460	$493.37_{0.97}$	$54.31_{0.93}$	$2.066_{0.024}$	$18.127_{0.084}$	$0.677_{0.016}$
450	$513.26_{0.50}$	$43.98_{0.45}$	$1.735_{0.025}$	$19.667_{0.065}$	$0.718_{0.013}$
440	$531.38_{0.38}$	$35.74_{0.41}$	$1.447_{0.023}$	$21.022_{0.031}$	$0.75_{0.01}$
430	$548.11_{0.47}$	$29.01_{0.58}$	$1.197_{0.020}$	$22.242_{0.067}$	$0.79_{0.01}$
420	$563.44_{0.72}$	$23.48_{0.64}$	$0.981_{0.015}$	$23.34_{0.12}$	$0.81_{0.01}$
410	$577.72_{0.97}$	$18.92_{0.59}$	$0.796_{0.011}$	$24.35_{0.16}$	$0.840_{0.015}$
400	$591.5_{1.4}$	$15.15_{0.47}$	$0.6386_{0.0070}$	$25.30_{0.20}$	$0.863_{0.018}$
390	$604.8_{1.9}$	$12.03_{0.32}$	$0.5062_{0.0042}$	$26.19_{0.22}$	$0.884_{0.018}$
380	$617.3_{1.8}$	$9.46_{0.18}$	$0.3957_{0.0027}$	$27.01_{0.19}$	$0.902_{0.014}$
370	$629.1_{1.5}$	$7.353_{0.077}$	$0.3048_{0.0022}$	$27.78_{0.14}$	$0.92_{0.01}$
360	$640.7_{1.8}$	$5.637_{0.054}$	$0.2308_{0.0024}$	$28.52_{0.12}$	$0.932_{0.016}$
350	$652.0_{1.7}$	$4.255_{0.079}$	$0.1717_{0.0030}$	$29.229_{0.082}$	$0.944_{0.032}$

Table SI.XX: GCMC-MBAR results for 1-hexyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
490	$433.3_{7.6}$	$81.5_{3.5}$	$2.435_{0.037}$	$15.09_{0.17}$	$0.602_{0.017}$
480	$461.1_{2.9}$	$65.5_{2.5}$	$2.080_{0.022}$	$17.26_{0.14}$	$0.654_{0.019}$
470	$483.65_{0.76}$	$53.2_{1.3}$	$1.766_{0.014}$	$19.07_{0.11}$	$0.698_{0.013}$
460	$502.56_{0.55}$	$43.45_{0.54}$	$1.489_{0.010}$	$20.586_{0.041}$	$0.74_{0.01}$
450	$519.59_{0.69}$	$35.56_{0.36}$	$1.2468_{0.0083}$	$21.936_{0.038}$	$0.77_{0.01}$
440	$536.0_{1.0}$	$29.06_{0.41}$	$1.0351_{0.0065}$	$23.201_{0.073}$	$0.80_{0.01}$
430	$551.44_{0.96}$	$23.66_{0.47}$	$0.8514_{0.0043}$	$24.36_{0.10}$	$0.827_{0.014}$
420	$565.34_{0.57}$	$19.17_{0.42}$	$0.6934_{0.0031}$	$25.40_{0.11}$	$0.851_{0.019}$
410	$578.56_{0.49}$	$15.43_{0.30}$	$0.5583_{0.0037}$	$26.378_{0.089}$	$0.872_{0.021}$
400	$591.16_{0.88}$	$12.31_{0.18}$	$0.4443_{0.0049}$	$27.296_{0.062}$	$0.892_{0.022}$
390	$602.46_{0.56}$	$9.72_{0.10}$	$0.3488_{0.0056}$	$28.119_{0.069}$	$0.909_{0.022}$
380	$612.80_{0.61}$	$7.587_{0.080}$	$0.2702_{0.0059}$	$28.86_{0.10}$	$0.926_{0.027}$
370	$624.2_{1.5}$	$5.851_{0.083}$	$0.2061_{0.0060}$	$29.65_{0.15}$	$0.940_{0.037}$
360	$637.2_{1.1}$	$4.445_{0.086}$	$0.1543_{0.0064}$	$30.50_{0.19}$	$0.953_{0.053}$

Table SI.XXI: GCMC-MBAR results for 2-hexyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
500	$467.54_{0.99}$	$66.01_{0.94}$	$2.140_{0.019}$	$17.945_{0.093}$	$0.64_{0.01}$
490	$488.44_{0.86}$	$53.67_{0.62}$	$1.825_{0.016}$	$19.766_{0.070}$	$0.69_{0.01}$
480	$506.42_{0.87}$	$44.07_{0.54}$	$1.547_{0.012}$	$21.286_{0.048}$	$0.72_{0.01}$
470	$523.2_{1.3}$	$36.28_{0.58}$	$1.3025_{0.0090}$	$22.660_{0.039}$	$0.75_{0.01}$
460	$539.7_{1.9}$	$29.82_{0.56}$	$1.0881_{0.0054}$	$23.957_{0.044}$	$0.784_{0.012}$
450	$555.2_{1.7}$	$24.42_{0.46}$	$0.9014_{0.0029}$	$25.157_{0.042}$	$0.810_{0.015}$
440	$569.3_{1.2}$	$19.91_{0.33}$	$0.7397_{0.0032}$	$26.240_{0.053}$	$0.834_{0.016}$
430	$582.7_{1.3}$	$16.13_{0.20}$	$0.6010_{0.0043}$	$27.253_{0.058}$	$0.856_{0.016}$
420	$596.1_{1.2}$	$12.96_{0.11}$	$0.4831_{0.0051}$	$28.236_{0.062}$	$0.876_{0.015}$
410	$608.84_{0.86}$	$10.324_{0.065}$	$0.3835_{0.0054}$	$29.161_{0.058}$	$0.895_{0.014}$
400	$620.4_{1.4}$	$8.136_{0.057}$	$0.3005_{0.0053}$	$30.003_{0.073}$	$0.912_{0.015}$
390	$631.7_{1.6}$	$6.340_{0.060}$	$0.2321_{0.0051}$	$30.807_{0.083}$	$0.928_{0.018}$
380	$642.43_{0.53}$	$4.878_{0.064}$	$0.1766_{0.0049}$	$31.568_{0.066}$	$0.941_{0.026}$
370	$652.13_{0.63}$	$3.702_{0.067}$	$0.1322_{0.0047}$	$32.24_{0.12}$	$0.953_{0.040}$

Table SI.XXII: GCMC-MBAR results for 1-heptyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
520	$437.2_{2.6}$	$77.0_{2.2}$	$2.122_{0.017}$	$16.99_{0.12}$	$0.613_{0.013}$
510	$462.1_{1.5}$	$62.4_{1.4}$	$1.822_{0.011}$	$19.188_{0.099}$	$0.662_{0.013}$
500	$483.1_{1.1}$	$51.09_{0.74}$	$1.5560_{0.0076}$	$21.050_{0.060}$	$0.70_{0.01}$
490	$501.4_{1.0}$	$42.07_{0.40}$	$1.3207_{0.0063}$	$22.662_{0.042}$	$0.74_{0.01}$
480	$518.0_{1.1}$	$34.69_{0.32}$	$1.1135_{0.0053}$	$24.100_{0.057}$	$0.77_{0.01}$
470	$533.2_{1.2}$	$28.57_{0.30}$	$0.9318_{0.0044}$	$25.399_{0.069}$	$0.80_{0.01}$
460	$547.3_{1.1}$	$23.46_{0.24}$	$0.7736_{0.0042}$	$26.588_{0.061}$	$0.83_{0.01}$
450	$560.81_{1.0}$	$19.18_{0.16}$	$0.6364_{0.0046}$	$27.703_{0.047}$	$0.853_{0.011}$
440	$573.98_{0.87}$	$15.59_{0.12}$	$0.5185_{0.0052}$	$28.768_{0.035}$	$0.874_{0.015}$
430	$586.56_{0.51}$	$12.58_{0.10}$	$0.4179_{0.0057}$	$29.770_{0.052}$	$0.894_{0.019}$
420	$598.39_{0.36}$	$10.06_{0.10}$	$0.3329_{0.0061}$	$30.703_{0.082}$	$0.911_{0.024}$
410	$609.60_{0.41}$	$7.97_{0.12}$	$0.2618_{0.0064}$	$31.576_{0.098}$	$0.927_{0.030}$
400	$620.43_{0.28}$	$6.24_{0.16}$	$0.2031_{0.0065}$	$32.41_{0.13}$	$0.941_{0.042}$
390	$631.53_{0.29}$	$4.82_{0.19}$	$0.1551_{0.0067}$	$33.24_{0.20}$	$0.953_{0.062}$

Table SI.XXIII: GCMC-MBAR results for 1-octyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
550	$424.4_{3.1}$	$85.7_{2.4}$	$2.025_{0.020}$	$16.856_{0.096}$	$0.570_{0.012}$
540	$449.6_{2.4}$	$68.1_{1.6}$	$1.745_{0.012}$	$19.509_{0.082}$	$0.629_{0.011}$
530	$471.3_{1.4}$	$55.11_{0.94}$	$1.4972_{0.0080}$	$21.746_{0.072}$	$0.68_{0.01}$
520	$490.29_{0.85}$	$45.27_{0.56}$	$1.2786_{0.0057}$	$23.612_{0.065}$	$0.72_{0.01}$
510	$507.10_{0.64}$	$37.43_{0.38}$	$1.0854_{0.0043}$	$25.219_{0.062}$	$0.75_{0.01}$
500	$522.35_{0.60}$	$30.99_{0.30}$	$0.9152_{0.0033}$	$26.653_{0.065}$	$0.78_{0.01}$
490	$536.64_{0.61}$	$25.61_{0.25}$	$0.7660_{0.0026}$	$27.971_{0.072}$	$0.81_{0.01}$
480	$550.42_{0.57}$	$21.08_{0.21}$	$0.6360_{0.0023}$	$29.216_{0.075}$	$0.83_{0.01}$
470	$563.64_{0.48}$	$17.26_{0.16}$	$0.5234_{0.0024}$	$30.392_{0.075}$	$0.85_{0.01}$
460	$576.05_{0.40}$	$14.05_{0.13}$	$0.4265_{0.0026}$	$31.485_{0.074}$	$0.875_{0.012}$
450	$587.80_{0.34}$	$11.35_{0.11}$	$0.3440_{0.0029}$	$32.510_{0.077}$	$0.893_{0.014}$
440	$599.21_{0.42}$	$9.09_{0.11}$	$0.2743_{0.0031}$	$33.488_{0.090}$	$0.909_{0.018}$
430	$610.23_{0.53}$	$7.210_{0.099}$	$0.2160_{0.0034}$	$34.42_{0.11}$	$0.923_{0.023}$
420	$620.82_{0.50}$	$5.656_{0.089}$	$0.1678_{0.0037}$	$35.31_{0.12}$	$0.936_{0.031}$
410	$631.54_{0.36}$	$4.383_{0.076}$	$0.1285_{0.0039}$	$36.19_{0.15}$	$0.948_{0.041}$

Table SI.XXIV: GCMC-MBAR results for 1-nonyne with the iMiPPE force field (optimal ψ value from Table SI.V). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
570	$427.1_{1.4}$	$80.51_{0.73}$	$1.776_{0.010}$	$18.58_{0.12}$	$0.58_{0.01}$
560	$450.90_{0.83}$	$64.43_{0.54}$	$1.5311_{0.0083}$	$21.279_{0.086}$	$0.63_{0.01}$
550	$471.72_{0.65}$	$52.31_{0.42}$	$1.3149_{0.0067}$	$23.583_{0.093}$	$0.68_{0.01}$
540	$489.89_{0.61}$	$43.02_{0.39}$	$1.1239_{0.0056}$	$25.52_{0.11}$	$0.72_{0.01}$
530	$506.26_{0.52}$	$35.61_{0.32}$	$0.9552_{0.0047}$	$27.21_{0.11}$	$0.76_{0.01}$
520	$521.47_{0.54}$	$29.51_{0.25}$	$0.8067_{0.0040}$	$28.739_{0.079}$	$0.79_{0.01}$
510	$535.75_{0.66}$	$24.42_{0.23}$	$0.6762_{0.0031}$	$30.149_{0.058}$	$0.81_{0.01}$
500	$549.10_{0.71}$	$20.14_{0.24}$	$0.5626_{0.0022}$	$31.452_{0.059}$	$0.83_{0.01}$
490	$561.65_{0.67}$	$16.54_{0.23}$	$0.4641_{0.0013}$	$32.666_{0.073}$	$0.856_{0.010}$
480	$573.68_{0.55}$	$13.50_{0.21}$	$0.37923_{0.00095}$	$33.811_{0.085}$	$0.875_{0.014}$
470	$585.26_{0.38}$	$10.94_{0.18}$	$0.3068_{0.0015}$	$34.896_{0.092}$	$0.892_{0.018}$
460	$596.35_{0.30}$	$8.79_{0.16}$	$0.2455_{0.0022}$	$35.92_{0.10}$	$0.907_{0.023}$
450	$607.27_{0.32}$	$7.00_{0.14}$	$0.1942_{0.0030}$	$36.92_{0.13}$	$0.921_{0.031}$
440	$618.21_{0.22}$	$5.52_{0.12}$	$0.1516_{0.0036}$	$37.91_{0.16}$	$0.933_{0.041}$
430	$628.74_{0.19}$	$4.29_{0.11}$	$0.1166_{0.0042}$	$38.85_{0.22}$	$0.944_{0.055}$
420	$638.50_{0.35}$	$3.296_{0.094}$	$0.0885_{0.0047}$	$39.72_{0.29}$	$0.955_{0.074}$

SI.IV Case study: Cyclohexane optimization

SI.IV.1 Minimum number of effective snapshots

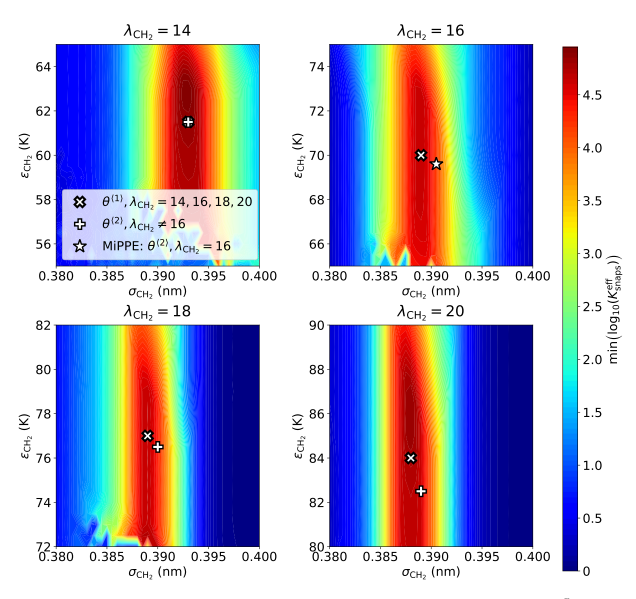


Figure SI.1: Second iteration minimum number of effective snapshots $(\min(K_{\mathrm{snaps}}^{\mathrm{eff}}))$ with respect to $\epsilon_{\mathrm{CH_2}}$ and $\sigma_{\mathrm{CH_2}}$ for cyclohexane. Optimization has converged as $\min(K_{\mathrm{snaps}}^{\mathrm{eff}}) \gg 50$ for the optimal $\epsilon_{\mathrm{CH_2}}$, $\sigma_{\mathrm{CH_2}}$, $\lambda_{\mathrm{CH_2}}$ parameter set. Top-left, top-right, bottom-left, and bottom-right panels correspond $\lambda_{\mathrm{CH_2}} = 14$, $\lambda_{\mathrm{CH_2}} = 16$, $\lambda_{\mathrm{CH_2}} = 18$, and $\lambda_{\mathrm{CH_2}} = 12$, respectively. White star represents the optimal parameter set, i.e., the lowest value of S, for a given $\lambda_{\mathrm{CH_2}}$

SI.IV.2 Tabulated phase equilibria for iterations

Table SI.XXV: GCMC-MBAR results for the MiPPE force field (second iteration, $\theta^{\langle 2 \rangle}$ $\lambda_{\rm CH_2}=16$). Subscripts correspond to the 95% confidence interval computed with twenty independent replicate GCMC simulations at each state point.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	$710.9_{1.7}$	$3.552_{0.027}$	$0.1213_{0.0012}$	$29.648_{0.084}$	$0.960_{0.012}$
370	$700.9_{2.0}$	$4.611_{0.032}$	$0.1604_{0.0015}$	$29.106_{0.098}$	$0.952_{0.011}$
380	$690.8_{1.8}$	$5.898_{0.039}$	$0.2085_{0.0018}$	$28.548_{0.088}$	$0.942_{0.010}$
390	$680.5_{1.5}$	$7.446_{0.048}$	$0.2670_{0.0022}$	$27.967_{0.071}$	$0.9306_{0.0097}$
400	$669.9_{1.4}$	$9.291_{0.056}$	$0.3372_{0.0026}$	$27.359_{0.067}$	$0.9183_{0.0090}$
410	$659.0_{1.4}$	$11.474_{0.065}$	$0.4205_{0.0030}$	$26.720_{0.069}$	$0.9048_{0.0083}$
420	$647.8_{1.6}$	$14.040_{0.075}$	$0.5185_{0.0034}$	$26.050_{0.074}$	$0.8899_{0.0075}$
430	$636.3_{1.8}$	$17.043_{0.085}$	$0.6325_{0.0039}$	$25.346_{0.082}$	$0.8736_{0.0069}$
440	$624.3_{1.8}$	$20.543_{0.096}$	$0.7642_{0.0044}$	$24.600_{0.080}$	$0.8558_{0.0064}$
450	$611.8_{1.6}$	$24.61_{0.11}$	$0.9152_{0.0051}$	$23.807_{0.069}$	$0.8363_{0.0059}$
460	$598.7_{1.3}$	$29.35_{0.11}$	$1.0871_{0.0058}$	$22.958_{0.058}$	$0.8151_{0.0054}$
470	$584.9_{1.1}$	$34.85_{0.12}$	$1.2816_{0.0066}$	$22.044_{0.055}$	$0.7920_{0.0049}$
480	$570.25_{0.93}$	$41.29_{0.17}$	$1.5006_{0.0072}$	$21.051_{0.058}$	$0.7665_{0.0048}$
490	$554.49_{0.93}$	$48.88_{0.31}$	$1.7460_{0.0075}$	$19.956_{0.063}$	$0.7379_{0.0057}$
500	$537.3_{1.2}$	$58.02_{0.54}$	$2.0200_{0.0076}$	$18.718_{0.064}$	$0.7049_{0.0071}$
510	$518.0_{2.0}$	$69.36_{0.85}$	$2.3254_{0.0084}$	$17.276_{0.076}$	$0.6654_{0.0085}$
520	$496.3_{3.0}$	$83.9_{1.4}$	$2.666_{0.011}$	$15.58_{0.13}$	$0.619_{0.011}$

Table SI.XXVI: GCMC-MBAR results for the TraPPE force field (zeroth iteration, $\theta^{\langle 0 \rangle}$). Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	$709.02_{0.33}$	$4.775_{0.062}$	$0.161_{0.016}$	$26.916_{0.050}$	$0.946_{0.096}$
370	$698.93_{0.39}$	$6.067_{0.072}$	$0.208_{0.012}$	$26.440_{0.042}$	$0.937_{0.055}$
380	$688.03_{0.44}$	$7.610_{0.088}$	$0.26_{0.01}$	$25.919_{0.036}$	$0.927_{0.026}$
390	$677.14_{0.49}$	$9.43_{0.11}$	$0.33_{0.01}$	$25.382_{0.033}$	$0.916_{0.015}$
400	$666.43_{0.55}$	$11.57_{0.12}$	$0.41_{0.01}$	$24.835_{0.030}$	$0.904_{0.023}$
410	$655.56_{0.39}$	$14.07_{0.13}$	$0.507_{0.017}$	$24.266_{0.023}$	$0.890_{0.032}$
420	$644.33_{0.48}$	$16.97_{0.13}$	$0.616_{0.025}$	$23.665_{0.027}$	$0.875_{0.036}$
430	$632.76_{0.54}$	$20.32_{0.12}$	$0.741_{0.031}$	$23.030_{0.030}$	$0.858_{0.037}$
440	$620.79_{0.67}$	$24.18_{0.26}$	$0.883_{0.039}$	$22.356_{0.051}$	$0.840_{0.038}$
450	$608.2_{1.2}$	$28.64_{0.96}$	$1.045_{0.052}$	$21.63_{0.10}$	$0.821_{0.049}$
460	$595.0_{2.9}$	$33.8_{2.4}$	$1.227_{0.098}$	$20.85_{0.20}$	$0.799_{0.085}$
470	$581.0_{4.6}$	$39.7_{3.8}$	$1.43_{0.17}$	$20.01_{0.30}$	$0.78_{0.12}$
480	$566.2_{4.8}$	$46.7_{4.6}$	$1.66_{0.24}$	$19.08_{0.33}$	$0.75_{0.13}$
490	$550.4_{4.5}$	$54.8_{4.7}$	$1.91_{0.29}$	$18.06_{0.38}$	$0.72_{0.13}$
500	$533.2_{5.4}$	$64.6_{3.8}$	$2.19_{0.38}$	$16.913_{0.088}$	$0.69_{0.13}$
510	$513.8_{5.3}$	$76.6_{5.4}$	$2.50_{0.37}$	$15.59_{0.33}$	$0.65_{0.11}$
520	$491.6_{3.8}$	$91.4_{7.6}$	$2.85_{0.28}$	$14.08_{0.34}$	$0.607_{0.078}$

Table SI.XXVII: GCMC-MBAR results for the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\text{CH}_2}=14$ force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}} (\mathrm{kg/m^3})$	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	$704.93_{0.26}$	$3.743_{0.073}$	$0.127_{0.044}$	$28.78_{0.10}$	$0.96_{0.33}$
370	$695.36_{0.63}$	$4.822_{0.057}$	$0.167_{0.048}$	$28.28_{0.11}$	$0.95_{0.27}$
380	$684.68_{0.79}$	$6.125_{0.057}$	$0.216_{0.049}$	$27.73_{0.11}$	$0.94_{0.21}$
390	$673.71_{0.68}$	$7.680_{0.084}$	$0.274_{0.048}$	$27.144_{0.099}$	$0.93_{0.16}$
400	$663.17_{0.45}$	$9.52_{0.13}$	$0.345_{0.044}$	$26.568_{0.086}$	$0.92_{0.12}$
410	$652.73_{0.24}$	$11.68_{0.17}$	$0.427_{0.039}$	$25.980_{0.062}$	$0.903_{0.084}$
420	$641.98_{0.82}$	$14.21_{0.23}$	$0.524_{0.033}$	$25.360_{0.039}$	$0.889_{0.058}$
430	$630.8_{1.2}$	$17.15_{0.30}$	$0.636_{0.032}$	$24.703_{0.036}$	$0.873_{0.046}$
440	$619.3_{1.1}$	$20.57_{0.39}$	$0.765_{0.041}$	$24.005_{0.044}$	$0.856_{0.049}$
450	$607.24_{0.80}$	$24.53_{0.54}$	$0.912_{0.062}$	$23.261_{0.063}$	$0.837_{0.060}$
460	$594.62_{0.73}$	$29.11_{0.77}$	$1.080_{0.093}$	$22.463_{0.085}$	$0.816_{0.074}$
470	$581.33_{0.87}$	$34.4_{1.1}$	$1.27_{0.14}$	$21.60_{0.11}$	$0.793_{0.091}$
480	$567.3_{1.1}$	$40.6_{1.6}$	$1.48_{0.21}$	$20.68_{0.13}$	$0.77_{0.11}$
490	$552.4_{1.5}$	$47.9_{2.1}$	$1.72_{0.30}$	$19.66_{0.14}$	$0.74_{0.13}$
500	$536.3_{2.5}$	$56.5_{2.6}$	$1.98_{0.41}$	$18.52_{0.14}$	$0.71_{0.15}$
510	$518.2_{4.3}$	$67.2_{2.8}$	$2.27_{0.54}$	$17.20_{0.16}$	$0.67_{0.16}$

Table SI.XXVIII: GCMC-MBAR results for the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\rm CH_2}=16$ force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}} (\mathrm{kg/m^3})$	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	$713.64_{0.38}$	$3.583_{0.049}$	$0.122_{0.039}$	$29.59_{0.13}$	$0.96_{0.31}$
370	$704.1_{1.2}$	$4.647_{0.057}$	$0.162_{0.036}$	$29.065_{0.088}$	$0.95_{0.21}$
380	$695.0_{2.4}$	$5.943_{0.071}$	$0.210_{0.032}$	$28.551_{0.098}$	$0.94_{0.14}$
390	$685.2_{2.3}$	$7.505_{0.090}$	$0.269_{0.027}$	$27.993_{0.084}$	$0.931_{0.094}$
400	$674.3_{1.1}$	$9.37_{0.11}$	$0.340_{0.024}$	$27.376_{0.037}$	$0.918_{0.066}$
410	$662.91_{0.76}$	$11.57_{0.12}$	$0.424_{0.026}$	$26.717_{0.040}$	$0.905_{0.056}$
420	$651.2_{1.0}$	$14.16_{0.14}$	$0.523_{0.032}$	$26.027_{0.065}$	$0.890_{0.055}$
430	$639.4_{1.1}$	$17.19_{0.18}$	$0.638_{0.040}$	$25.310_{0.078}$	$0.873_{0.056}$
440	$627.31_{0.89}$	$20.71_{0.22}$	$0.770_{0.050}$	$24.565_{0.072}$	$0.856_{0.056}$
450	$614.82_{0.58}$	$24.79_{0.24}$	$0.922_{0.059}$	$23.777_{0.051}$	$0.837_{0.055}$
460	$601.71_{0.56}$	$29.53_{0.25}$	$1.095_{0.067}$	$22.935_{0.053}$	$0.816_{0.050}$
470	$587.98_{0.65}$	$35.03_{0.48}$	$1.291_{0.067}$	$22.033_{0.094}$	$0.794_{0.042}$
480	$573.59_{0.74}$	$41.45_{0.99}$	$1.511_{0.063}$	$21.06_{0.15}$	$0.768_{0.037}$
490	$558.19_{0.80}$	$49.0_{1.7}$	$1.757_{0.086}$	$19.99_{0.20}$	$0.740_{0.045}$
500	$541.27_{0.93}$	$58.2_{2.6}$	$2.03_{0.16}$	$18.78_{0.24}$	$0.707_{0.064}$
510	$522.1_{1.7}$	$69.5_{3.4}$	$2.34_{0.28}$	$17.36_{0.25}$	$0.668_{0.087}$
520	$499.8_{2.6}$	83.7 _{3.9}	$2.68_{0.43}$	$15.71_{0.22}$	$0.62_{0.10}$

Table SI.XXIX: GCMC-MBAR results for the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\text{CH}_2} = 18$ force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	$713.60_{0.89}$	$3.294_{0.099}$	$0.113_{0.020}$	$30.527_{0.066}$	$0.96_{0.17}$
370	$702.86_{0.81}$	$4.306_{0.092}$	$0.150_{0.015}$	$29.921_{0.044}$	$0.95_{0.10}$
380	$692.51_{0.62}$	$5.545_{0.081}$	$0.196_{0.013}$	$29.327_{0.039}$	$0.943_{0.063}$
390	$682.23_{0.53}$	$7.043_{0.088}$	$0.253_{0.011}$	$28.725_{0.043}$	$0.932_{0.043}$
400	$671.99_{0.38}$	$8.84_{0.13}$	$0.32_{0.01}$	$28.110_{0.043}$	$0.920_{0.030}$
410	$661.74_{0.33}$	$10.98_{0.17}$	$0.403_{0.013}$	$27.478_{0.038}$	$0.906_{0.033}$
420	$650.92_{0.46}$	$13.51_{0.21}$	$0.499_{0.024}$	$26.801_{0.037}$	$0.891_{0.046}$
430	$639.09_{0.44}$	$16.48_{0.24}$	$0.613_{0.039}$	$26.056_{0.039}$	$0.875_{0.057}$
440	$626.49_{0.39}$	$19.96_{0.27}$	$0.744_{0.055}$	$25.253_{0.042}$	$0.857_{0.065}$
450	$613.53_{0.38}$	$24.02_{0.35}$	$0.894_{0.073}$	$24.409_{0.048}$	$0.837_{0.069}$
460	$600.27_{0.57}$	$28.75_{0.70}$	$1.066_{0.096}$	$23.525_{0.086}$	$0.816_{0.076}$
470	$586.50_{0.90}$	$34.3_{1.6}$	$1.26_{0.14}$	$22.58_{0.16}$	$0.793_{0.095}$
480	$571.9_{1.5}$	$40.7_{2.6}$	$1.48_{0.23}$	$21.56_{0.21}$	$0.77_{0.13}$
490	$556.3_{2.5}$	$48.4_{3.2}$	$1.73_{0.35}$	$20.44_{0.25}$	$0.74_{0.16}$
500	$538.9_{3.0}$	$57.7_{3.1}$	$2.01_{0.46}$	$19.16_{0.25}$	$0.71_{0.16}$
510	$519.1_{2.0}$	$69.1_{2.7}$	$2.32_{0.54}$	$17.67_{0.18}$	$0.67_{0.16}$
520	$495.9_{1.6}$	$83.5_{2.4}$	$2.67_{0.62}$	$15.93_{0.12}$	$0.62_{0.15}$

Table SI.XXX: GCMC-MBAR results for the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\rm CH_2}=20$ force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	P _{vap} (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
360	721.21 _{0.17}	$2.885_{0.047}$	$0.099_{0.059}$	$31.72_{0.18}$	$0.96_{0.57}$
370	$711.38_{0.48}$	$3.812_{0.043}$	$0.133_{0.056}$	$31.13_{0.11}$	$0.96_{0.40}$
380	$701.58_{0.81}$	$4.955_{0.041}$	$0.176_{0.054}$	$30.545_{0.066}$	$0.95_{0.29}$
390	$691.47_{0.70}$	$6.351_{0.048}$	$0.229_{0.054}$	$29.931_{0.055}$	$0.94_{0.22}$
400	$680.14_{0.36}$	$8.033_{0.065}$	$0.293_{0.055}$	$29.243_{0.046}$	$0.92_{0.17}$
410	$668.26_{0.63}$	$10.039_{0.086}$	$0.370_{0.057}$	$28.510_{0.057}$	$0.91_{0.14}$
420	$656.5_{1.6}$	$12.41_{0.11}$	$0.462_{0.061}$	$27.77_{0.10}$	$0.90_{0.12}$
430	$644.5_{2.3}$	$15.21_{0.13}$	$0.570_{0.065}$	$27.00_{0.14}$	$0.88_{0.10}$
440	$631.8_{1.8}$	$18.50_{0.16}$	$0.695_{0.068}$	$26.17_{0.12}$	$0.864_{0.085}$
450	$618.8_{1.1}$	$22.34_{0.20}$	$0.840_{0.072}$	$25.308_{0.082}$	$0.845_{0.072}$
460	$605.44_{0.73}$	$26.84_{0.33}$	$1.005_{0.072}$	$24.399_{0.086}$	$0.824_{0.060}$
470	$591.81_{0.70}$	$32.11_{0.65}$	$1.194_{0.069}$	$23.44_{0.13}$	$0.801_{0.049}$
480	$577.60_{0.68}$	$38.3_{1.2}$	$1.408_{0.074}$	$22.41_{0.19}$	$0.775_{0.048}$
490	$562.27_{0.91}$	$45.7_{2.1}$	$1.65_{0.12}$	$21.28_{0.23}$	$0.746_{0.064}$
500	$545.2_{2.6}$	$54.6_{2.9}$	$1.92_{0.23}$	$20.00_{0.19}$	$0.713_{0.092}$
510	$526.1_{5.6}$	$65.5_{3.4}$	$2.23_{0.38}$	$18.526_{0.093}$	$0.68_{0.12}$
520	$504.1_{9.3}$	$79.2_{3.3}$	$2.57_{0.57}$	$16.81_{0.22}$	$0.63_{0.14}$

SI.V Compressibility factor

SI.V.1 Validation of GCMC-MBAR

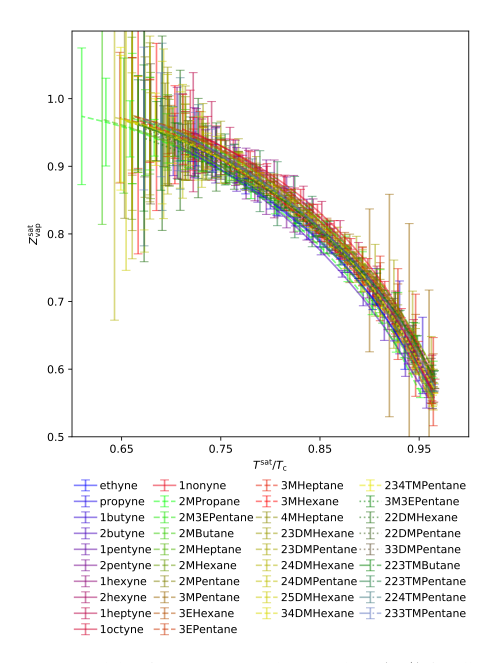


Figure SI.2: Compressibility factor in saturated vapor phase $(Z_{\rm vap}^{\rm sat})$ for all compounds simulated in Mick et al. and Soroush Barhaghi et al. Note that symmetric (normal) 95% confidence intervals are ill-suited when $Z_{\rm vap}^{\rm sat} \approx 1$, as this assumption can result in $Z_{\rm vap}^{\rm sat} > 1$.

SI.V.2 Case study: Cyclohexane optimization

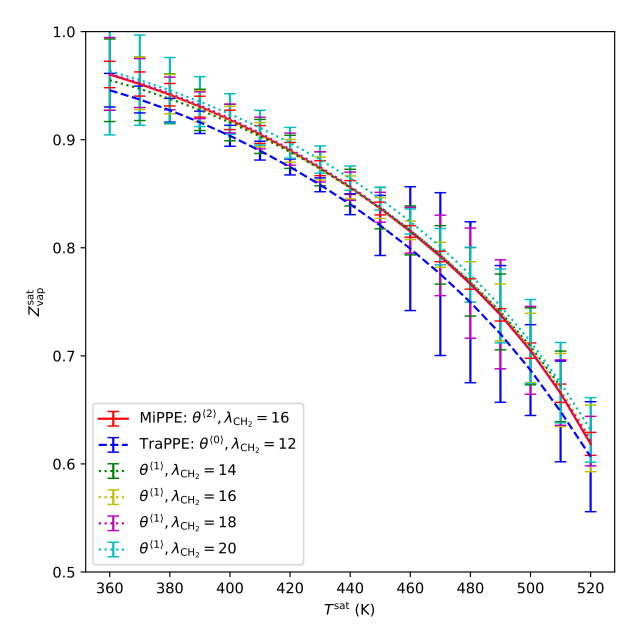


Figure SI.3: Compressibility factor in saturated vapor phase $(Z_{\mathrm{vap}}^{\mathrm{sat}})$ for iterations of cyclohexane optimization. Note that symmetric (normal) 95% confidence intervals are ill-suited when $Z_{\mathrm{vap}}^{\mathrm{sat}} \approx 1$, as this assumption can result in $Z_{\mathrm{vap}}^{\mathrm{sat}} > 1$.

SI.VI Tabulated phase equilibria for validation of GCMC-MBAR

SI.VI.1 Branched alkanes

Table SI.XXXI: GCMC-MBAR results for 2-methylpropane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
390	$388.8_{8.4}$	$87.1_{1.6}$	$2.742_{0.026}$	$9.76_{0.34}$	$0.56_{0.01}$
380	$416.6_{4.6}$	$66.4_{1.2}$	$2.292_{0.026}$	$11.69_{0.17}$	$0.63_{0.01}$
370	$439.7_{1.3}$	$51.95_{0.77}$	$1.903_{0.025}$	$13.219_{0.077}$	$0.692_{0.011}$
360	$459.18_{0.47}$	$41.22_{0.53}$	$1.566_{0.023}$	$14.442_{0.060}$	$0.74_{0.01}$
350	$476.39_{0.48}$	$32.82_{0.53}$	$1.276_{0.019}$	$15.485_{0.049}$	$0.78_{0.01}$
340	$492.09_{0.59}$	$26.09_{0.54}$	$1.028_{0.015}$	$16.404_{0.052}$	$0.81_{0.01}$
330	$506.65_{0.67}$	$20.62_{0.48}$	$0.817_{0.011}$	$17.227_{0.055}$	$0.84_{0.01}$
320	$520.48_{0.53}$	$16.16_{0.39}$	$0.6403_{0.0069}$	$17.980_{0.058}$	$0.865_{0.013}$
310	$533.93_{0.54}$	$12.53_{0.27}$	$0.4935_{0.0043}$	$18.686_{0.062}$	$0.889_{0.015}$
300	$546.58_{0.63}$	$9.57_{0.17}$	$0.3733_{0.0032}$	$19.332_{0.062}$	$0.909_{0.016}$
290	$558.46_{0.48}$	$7.19_{0.12}$	$0.2765_{0.0030}$	$19.925_{0.061}$	$0.927_{0.019}$
280	$570.30_{0.67}$	$5.300_{0.094}$	$0.2000_{0.0032}$	$20.498_{0.077}$	$0.942_{0.027}$
270	$582.5_{1.0}$	$3.815_{0.083}$	$0.1408_{0.0037}$	$21.07_{0.11}$	$0.955_{0.042}$
260	$593.47_{0.60}$	$2.672_{0.071}$	$0.0959_{0.0043}$	$21.57_{0.14}$	$0.965_{0.065}$
250	$603.0_{1.0}$	$1.815_{0.057}$	$0.0632_{0.0048}$	$22.01_{0.22}$	$0.97_{0.10}$

Table SI.XXXII: GCMC-MBAR results for 2-methylbutane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
440	$399.8_{3.5}$	$86.6_{2.1}$	$2.528_{0.020}$	$11.558_{0.090}$	$0.58_{0.01}$
430	$426.9_{2.6}$	$68.9_{1.7}$	$2.149_{0.011}$	$13.44_{0.11}$	$0.629_{0.012}$
420	$449.4_{1.4}$	$55.1_{1.2}$	$1.8149_{0.0059}$	$15.07_{0.11}$	$0.680_{0.012}$
410	$468.58_{0.70}$	$44.49_{0.66}$	$1.5225_{0.0039}$	$16.438_{0.094}$	$0.724_{0.010}$
400	$485.59_{0.52}$	$36.12_{0.31}$	$1.2671_{0.0037}$	$17.615_{0.066}$	$0.76_{0.01}$
390	$501.08_{0.68}$	$29.33_{0.15}$	$1.0450_{0.0036}$	$18.654_{0.048}$	$0.79_{0.01}$
380	$515.41_{0.89}$	$23.73_{0.12}$	$0.8532_{0.0034}$	$19.590_{0.055}$	$0.82_{0.01}$
370	$528.88_{0.98}$	$19.09_{0.11}$	$0.6887_{0.0031}$	$20.447_{0.064}$	$0.85_{0.01}$
360	$541.7_{1.0}$	$15.230_{0.098}$	$0.5489_{0.0029}$	$21.238_{0.067}$	$0.87_{0.01}$
350	$553.9_{1.1}$	$12.03_{0.10}$	$0.4315_{0.0028}$	$21.976_{0.071}$	$0.889_{0.010}$
340	$565.8_{1.3}$	$9.40_{0.11}$	$0.3341_{0.0029}$	$22.678_{0.085}$	$0.907_{0.016}$
330	$577.7_{1.7}$	$7.24_{0.12}$	$0.2541_{0.0033}$	$23.35_{0.12}$	$0.923_{0.025}$
320	$588.9_{1.7}$	$5.48_{0.14}$	$0.1896_{0.0039}$	$23.98_{0.16}$	$0.938_{0.040}$
310	$598.9_{1.2}$	$4.08_{0.14}$	$0.1385_{0.0047}$	$24.53_{0.20}$	$0.950_{0.064}$
300	$608.6_{1.2}$	$2.98_{0.14}$	$0.0988_{0.0055}$	$25.06_{0.28}$	$0.960_{0.10}$
290	$618.9_{1.3}$	$2.12_{0.13}$	$0.0686_{0.0064}$	$25.61_{0.41}$	$0.97_{0.16}$

Table SI.XXXIII: GCMC-MBAR results for 2-methylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}} (\mathrm{kg/m^3})$	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	$422.20_{0.56}$	$76.7_{1.5}$	$2.068_{0.017}$	$14.14_{0.11}$	$0.59_{0.01}$
460	$444.71_{0.79}$	$61.2_{1.2}$	$1.761_{0.013}$	$16.06_{0.12}$	$0.65_{0.01}$
450	$464.38_{0.87}$	$49.31_{0.88}$	$1.4918_{0.0096}$	$17.71_{0.11}$	$0.70_{0.01}$
440	$481.69_{0.77}$	$40.17_{0.64}$	$1.2557_{0.0068}$	$19.106_{0.090}$	$0.74_{0.01}$
430	$497.34_{0.70}$	$32.86_{0.47}$	$1.0491_{0.0045}$	$20.318_{0.074}$	$0.77_{0.01}$
420	$511.89_{0.78}$	$26.86_{0.36}$	$0.8690_{0.0027}$	$21.408_{0.068}$	$0.80_{0.01}$
410	$525.69_{0.84}$	$21.87_{0.29}$	$0.7130_{0.0017}$	$22.412_{0.069}$	$0.824_{0.011}$
400	$538.82_{0.88}$	$17.70_{0.25}$	$0.5788_{0.0020}$	$23.344_{0.069}$	$0.848_{0.014}$
390	$551.23_{0.87}$	$14.21_{0.22}$	$0.4645_{0.0031}$	$24.207_{0.075}$	$0.869_{0.018}$
380	$562.89_{0.80}$	$11.30_{0.19}$	$0.3681_{0.0042}$	$25.004_{0.096}$	$0.888_{0.024}$
370	$573.96_{0.88}$	$8.90_{0.16}$	$0.2878_{0.0051}$	$25.75_{0.14}$	$0.906_{0.032}$
360	$584.8_{1.1}$	$6.92_{0.14}$	$0.2215_{0.0059}$	$26.46_{0.18}$	$0.922_{0.042}$
350	$595.6_{1.2}$	$5.31_{0.12}$	$0.1677_{0.0065}$	$27.15_{0.22}$	$0.935_{0.056}$
340	$606.3_{1.2}$	$4.01_{0.10}$	$0.1245_{0.0070}$	$27.83_{0.27}$	$0.947_{0.076}$
330	$616.4_{1.0}$	$2.969_{0.085}$	$0.0906_{0.0074}$	$28.45_{0.32}$	$0.96_{0.10}$
320	$625.82_{0.57}$	$2.156_{0.068}$	$0.0644_{0.0078}$	$29.04_{0.40}$	$0.97_{0.14}$

Table SI.XXXIV: GCMC-MBAR results for 2-methylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
510	406.8 _{3.4}	88.4 _{2.0}	$2.092_{0.011}$	$14.14_{0.12}$	$0.56_{0.01}$
500	$431.2_{2.2}$	$70.0_{1.6}$	$1.7989_{0.0052}$	$16.506_{0.085}$	$0.620_{0.013}$
490	$452.1_{1.2}$	$56.5_{1.0}$	$1.5403_{0.0041}$	$18.472_{0.088}$	$0.671_{0.013}$
480	$470.38_{0.73}$	$46.30_{0.50}$	$1.3120_{0.0056}$	$20.092_{0.065}$	$0.712_{0.010}$
470	$486.88_{0.50}$	$38.21_{0.17}$	$1.1104_{0.0062}$	$21.497_{0.034}$	$0.75_{0.01}$
460	$502.20_{0.37}$	$31.56_{0.16}$	$0.9327_{0.0058}$	$22.762_{0.022}$	$0.77_{0.01}$
450	$516.52_{0.39}$	$26.01_{0.17}$	$0.7772_{0.0050}$	$23.921_{0.024}$	$0.80_{0.01}$
440	$530.01_{0.34}$	$21.34_{0.15}$	$0.6417_{0.0043}$	$24.994_{0.024}$	$0.82_{0.01}$
430	$542.86_{0.37}$	$17.41_{0.14}$	$0.5245_{0.0039}$	$25.998_{0.031}$	$0.84_{0.01}$
420	$554.93_{0.46}$	$14.10_{0.13}$	$0.4241_{0.0037}$	$26.928_{0.041}$	$0.86_{0.01}$
410	$566.22_{0.43}$	$11.32_{0.14}$	$0.3390_{0.0036}$	$27.789_{0.046}$	$0.881_{0.014}$
400	$577.21_{0.43}$	$8.99_{0.14}$	$0.2674_{0.0037}$	$28.610_{0.060}$	$0.896_{0.020}$
390	$588.02_{0.38}$	$7.06_{0.15}$	$0.2080_{0.0039}$	$29.403_{0.089}$	$0.910_{0.030}$
380	$598.35_{0.22}$	$5.47_{0.16}$	$0.1593_{0.0044}$	$30.15_{0.14}$	$0.923_{0.044}$
370	$608.04_{0.19}$	$4.18_{0.16}$	$0.1200_{0.0050}$	$30.85_{0.20}$	$0.935_{0.064}$
360	$617.49_{0.27}$	$3.14_{0.16}$	$0.0888_{0.0057}$	$31.52_{0.28}$	$0.945_{0.093}$
350	$627.22_{0.26}$	$2.32_{0.15}$	$0.0644_{0.0065}$	$32.20_{0.39}$	$0.96_{0.13}$

Table SI.XXXV: GCMC-MBAR results for 2-methylheptane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$401.3_{3.7}$	$88.7_{2.6}$	$1.947_{0.032}$	$15.17_{0.13}$	$0.56_{0.01}$
530	$426.3_{2.7}$	$71.5_{2.6}$	$1.683_{0.023}$	$17.60_{0.12}$	$0.610_{0.015}$
520	$448.1_{1.4}$	$58.0_{2.1}$	$1.447_{0.015}$	$19.75_{0.16}$	$0.660_{0.020}$
510	$467.12_{0.68}$	$47.5_{1.5}$	$1.2389_{0.0097}$	$21.59_{0.16}$	$0.702_{0.019}$
500	$483.89_{0.51}$	$39.23_{0.90}$	$1.0545_{0.0068}$	$23.17_{0.13}$	$0.738_{0.016}$
490	$499.04_{0.51}$	$32.49_{0.54}$	$0.8917_{0.0055}$	$24.571_{0.096}$	$0.770_{0.012}$
480	$513.07_{0.62}$	$26.88_{0.35}$	$0.7488_{0.0048}$	$25.842_{0.079}$	$0.80_{0.01}$
470	$526.33_{0.63}$	$22.18_{0.26}$	$0.6238_{0.0042}$	$27.019_{0.072}$	$0.82_{0.01}$
460	$538.86_{0.70}$	$18.22_{0.22}$	$0.5151_{0.0038}$	$28.112_{0.079}$	$0.844_{0.010}$
450	$550.62_{0.83}$	$14.88_{0.19}$	$0.4215_{0.0036}$	$29.125_{0.091}$	$0.865_{0.013}$
440	$561.64_{0.82}$	$12.07_{0.16}$	$0.3414_{0.0037}$	$30.06_{0.10}$	$0.883_{0.017}$
430	$572.07_{0.80}$	$9.71_{0.14}$	$0.2735_{0.0040}$	$30.94_{0.11}$	$0.900_{0.022}$
420	$582.24_{0.77}$	$7.74_{0.11}$	$0.2165_{0.0043}$	$31.78_{0.13}$	$0.915_{0.028}$
410	$592.62_{0.67}$	$6.108_{0.092}$	$0.1691_{0.0045}$	$32.62_{0.14}$	$0.927_{0.035}$
400	$603.09_{0.65}$	$4.758_{0.074}$	$0.1302_{0.0047}$	$33.44_{0.16}$	$0.940_{0.044}$
390	$612.67_{0.70}$	$3.655_{0.061}$	$0.0986_{0.0049}$	$34.20_{0.19}$	$0.950_{0.056}$
380	$621.39_{0.62}$	$2.766_{0.052}$	$0.0734_{0.0049}$	$34.89_{0.24}$	$0.960_{0.071}$
370	$630.41_{0.64}$	$2.057_{0.045}$	$0.0536_{0.0049}$	$35.58_{0.31}$	$0.967_{0.093}$

Table SI.XXXVI: GCMC-MBAR results for 3-methylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

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$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
480	$415.8_{1.5}$	84_{24}	$2.31_{0.24}$	$13.7_{1.3}$	$0.60_{0.12}$
470	$440.0_{2.3}$	67_{24}	$1.99_{0.14}$	$15.6_{1.7}$	$0.65_{0.17}$
460	$459.8_{2.5}$	55_{18}	$1.698_{0.059}$	$17.2_{1.6}$	$0.69_{0.16}$
450	$477.1_{1.9}$	$45.4_{7.6}$	$1.443_{0.012}$	$18.52_{0.94}$	$0.73_{0.11}$
440	$493.2_{1.2}$	$37.5_{1.5}$	$1.217_{0.014}$	$19.74_{0.27}$	$0.764_{0.036}$
430	$508.9_{1.4}$	$30.95_{0.38}$	$1.019_{0.015}$	$20.87_{0.11}$	$0.794_{0.013}$
420	$523.6_{1.1}$	$25.42_{0.36}$	$0.845_{0.014}$	$21.917_{0.094}$	$0.821_{0.016}$
410	$536.8_{1.4}$	$20.78_{0.32}$	$0.695_{0.014}$	$22.85_{0.12}$	$0.845_{0.020}$
400	$548.9_{1.7}$	$16.89_{0.24}$	$0.565_{0.013}$	$23.69_{0.15}$	$0.867_{0.021}$
390	$560.5_{1.5}$	$13.63_{0.21}$	$0.455_{0.012}$	$24.49_{0.13}$	$0.887_{0.018}$
380	$571.8_{1.2}$	$10.88_{0.20}$	$0.361_{0.010}$	$25.250_{0.099}$	$0.905_{0.016}$
370	$582.3_{1.1}$	$8.59_{0.18}$	$0.2829_{0.0089}$	$25.950_{0.093}$	$0.922_{0.018}$
360	$593.0_{1.1}$	$6.70_{0.16}$	$0.2185_{0.0077}$	$26.64_{0.12}$	$0.939_{0.024}$
350	$603.8_{1.0}$	$5.15_{0.14}$	$0.1661_{0.0067}$	$27.33_{0.14}$	$0.955_{0.035}$

Table SI.XXXVII: GCMC-MBAR results for 3-methylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
520	398.4 _{4.6}	98.1 _{1.2}	$2.307_{0.018}$	$13.25_{0.29}$	$0.55_{0.01}$
510	$426.2_{3.4}$	$77.3_{1.3}$	$1.996_{0.015}$	$15.86_{0.21}$	$0.61_{0.01}$
500	$449.0_{1.8}$	$62.6_{1.1}$	$1.720_{0.012}$	$17.92_{0.14}$	$0.66_{0.01}$
490	$467.85_{0.87}$	$51.64_{0.76}$	$1.474_{0.011}$	$19.56_{0.10}$	$0.70_{0.01}$
480	$484.12_{0.71}$	$42.82_{0.51}$	$1.2562_{0.0093}$	$20.963_{0.074}$	$0.74_{0.01}$
470	$498.93_{0.76}$	$35.54_{0.44}$	$1.0638_{0.0075}$	$22.215_{0.069}$	$0.77_{0.01}$
460	$512.9_{1.0}$	$29.46_{0.42}$	$0.8944_{0.0056}$	$23.366_{0.088}$	$0.80_{0.01}$
450	$526.2_{1.3}$	$24.34_{0.36}$	$0.7463_{0.0040}$	$24.442_{0.093}$	$0.82_{0.01}$
440	$539.09_{0.79}$	$20.03_{0.26}$	$0.6171_{0.0029}$	$25.454_{0.048}$	$0.84_{0.01}$
430	$551.25_{0.90}$	$16.38_{0.15}$	$0.5055_{0.0025}$	$26.397_{0.089}$	$0.86_{0.01}$
420	$563.0_{1.6}$	$13.302_{0.087}$	$0.4097_{0.0025}$	$27.29_{0.16}$	$0.88_{0.01}$
410	$574.5_{1.4}$	$10.70_{0.12}$	$0.3283_{0.0024}$	$28.15_{0.18}$	$0.90_{0.01}$
400	$585.3_{1.3}$	$8.53_{0.18}$	$0.2598_{0.0021}$	$28.96_{0.19}$	$0.918_{0.019}$
390	$595.7_{2.4}$	$6.72_{0.26}$	$0.2028_{0.0021}$	$29.72_{0.30}$	$0.933_{0.039}$
380	$605.9_{2.8}$	$5.23_{0.33}$	$0.1560_{0.0032}$	$30.45_{0.41}$	$0.947_{0.073}$
370	$615.4_{1.3}$	$4.01_{0.38}$	$0.1181_{0.0053}$	$31.13_{0.47}$	$0.96_{0.12}$
360	$624.92_{0.55}$	$3.04_{0.40}$	$0.0878_{0.0081}$	$31.78_{0.63}$	$0.97_{0.20}$

Table SI.XXXVIII: GCMC-MBAR results for 3-methylheptane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$417.2_{1.2}$	83 ₁₂	$1.890_{0.063}$	$16.23_{0.82}$	$0.582_{0.065}$
530	$440.3_{1.2}$	$66.7_{9.0}$	$1.634_{0.033}$	$18.58_{0.89}$	$0.635_{0.076}$
520	$460.5_{1.2}$	$54.5_{5.6}$	$1.407_{0.012}$	$20.58_{0.75}$	$0.681_{0.067}$
510	$478.1_{1.2}$	$45.1_{2.8}$	$1.2045_{0.0059}$	$22.26_{0.50}$	$0.719_{0.048}$
500	$494.01_{0.92}$	$37.5_{1.1}$	$1.0252_{0.0098}$	$23.73_{0.26}$	$0.752_{0.028}$
490	$508.73_{0.65}$	$31.13_{0.27}$	$0.867_{0.011}$	$25.068_{0.10}$	$0.781_{0.013}$
480	$522.52_{0.51}$	$25.80_{0.29}$	$0.728_{0.010}$	$26.295_{0.066}$	$0.81_{0.01}$
470	$535.46_{0.54}$	$21.31_{0.32}$	$0.6061_{0.0088}$	$27.431_{0.087}$	$0.83_{0.01}$
460	$547.59_{0.69}$	$17.52_{0.28}$	$0.5005_{0.0075}$	$28.484_{0.098}$	$0.85_{0.01}$
450	$559.13_{0.85}$	$14.31_{0.24}$	$0.4094_{0.0063}$	$29.47_{0.11}$	$0.87_{0.01}$
440	$570.31_{0.88}$	$11.61_{0.22}$	$0.3315_{0.0051}$	$30.41_{0.11}$	$0.89_{0.01}$
430	$581.03_{0.75}$	$9.34_{0.20}$	$0.2656_{0.0041}$	$31.30_{0.11}$	$0.91_{0.01}$
420	$591.11_{0.65}$	$7.44_{0.18}$	$0.2102_{0.0031}$	$32.12_{0.12}$	$0.924_{0.015}$
410	$600.65_{0.78}$	$5.87_{0.16}$	$0.1642_{0.0025}$	$32.89_{0.15}$	$0.937_{0.022}$
400	$610.1_{1.1}$	$4.58_{0.14}$	$0.1265_{0.0021}$	$33.65_{0.20}$	$0.949_{0.032}$
390	$620.1_{1.3}$	$3.52_{0.12}$	$0.0959_{0.0022}$	$34.43_{0.24}$	$0.959_{0.044}$
380	$630.35_{0.84}$	$2.67_{0.10}$	$0.0714_{0.0025}$	$35.23_{0.26}$	$0.967_{0.062}$
370	$640.62_{0.56}$	$1.986_{0.080}$	$0.0522_{0.0029}$	$36.01_{0.30}$	$0.975_{0.085}$

Table SI.XXXIX: GCMC-MBAR results for 3-ethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
520	$405.9_{2.5}$	$92.5_{3.7}$	$2.279_{0.020}$	$13.93_{0.27}$	$0.571_{0.018}$
510	$433.2_{1.4}$	$75.3_{2.9}$	$1.975_{0.011}$	$16.14_{0.27}$	$0.620_{0.021}$
500	$456.01_{0.97}$	$61.6_{1.8}$	$1.7031_{0.0070}$	$18.08_{0.24}$	$0.667_{0.019}$
490	$474.95_{0.93}$	$50.83_{0.86}$	$1.4607_{0.0069}$	$19.70_{0.17}$	$0.707_{0.013}$
480	$491.46_{0.83}$	$42.20_{0.30}$	$1.2454_{0.0072}$	$21.10_{0.11}$	$0.74_{0.01}$
470	$506.63_{0.68}$	$35.08_{0.18}$	$1.0550_{0.0067}$	$22.354_{0.060}$	$0.77_{0.01}$
460	$521.02_{0.50}$	$29.12_{0.22}$	$0.8870_{0.0058}$	$23.514_{0.046}$	$0.80_{0.01}$
450	$534.61_{0.44}$	$24.08_{0.23}$	$0.7398_{0.0048}$	$24.589_{0.057}$	$0.82_{0.01}$
440	$547.23_{0.42}$	$19.82_{0.21}$	$0.6115_{0.0037}$	$25.575_{0.058}$	$0.85_{0.01}$
430	$559.10_{0.45}$	$16.21_{0.17}$	$0.5007_{0.0028}$	$26.491_{0.043}$	$0.87_{0.01}$
420	$570.66_{0.61}$	$13.16_{0.11}$	$0.4058_{0.0023}$	$27.365_{0.031}$	$0.88_{0.01}$
410	$582.02_{0.53}$	$10.587_{0.082}$	$0.3251_{0.0022}$	$28.205_{0.034}$	$0.90_{0.01}$
400	$592.79_{0.41}$	$8.438_{0.086}$	$0.2572_{0.0023}$	$28.992_{0.054}$	$0.918_{0.015}$
390	$602.92_{0.51}$	$6.653_{0.098}$	$0.2007_{0.0026}$	$29.724_{0.089}$	$0.932_{0.023}$
380	$612.81_{0.53}$	$5.18_{0.10}$	$0.1544_{0.0030}$	$30.43_{0.12}$	$0.945_{0.033}$
370	$622.47_{0.37}$	$3.982_{0.097}$	$0.1168_{0.0035}$	$31.10_{0.15}$	$0.956_{0.048}$
360	$631.88_{0.27}$	$3.012_{0.090}$	$0.0868_{0.0040}$	$31.75_{0.21}$	$0.965_{0.068}$
350	$641.27_{0.30}$	$2.238_{0.081}$	$0.0632_{0.0044}$	$32.39_{0.29}$	$0.972_{0.097}$

Table SI.XL: GCMC-MBAR results for 3-ethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(K)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$425.4_{1.8}$	$79.4_{4.9}$	$1.866_{0.043}$	$16.85_{0.40}$	$0.598_{0.025}$
530	$449.1_{1.4}$	$64.7_{4.4}$	$1.614_{0.029}$	$19.05_{0.45}$	$0.647_{0.034}$
520	$468.9_{1.0}$	$53.1_{3.2}$	$1.390_{0.017}$	$20.94_{0.42}$	$0.691_{0.035}$
510	$486.13_{0.88}$	$44.0_{2.0}$	$1.1907_{0.0092}$	$22.56_{0.34}$	$0.728_{0.029}$
500	$501.71_{0.68}$	$36.6_{1.1}$	$1.0141_{0.0047}$	$23.99_{0.23}$	$0.760_{0.021}$
490	$516.13_{0.49}$	$30.49_{0.54}$	$0.8581_{0.0029}$	$25.28_{0.14}$	$0.789_{0.013}$
480	$529.64_{0.54}$	$25.32_{0.27}$	$0.7207_{0.0024}$	$26.475_{0.089}$	$0.81_{0.01}$
470	$542.43_{0.64}$	$20.95_{0.17}$	$0.6006_{0.0021}$	$27.583_{0.069}$	$0.84_{0.01}$
460	$554.62_{0.65}$	$17.25_{0.13}$	$0.4962_{0.0019}$	$28.624_{0.065}$	$0.86_{0.01}$
450	$566.22_{0.66}$	$14.11_{0.11}$	$0.4063_{0.0017}$	$29.600_{0.066}$	$0.88_{0.01}$
440	$577.33_{0.67}$	$11.466_{0.088}$	$0.3292_{0.0016}$	$30.521_{0.065}$	$0.90_{0.01}$
430	$588.23_{0.68}$	$9.237_{0.067}$	$0.2638_{0.0016}$	$31.409_{0.064}$	$0.91_{0.01}$
420	$598.94_{0.67}$	$7.369_{0.055}$	$0.2088_{0.0018}$	$32.268_{0.063}$	$0.927_{0.013}$
410	$609.11_{0.53}$	$5.815_{0.055}$	$0.1630_{0.0019}$	$33.074_{0.060}$	$0.939_{0.019}$
400	$618.75_{0.46}$	$4.534_{0.059}$	$0.1255_{0.0022}$	$33.831_{0.070}$	$0.950_{0.028}$
390	$628.18_{0.46}$	$3.487_{0.060}$	$0.0950_{0.0025}$	$34.56_{0.10}$	$0.960_{0.041}$
380	$637.85_{0.36}$	$2.641_{0.057}$	$0.0707_{0.0028}$	$35.31_{0.15}$	$0.968_{0.060}$
370	$647.74_{0.30}$	$1.965_{0.051}$	$0.0515_{0.0031}$	$36.05_{0.23}$	$0.974_{0.087}$

Table SI.XLI: GCMC-MBAR results for 4-methylheptane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$418.9_{1.9}$	$80.1_{1.6}$	$1.858_{0.017}$	$16.62_{0.17}$	$0.59_{0.01}$
530	$442.2_{1.4}$	$65.0_{1.3}$	$1.606_{0.015}$	$18.87_{0.15}$	$0.640_{0.010}$
520	$462.3_{1.1}$	$53.3_{1.1}$	$1.382_{0.013}$	$20.81_{0.15}$	$0.685_{0.011}$
510	$479.9_{1.1}$	$44.10_{0.88}$	$1.183_{0.010}$	$22.48_{0.16}$	$0.722_{0.010}$
500	$495.8_{1.0}$	$36.63_{0.73}$	$1.0060_{0.0083}$	$23.95_{0.16}$	$0.755_{0.011}$
490	$510.3_{1.0}$	$30.43_{0.61}$	$0.8501_{0.0062}$	$25.28_{0.16}$	$0.783_{0.011}$
480	$523.91_{0.96}$	$25.22_{0.47}$	$0.7132_{0.0044}$	$26.49_{0.15}$	$0.809_{0.011}$
470	$536.77_{0.78}$	$20.83_{0.32}$	$0.5936_{0.0029}$	$27.62_{0.11}$	$0.83_{0.01}$
460	$548.98_{0.49}$	$17.12_{0.19}$	$0.4897_{0.0020}$	$28.679_{0.071}$	$0.85_{0.01}$
450	$560.48_{0.39}$	$13.982_{0.094}$	$0.4002_{0.0014}$	$29.660_{0.044}$	$0.87_{0.01}$
440	$571.22_{0.45}$	$11.338_{0.062}$	$0.3238_{0.0012}$	$30.567_{0.048}$	$0.89_{0.01}$
430	$581.42_{0.48}$	$9.118_{0.092}$	$0.2591_{0.0013}$	$31.419_{0.060}$	$0.908_{0.012}$
420	$591.67_{0.57}$	$7.26_{0.12}$	$0.2049_{0.0018}$	$32.257_{0.078}$	$0.923_{0.023}$
410	$602.13_{0.53}$	$5.72_{0.14}$	$0.1598_{0.0025}$	$33.10_{0.12}$	$0.936_{0.037}$
400	$612.18_{0.47}$	$4.45_{0.14}$	$0.1228_{0.0033}$	$33.90_{0.18}$	$0.947_{0.056}$
390	$621.73_{0.54}$	$3.42_{0.14}$	$0.0928_{0.0042}$	$34.65_{0.25}$	$0.957_{0.082}$
380	$631.63_{0.45}$	$2.58_{0.12}$	$0.0690_{0.0050}$	$35.41_{0.37}$	$0.97_{0.12}$
370	$641.49_{0.32}$	$1.92_{0.11}$	$0.0502_{0.0057}$	$36.15_{0.52}$	$0.97_{0.17}$

Table SI.XLII: GCMC-MBAR results for 2,3-dimethylbutane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
480	$411.7_{1.1}$	$88.4_{2.2}$	$2.369_{0.027}$	$12.87_{0.14}$	$0.58_{0.01}$
470	$437.1_{1.0}$	$71.6_{2.1}$	$2.037_{0.019}$	$14.80_{0.17}$	$0.627_{0.013}$
460	$459.23_{0.86}$	$58.1_{1.6}$	$1.742_{0.013}$	$16.53_{0.18}$	$0.676_{0.015}$
450	$478.07_{0.79}$	$47.5_{1.1}$	$1.4813_{0.0074}$	$18.00_{0.16}$	$0.718_{0.014}$
440	$494.54_{0.81}$	$39.07_{0.72}$	$1.2513_{0.0041}$	$19.26_{0.13}$	$0.754_{0.012}$
430	$509.51_{0.77}$	$32.20_{0.43}$	$1.0493_{0.0024}$	$20.37_{0.10}$	$0.79_{0.01}$
420	$523.51_{0.64}$	$26.49_{0.25}$	$0.8726_{0.0019}$	$21.382_{0.079}$	$0.81_{0.01}$
410	$536.78_{0.54}$	$21.70_{0.15}$	$0.7189_{0.0020}$	$22.310_{0.063}$	$0.84_{0.01}$
400	$549.39_{0.59}$	$17.68_{0.10}$	$0.5863_{0.0021}$	$23.171_{0.060}$	$0.86_{0.01}$
390	$561.38_{0.73}$	$14.296_{0.082}$	$0.4727_{0.0023}$	$23.971_{0.062}$	$0.88_{0.01}$
380	$572.70_{0.68}$	$11.455_{0.085}$	$0.3764_{0.0025}$	$24.712_{0.056}$	$0.896_{0.011}$
370	$583.42_{0.40}$	$9.083_{0.092}$	$0.2957_{0.0028}$	$25.402_{0.050}$	$0.912_{0.016}$
360	$593.75_{0.21}$	$7.115_{0.095}$	$0.2289_{0.0033}$	$26.054_{0.064}$	$0.926_{0.024}$
350	$603.97_{0.20}$	$5.497_{0.091}$	$0.1742_{0.0037}$	$26.686_{0.091}$	$0.938_{0.034}$
340	$614.11_{0.22}$	$4.179_{0.082}$	$0.1302_{0.0042}$	$27.30_{0.13}$	$0.950_{0.047}$
330	$623.72_{0.26}$	$3.121_{0.069}$	$0.0953_{0.0047}$	$27.87_{0.18}$	$0.959_{0.066}$
320	$632.83_{0.34}$	$2.284_{0.057}$	$0.0682_{0.0050}$	$28.41_{0.25}$	$0.968_{0.092}$

Table SI.XLIII: GCMC-MBAR results for 2,3-dimethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$422.2_{1.1}$	$84.5_{3.7}$	$1.948_{0.048}$	$15.98_{0.12}$	$0.586_{0.012}$
530	$445.87_{0.82}$	$68.6_{3.6}$	$1.689_{0.036}$	$18.23_{0.22}$	$0.638_{0.022}$
520	$466.30_{0.66}$	$56.3_{3.0}$	$1.458_{0.024}$	$20.17_{0.26}$	$0.684_{0.027}$
510	$484.13_{0.59}$	$46.7_{2.1}$	$1.253_{0.015}$	$21.81_{0.23}$	$0.722_{0.026}$
500	$499.99_{0.52}$	$39.0_{1.4}$	$1.0703_{0.0092}$	$23.24_{0.18}$	$0.755_{0.022}$
490	$514.34_{0.48}$	$32.51_{0.80}$	$0.9084_{0.0054}$	$24.51_{0.13}$	$0.783_{0.016}$
480	$527.67_{0.54}$	$27.08_{0.47}$	$0.7657_{0.0032}$	$25.677_{0.096}$	$0.809_{0.012}$
470	$540.41_{0.64}$	$22.49_{0.30}$	$0.6406_{0.0019}$	$26.765_{0.082}$	$0.833_{0.010}$
460	$552.77_{0.76}$	$18.58_{0.23}$	$0.5313_{0.0015}$	$27.797_{0.085}$	$0.854_{0.011}$
450	$564.84_{0.87}$	$15.26_{0.22}$	$0.4366_{0.0018}$	$28.78_{0.11}$	$0.874_{0.015}$
440	$576.56_{0.95}$	$12.43_{0.23}$	$0.3553_{0.0025}$	$29.73_{0.14}$	$0.892_{0.022}$
430	$587.55_{0.97}$	$10.05_{0.23}$	$0.2860_{0.0034}$	$30.60_{0.18}$	$0.909_{0.032}$
420	$597.72_{0.83}$	$8.05_{0.21}$	$0.2274_{0.0044}$	$31.40_{0.21}$	$0.924_{0.043}$
410	$607.46_{0.53}$	$6.39_{0.18}$	$0.1787_{0.0053}$	$32.16_{0.22}$	$0.937_{0.055}$
400	$617.12_{0.48}$	$5.01_{0.14}$	$0.1384_{0.0061}$	$32.90_{0.24}$	$0.949_{0.070}$
390	$626.72_{0.62}$	$3.88_{0.10}$	$0.1056_{0.0067}$	$33.64_{0.28}$	$0.959_{0.087}$
380	$636.43_{0.54}$	$2.959_{0.070}$	$0.0792_{0.0071}$	$34.36_{0.35}$	$0.97_{0.11}$
370	$646.47_{0.59}$	$2.219_{0.049}$	$0.0583_{0.0072}$	$35.10_{0.46}$	$0.98_{0.14}$

Table SI.XLIV: GCMC-MBAR results for 2,3-dimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	P _{vap} (MPa)	$\Delta H_{ m v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
510	$427.9_{1.6}$	81 ₁₀	$2.082_{0.091}$	$15.20_{0.53}$	$0.606_{0.042}$
500	$451.7_{1.0}$	$66.3_{9.6}$	$1.800_{0.057}$	$17.17_{0.70}$	$0.654_{0.061}$
490	$471.41_{0.86}$	$54.7_{7.1}$	$1.549_{0.028}$	$18.83_{0.70}$	$0.696_{0.065}$
480	488.48 _{0.89}	$45.5_{4.0}$	$1.3245_{0.0092}$	$20.25_{0.54}$	$0.731_{0.053}$
470	$503.90_{0.73}$	$37.9_{1.7}$	$1.1257_{0.0050}$	$21.50_{0.32}$	$0.762_{0.034}$
460	$518.28_{0.52}$	$31.50_{0.62}$	$0.9501_{0.0072}$	$22.64_{0.14}$	$0.790_{0.020}$
450	$532.03_{0.63}$	$26.12_{0.25}$	$0.7956_{0.0080}$	$23.709_{0.054}$	$0.816_{0.013}$
440	$545.26_{0.62}$	$21.57_{0.18}$	$0.6606_{0.0080}$	$24.711_{0.037}$	$0.839_{0.011}$
430	$557.77_{0.54}$	$17.70_{0.16}$	$0.5435_{0.0075}$	$25.646_{0.038}$	$0.860_{0.010}$
420	$569.33_{0.53}$	$14.43_{0.15}$	$0.4426_{0.0070}$	$26.504_{0.037}$	$0.880_{0.011}$
410	$580.03_{0.59}$	$11.68_{0.14}$	$0.3565_{0.0063}$	$27.290_{0.040}$	$0.897_{0.012}$
400	$590.33_{0.66}$	$9.36_{0.13}$	$0.2837_{0.0057}$	$28.032_{0.047}$	$0.913_{0.014}$
390	$600.50_{0.61}$	$7.43_{0.11}$	$0.2229_{0.0052}$	$28.747_{0.057}$	$0.927_{0.019}$
380	$610.33_{0.59}$	$5.831_{0.094}$	$0.1726_{0.0048}$	$29.424_{0.079}$	$0.939_{0.025}$
370	$620.01_{0.64}$	$4.512_{0.077}$	$0.1315_{0.0046}$	$30.08_{0.11}$	$0.950_{0.036}$
360	$630.25_{0.55}$	$3.438_{0.066}$	$0.0984_{0.0045}$	$30.75_{0.14}$	$0.959_{0.050}$
350	$640.58_{0.49}$	$2.572_{0.058}$	$0.0722_{0.0046}$	$31.42_{0.20}$	$0.966_{0.072}$

Table SI.XLV: GCMC-MBAR results for 2,4-dimethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$406.0_{3.8}$	97.8 _{2.1}	$2.090_{0.036}$	$14.21_{0.22}$	$0.54_{0.01}$
530	$430.3_{3.2}$	$78.1_{2.3}$	$1.810_{0.028}$	$16.71_{0.16}$	$0.60_{0.01}$
520	$452.1_{1.9}$	$63.1_{2.2}$	$1.562_{0.019}$	$18.90_{0.15}$	$0.654_{0.016}$
510	$470.9_{1.1}$	$51.8_{1.9}$	$1.342_{0.013}$	$20.72_{0.19}$	$0.698_{0.020}$
500	$487.5_{1.1}$	$42.9_{1.4}$	$1.1472_{0.0080}$	$22.26_{0.19}$	$0.734_{0.020}$
490	$502.5_{1.2}$	$35.72_{0.85}$	$0.9745_{0.0054}$	$23.60_{0.16}$	$0.765_{0.017}$
480	$516.3_{1.4}$	$29.69_{0.47}$	$0.8222_{0.0043}$	$24.83_{0.13}$	$0.793_{0.012}$
470	$529.5_{1.6}$	$24.62_{0.27}$	$0.6886_{0.0037}$	$25.96_{0.11}$	$0.82_{0.01}$
460	$542.1_{1.7}$	$20.34_{0.18}$	$0.5720_{0.0031}$	$27.026_{0.098}$	$0.84_{0.01}$
450	$554.3_{1.8}$	$16.71_{0.13}$	$0.4708_{0.0026}$	$28.031_{0.098}$	$0.86_{0.01}$
440	$566.1_{1.8}$	$13.63_{0.10}$	$0.3836_{0.0024}$	$28.98_{0.10}$	$0.88_{0.01}$
430	$577.4_{1.6}$	$11.03_{0.11}$	$0.3092_{0.0023}$	$29.89_{0.11}$	$0.895_{0.010}$
420	$588.1_{1.4}$	$8.85_{0.15}$	$0.2463_{0.0022}$	$30.73_{0.13}$	$0.910_{0.019}$
410	$598.1_{1.1}$	$7.02_{0.19}$	$0.1937_{0.0024}$	$31.51_{0.17}$	$0.924_{0.033}$
400	$607.97_{0.72}$	$5.51_{0.22}$	$0.1503_{0.0031}$	$32.27_{0.22}$	$0.937_{0.056}$
390	$617.69_{0.31}$	$4.27_{0.23}$	$0.1148_{0.0042}$	$33.01_{0.29}$	$0.947_{0.088}$
380	$627.04_{0.22}$	$3.26_{0.22}$	$0.0862_{0.0055}$	$33.71_{0.42}$	$0.96_{0.14}$
370	$635.81_{0.19}$	$2.45_{0.20}$	$0.0636_{0.0068}$	$34.36_{0.62}$	$0.97_{0.20}$
360	$644.04_{0.32}$	$1.80_{0.18}$	$0.0460_{0.0080}$	$34.97_{0.91}$	$0.97_{0.30}$

Table SI.XLVI: GCMC-MBAR results for 2,4-dimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(K)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
500	$420.0_{1.7}$	$81.9_{2.0}$	$2.003_{0.018}$	$14.77_{0.12}$	$0.590_{0.011}$
490	$443.0_{1.0}$	$65.6_{1.4}$	$1.722_{0.014}$	$16.87_{0.12}$	$0.646_{0.011}$
480	$462.83_{0.68}$	$53.40_{0.95}$	$1.473_{0.011}$	$18.632_{0.099}$	$0.69_{0.01}$
470	$480.50_{0.59}$	$43.96_{0.74}$	$1.2527_{0.0088}$	$20.125_{0.093}$	$0.73_{0.01}$
460	$496.63_{0.64}$	$36.33_{0.64}$	$1.0583_{0.0067}$	$21.44_{0.10}$	$0.763_{0.010}$
450	$511.50_{0.67}$	$30.02_{0.54}$	$0.8873_{0.0049}$	$22.62_{0.10}$	$0.792_{0.012}$
440	$525.28_{0.65}$	$24.73_{0.40}$	$0.7377_{0.0038}$	$23.700_{0.095}$	$0.817_{0.012}$
430	$538.33_{0.62}$	$20.28_{0.26}$	$0.6076_{0.0032}$	$24.704_{0.085}$	$0.840_{0.011}$
420	$550.86_{0.66}$	$16.52_{0.19}$	$0.4955_{0.0029}$	$25.650_{0.080}$	$0.86_{0.01}$
410	$562.59_{0.67}$	$13.35_{0.21}$	$0.3996_{0.0024}$	$26.522_{0.078}$	$0.880_{0.014}$
400	$573.32_{0.65}$	$10.70_{0.23}$	$0.3184_{0.0024}$	$27.312_{0.092}$	$0.897_{0.023}$
390	$583.50_{0.57}$	$8.49_{0.25}$	$0.2504_{0.0031}$	$28.05_{0.13}$	$0.911_{0.037}$
380	$593.48_{0.38}$	$6.66_{0.25}$	$0.1942_{0.0044}$	$28.75_{0.18}$	$0.925_{0.056}$
370	$603.16_{0.41}$	$5.15_{0.26}$	$0.1483_{0.0059}$	$29.42_{0.26}$	$0.937_{0.084}$
360	$613.02_{0.44}$	$3.93_{0.28}$	$0.1113_{0.0076}$	$30.09_{0.36}$	$0.95_{0.13}$
350	$623.28_{0.28}$	$2.94_{0.28}$	$0.0819_{0.0093}$	$30.77_{0.52}$	$0.96_{0.18}$

Table SI.XLVII: GCMC-MBAR results for 2,5-dimethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
530	$405.4_{2.2}$	$92.0_{1.7}$	$1.963_{0.011}$	$14.59_{0.15}$	$0.55_{0.01}$
520	$429.1_{1.3}$	$73.2_{1.3}$	$1.6929_{0.0078}$	$17.04_{0.13}$	$0.61_{0.01}$
510	$450.43_{0.78}$	$59.06_{0.93}$	$1.4547_{0.0055}$	$19.17_{0.12}$	$0.66_{0.01}$
500	$469.12_{0.68}$	$48.33_{0.60}$	$1.2442_{0.0042}$	$20.967_{0.088}$	$0.71_{0.01}$
490	$485.77_{0.64}$	$39.90_{0.39}$	$1.0581_{0.0034}$	$22.495_{0.063}$	$0.74_{0.01}$
480	$500.89_{0.55}$	$33.04_{0.27}$	$0.8939_{0.0028}$	$23.843_{0.049}$	$0.77_{0.01}$
470	$514.91_{0.60}$	$27.35_{0.18}$	$0.7498_{0.0025}$	$25.066_{0.046}$	$0.80_{0.01}$
460	$528.20_{0.87}$	$22.57_{0.11}$	$0.6237_{0.0025}$	$26.198_{0.050}$	$0.83_{0.01}$
450	$540.9_{1.0}$	$18.541_{0.071}$	$0.5141_{0.0025}$	$27.259_{0.054}$	$0.85_{0.01}$
440	$553.03_{0.89}$	$15.132_{0.065}$	$0.4196_{0.0026}$	$28.254_{0.049}$	$0.87_{0.01}$
430	$564.47_{0.80}$	$12.255_{0.087}$	$0.3388_{0.0028}$	$29.185_{0.043}$	$0.883_{0.012}$
420	$575.38_{0.95}$	$9.83_{0.12}$	$0.2703_{0.0031}$	$30.060_{0.042}$	$0.899_{0.019}$
410	$586.0_{1.0}$	$7.81_{0.15}$	$0.2130_{0.0036}$	$30.900_{0.064}$	$0.914_{0.030}$
400	$596.47_{0.83}$	$6.13_{0.17}$	$0.1654_{0.0044}$	$31.71_{0.12}$	$0.927_{0.047}$
390	$606.50_{0.59}$	$4.75_{0.18}$	$0.1266_{0.0053}$	$32.48_{0.20}$	$0.939_{0.071}$
380	$615.90_{0.53}$	$3.63_{0.18}$	$0.0953_{0.0064}$	$33.20_{0.30}$	$0.95_{0.11}$
370	$624.82_{0.36}$	$2.73_{0.18}$	$0.0704_{0.0075}$	$33.87_{0.44}$	$0.96_{0.15}$
360	$633.58_{0.39}$	$2.01_{0.16}$	$0.0510_{0.0085}$	$34.52_{0.65}$	$0.97_{0.22}$

Table SI.XLVIII: GCMC-MBAR results for 3,4-dimethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
550	$416.97_{0.84}$	$91.7_{2.1}$	$2.100_{0.024}$	$15.29_{0.11}$	$0.57_{0.01}$
540	$441.75_{0.84}$	$75.2_{2.0}$	$1.828_{0.018}$	$17.53_{0.13}$	$0.619_{0.011}$
530	$462.97_{0.65}$	$61.9_{1.6}$	$1.584_{0.012}$	$19.50_{0.14}$	$0.664_{0.014}$
520	$481.27_{0.52}$	$51.4_{1.2}$	$1.3670_{0.0076}$	$21.20_{0.13}$	$0.703_{0.014}$
510	$497.59_{0.51}$	$42.88_{0.80}$	$1.1732_{0.0044}$	$22.68_{0.12}$	$0.737_{0.012}$
500	$512.45_{0.50}$	$35.87_{0.49}$	$1.0010_{0.0025}$	$23.994_{0.090}$	$0.77_{0.01}$
490	$526.21_{0.55}$	$30.00_{0.31}$	$0.8485_{0.0015}$	$25.192_{0.073}$	$0.79_{0.01}$
480	$539.20_{0.79}$	$25.02_{0.22}$	$0.7142_{0.0014}$	$26.302_{0.072}$	$0.82_{0.01}$
470	$551.6_{1.1}$	$20.79_{0.18}$	$0.5963_{0.0017}$	$27.341_{0.083}$	$0.84_{0.01}$
460	$563.5_{1.3}$	$17.18_{0.15}$	$0.4936_{0.0023}$	$28.319_{0.097}$	$0.858_{0.011}$
450	$574.7_{1.3}$	$14.10_{0.14}$	$0.4050_{0.0028}$	$29.24_{0.11}$	$0.877_{0.014}$
440	$585.4_{1.2}$	$11.49_{0.12}$	$0.3287_{0.0033}$	$30.09_{0.11}$	$0.893_{0.018}$
430	$595.6_{1.1}$	$9.29_{0.11}$	$0.2642_{0.0037}$	$30.90_{0.11}$	$0.909_{0.022}$
420	$605.36_{0.91}$	$7.437_{0.097}$	$0.2097_{0.0040}$	$31.66_{0.11}$	$0.922_{0.027}$
410	$614.74_{0.65}$	$5.893_{0.085}$	$0.1644_{0.0042}$	$32.38_{0.12}$	$0.935_{0.033}$
400	$624.02_{0.41}$	$4.615_{0.075}$	$0.1271_{0.0044}$	$33.08_{0.14}$	$0.946_{0.041}$
390	$633.41_{0.31}$	$3.566_{0.067}$	$0.0967_{0.0045}$	$33.78_{0.17}$	$0.955_{0.053}$
380	$642.95_{0.22}$	$2.715_{0.059}$	$0.0724_{0.0045}$	$34.48_{0.22}$	$0.964_{0.070}$
370	$652.67_{0.19}$	$2.032_{0.051}$	$0.0531_{0.0046}$	$35.18_{0.29}$	$0.970_{0.094}$

Table SI.XLIX: GCMC-MBAR results for 2,3,4-trimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
550	$415.7_{1.4}$	$96.1_{3.2}$	$2.199_{0.049}$	$14.71_{0.16}$	$0.57_{0.01}$
540	$442.1_{1.2}$	$79.3_{3.4}$	$1.920_{0.039}$	$16.92_{0.21}$	$0.616_{0.015}$
530	$464.77_{0.78}$	$65.5_{3.0}$	$1.669_{0.029}$	$18.91_{0.25}$	$0.661_{0.021}$
520	$483.71_{0.55}$	$54.4_{2.4}$	$1.444_{0.021}$	$20.61_{0.24}$	$0.701_{0.023}$
510	$499.94_{0.54}$	$45.5_{1.7}$	$1.243_{0.015}$	$22.07_{0.20}$	$0.736_{0.021}$
500	$514.37_{0.58}$	$38.2_{1.1}$	$1.064_{0.011}$	$23.34_{0.15}$	$0.766_{0.017}$
490	$527.58_{0.61}$	$32.03_{0.69}$	$0.9054_{0.0087}$	$24.49_{0.10}$	$0.793_{0.013}$
480	$540.01_{0.71}$	$26.83_{0.52}$	$0.7652_{0.0066}$	$25.541_{0.081}$	$0.816_{0.011}$
470	$552.10_{0.82}$	$22.39_{0.43}$	$0.6419_{0.0048}$	$26.539_{0.074}$	$0.838_{0.011}$
460	$564.06_{0.86}$	$18.59_{0.35}$	$0.5341_{0.0032}$	$27.497_{0.070}$	$0.858_{0.013}$
450	$575.70_{0.84}$	$15.35_{0.26}$	$0.4403_{0.0020}$	$28.409_{0.064}$	$0.876_{0.014}$
440	$586.68_{0.81}$	$12.58_{0.18}$	$0.3594_{0.0017}$	$29.258_{0.061}$	$0.892_{0.013}$
430	$596.87_{0.75}$	$10.22_{0.11}$	$0.2903_{0.0018}$	$30.041_{0.060}$	$0.907_{0.013}$
420	$606.38_{0.61}$	$8.235_{0.070}$	$0.2318_{0.0019}$	$30.766_{0.064}$	$0.921_{0.013}$
410	$615.73_{0.46}$	$6.567_{0.082}$	$0.1828_{0.0020}$	$31.466_{0.074}$	$0.933_{0.018}$
400	$625.40_{0.41}$	$5.18_{0.11}$	$0.1422_{0.0022}$	$32.176_{0.095}$	$0.944_{0.029}$
390	$634.86_{0.32}$	$4.03_{0.13}$	$0.1090_{0.0025}$	$32.86_{0.14}$	$0.953_{0.046}$
380	$643.49_{0.23}$	$3.09_{0.13}$	$0.0822_{0.0031}$	$33.48_{0.20}$	$0.962_{0.071}$
370	$652.26_{0.34}$	$2.33_{0.13}$	$0.0609_{0.0038}$	$34.10_{0.29}$	$0.97_{0.11}$

Table SI.L: GCMC-MBAR results for 2-methyl-3-ethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
540	$433.4_{2.7}$	$82.6_{3.0}$	$1.946_{0.032}$	$16.49_{0.24}$	$0.599_{0.012}$
530	$455.6_{2.0}$	$67.4_{2.8}$	$1.688_{0.024}$	$18.61_{0.30}$	$0.649_{0.019}$
520	$474.7_{1.3}$	$55.6_{2.2}$	$1.458_{0.016}$	$20.42_{0.29}$	$0.692_{0.021}$
510	$491.79_{0.85}$	$46.4_{1.5}$	$1.253_{0.011}$	$21.98_{0.23}$	$0.728_{0.019}$
500	$507.34_{0.64}$	$38.79_{0.93}$	$1.0711_{0.0081}$	$23.35_{0.17}$	$0.759_{0.015}$
490	$521.73_{0.53}$	$32.44_{0.55}$	$0.9094_{0.0062}$	$24.59_{0.11}$	$0.786_{0.011}$
480	$535.29_{0.40}$	$27.07_{0.36}$	$0.7666_{0.0049}$	$25.746_{0.076}$	$0.81_{0.01}$
470	$548.25_{0.30}$	$22.50_{0.28}$	$0.6411_{0.0038}$	$26.827_{0.056}$	$0.83_{0.01}$
460	$560.62_{0.34}$	$18.60_{0.23}$	$0.5316_{0.0028}$	$27.843_{0.047}$	$0.85_{0.01}$
450	$572.31_{0.35}$	$15.28_{0.20}$	$0.4368_{0.0020}$	$28.791_{0.049}$	$0.87_{0.01}$
440	$583.21_{0.31}$	$12.46_{0.17}$	$0.3554_{0.0014}$	$29.667_{0.055}$	$0.891_{0.012}$
430	$593.38_{0.30}$	$10.08_{0.14}$	$0.2861_{0.0014}$	$30.476_{0.064}$	$0.907_{0.015}$
420	$602.98_{0.33}$	$8.09_{0.13}$	$0.2277_{0.0018}$	$31.230_{0.085}$	$0.921_{0.020}$
410	$612.40_{0.54}$	$6.42_{0.11}$	$0.1789_{0.0022}$	$31.96_{0.12}$	$0.933_{0.026}$
400	$622.18_{0.63}$	$5.04_{0.10}$	$0.1387_{0.0027}$	$32.69_{0.15}$	$0.944_{0.036}$
390	$632.26_{0.45}$	$3.907_{0.092}$	$0.1058_{0.0031}$	$33.42_{0.18}$	$0.954_{0.049}$
380	$641.81_{0.31}$	$2.981_{0.084}$	$0.0794_{0.0036}$	$34.12_{0.22}$	$0.963_{0.069}$
370	$650.71_{0.52}$	$2.238_{0.074}$	$0.0585_{0.0039}$	$34.76_{0.31}$	$0.971_{0.096}$

Table SI.LI: GCMC-MBAR results for 3-methyl-3-ethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$\rho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
560	429.6 _{1.9}	$93.3_{3.0}$	$2.224_{0.025}$	$15.51_{0.15}$	$0.585_{0.013}$
550	$453.4_{1.3}$	$76.7_{2.3}$	$1.949_{0.017}$	$17.72_{0.13}$	$0.635_{0.015}$
540	$473.9_{1.5}$	$64.1_{1.4}$	$1.702_{0.012}$	$19.55_{0.10}$	$0.676_{0.012}$
530	$491.5_{1.3}$	$54.07_{0.82}$	$1.4787_{0.0089}$	$21.089_{0.076}$	$0.71_{0.01}$
520	$506.85_{0.72}$	$45.77_{0.59}$	$1.2780_{0.0068}$	$22.420_{0.057}$	$0.74_{0.01}$
510	$520.59_{0.37}$	$38.74_{0.54}$	$1.0984_{0.0049}$	$23.608_{0.064}$	$0.76_{0.01}$
500	$533.49_{0.41}$	$32.75_{0.51}$	$0.9383_{0.0034}$	$24.705_{0.078}$	$0.787_{0.011}$
490	$545.92_{0.68}$	$27.61_{0.41}$	$0.7960_{0.0027}$	$25.736_{0.080}$	$0.808_{0.012}$
480	$557.91_{0.83}$	$23.19_{0.27}$	$0.6704_{0.0029}$	$26.707_{0.072}$	$0.827_{0.012}$
470	$569.50_{0.65}$	$19.38_{0.14}$	$0.5599_{0.0033}$	$27.623_{0.058}$	$0.84_{0.01}$
460	$580.74_{0.49}$	$16.096_{0.094}$	$0.4634_{0.0033}$	$28.496_{0.048}$	$0.86_{0.01}$
450	$591.73_{0.55}$	$13.27_{0.13}$	$0.3799_{0.0029}$	$29.336_{0.044}$	$0.87_{0.01}$
440	$602.39_{0.47}$	$10.84_{0.16}$	$0.3082_{0.0025}$	$30.140_{0.046}$	$0.888_{0.011}$
430	$612.22_{0.39}$	$8.78_{0.18}$	$0.2472_{0.0021}$	$30.879_{0.073}$	$0.900_{0.017}$
420	$621.24_{0.63}$	$7.03_{0.20}$	$0.1959_{0.0020}$	$31.56_{0.12}$	$0.911_{0.026}$
410	$630.11_{0.71}$	$5.57_{0.23}$	$0.1533_{0.0023}$	$32.21_{0.17}$	$0.922_{0.041}$
400	$639.21_{0.40}$	$4.36_{0.28}$	$0.1182_{0.0032}$	$32.87_{0.24}$	$0.932_{0.068}$
390	$648.92_{0.26}$	$3.36_{0.32}$	$0.0898_{0.0047}$	$33.56_{0.35}$	$0.94_{0.11}$

Table SI.LII: GCMC-MBAR results for 2,2-dimethylbutane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	$412.4_{5.2}$	$85.4_{8.5}$	$2.32_{0.11}$	$12.77_{0.20}$	$0.598_{0.027}$
460	$438.1_{4.9}$	$69.3_{8.7}$	$1.996_{0.071}$	$14.62_{0.34}$	$0.649_{0.049}$
450	$459.2_{3.1}$	$56.9_{7.2}$	$1.709_{0.036}$	$16.16_{0.45}$	$0.691_{0.061}$
440	$477.5_{1.5}$	$47.1_{4.6}$	$1.453_{0.011}$	$17.47_{0.41}$	$0.727_{0.057}$
430	$494.03_{0.95}$	$39.0_{2.1}$	$1.2267_{0.0045}$	$18.63_{0.25}$	$0.759_{0.039}$
420	$508.92_{0.71}$	$32.17_{0.58}$	$1.0273_{0.0094}$	$19.664_{0.098}$	$0.788_{0.020}$
410	$522.56_{0.38}$	$26.47_{0.14}$	$0.853_{0.010}$	$20.599_{0.035}$	$0.81_{0.01}$
400	$535.52_{0.29}$	$21.68_{0.16}$	$0.7014_{0.0094}$	$21.466_{0.026}$	$0.84_{0.01}$
390	$548.22_{0.34}$	$17.64_{0.11}$	$0.5708_{0.0086}$	$22.291_{0.027}$	$0.860_{0.010}$
380	$560.63_{0.37}$	$14.238_{0.067}$	$0.4591_{0.0081}$	$23.076_{0.038}$	$0.879_{0.013}$
370	$572.47_{0.42}$	$11.382_{0.058}$	$0.3644_{0.0077}$	$23.813_{0.053}$	$0.897_{0.017}$
360	$583.49_{0.45}$	$8.997_{0.074}$	$0.2850_{0.0074}$	$24.488_{0.062}$	$0.912_{0.022}$
350	$593.61_{0.52}$	$7.021_{0.099}$	$0.2196_{0.0070}$	$25.098_{0.081}$	$0.926_{0.029}$
340	$603.19_{0.55}$	$5.40_{0.12}$	$0.1662_{0.0067}$	$25.66_{0.11}$	$0.939_{0.041}$
330	$612.67_{0.41}$	$4.09_{0.14}$	$0.1236_{0.0065}$	$26.22_{0.16}$	$0.950_{0.061}$
320	$622.10_{0.33}$	$3.03_{0.13}$	$0.0899_{0.0065}$	$26.75_{0.23}$	$0.959_{0.091}$
310	$631.64_{0.30}$	$2.21_{0.12}$	$0.0640_{0.0068}$	$27.29_{0.34}$	$0.97_{0.14}$

Table SI.LIII: GCMC-MBAR results for 2,2-dimethylhexane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
530	$405.4_{1.1}$	$88.4_{2.4}$	$1.979_{0.012}$	$14.84_{0.19}$	$0.580_{0.013}$
520	$431.48_{0.93}$	$72.7_{1.8}$	$1.7154_{0.0075}$	$16.99_{0.19}$	$0.623_{0.013}$
510	$453.8_{1.0}$	$59.8_{1.2}$	$1.4785_{0.0047}$	$18.93_{0.16}$	$0.666_{0.012}$
500	$472.8_{1.0}$	$49.35_{0.64}$	$1.2675_{0.0038}$	$20.62_{0.13}$	$0.71_{0.01}$
490	$489.55_{0.82}$	$40.92_{0.33}$	$1.0801_{0.0034}$	$22.090_{0.076}$	$0.74_{0.01}$
480	$504.38_{0.54}$	$33.96_{0.19}$	$0.9145_{0.0030}$	$23.386_{0.037}$	$0.77_{0.01}$
470	$517.67_{0.43}$	$28.16_{0.15}$	$0.7689_{0.0025}$	$24.539_{0.032}$	$0.80_{0.01}$
460	$530.06_{0.45}$	$23.29_{0.13}$	$0.6417_{0.0022}$	$25.596_{0.035}$	$0.82_{0.01}$
450	$542.23_{0.55}$	$19.19_{0.13}$	$0.5309_{0.0021}$	$26.601_{0.035}$	$0.84_{0.01}$
440	$554.29_{0.63}$	$15.71_{0.16}$	$0.4353_{0.0023}$	$27.566_{0.050}$	$0.865_{0.012}$
430	$565.90_{0.55}$	$12.78_{0.20}$	$0.3531_{0.0029}$	$28.479_{0.074}$	$0.883_{0.020}$
420	$577.12_{0.44}$	$10.30_{0.25}$	$0.2833_{0.0040}$	$29.35_{0.10}$	$0.899_{0.033}$
410	$588.18_{0.45}$	$8.23_{0.27}$	$0.2243_{0.0053}$	$30.19_{0.16}$	$0.914_{0.050}$
400	$598.75_{0.43}$	$6.49_{0.27}$	$0.1753_{0.0068}$	$30.99_{0.23}$	$0.927_{0.073}$
390	$608.66_{0.49}$	$5.06_{0.26}$	$0.1349_{0.0082}$	$31.74_{0.32}$	$0.94_{0.10}$
380	$618.76_{0.62}$	$3.89_{0.23}$	$0.1021_{0.0096}$	$32.48_{0.43}$	$0.95_{0.14}$
370	$629.14_{0.47}$	$2.94_{0.20}$	$0.076_{0.011}$	$33.23_{0.60}$	$0.96_{0.20}$

Table SI.LIV: GCMC-MBAR results for 2,2-dimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\rm vap}^{\rm sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
500	$411.2_{4.8}$	$88.4_{3.0}$	$2.127_{0.025}$	$13.748_{0.081}$	$0.580_{0.013}$
490	$436.8_{4.1}$	$71.5_{2.2}$	$1.835_{0.014}$	$15.833_{0.068}$	$0.631_{0.015}$
480	$457.9_{2.5}$	$58.3_{1.4}$	$1.5750_{0.0068}$	$17.601_{0.049}$	$0.678_{0.014}$
470	$476.0_{1.4}$	$48.10_{0.81}$	$1.3443_{0.0036}$	$19.097_{0.040}$	$0.717_{0.012}$
460	$492.22_{0.92}$	$39.84_{0.47}$	$1.1402_{0.0035}$	$20.399_{0.036}$	$0.75_{0.01}$
450	$506.85_{0.87}$	$33.01_{0.29}$	$0.9600_{0.0041}$	$21.553_{0.034}$	$0.78_{0.01}$
440	$520.4_{1.0}$	$27.29_{0.18}$	$0.8020_{0.0047}$	$22.600_{0.037}$	$0.80_{0.01}$
430	$533.27_{0.96}$	$22.47_{0.10}$	$0.6642_{0.0050}$	$23.572_{0.034}$	$0.83_{0.01}$
420	$545.67_{0.39}$	$18.398_{0.066}$	$0.5448_{0.0050}$	$24.481_{0.029}$	$0.85_{0.01}$
410	$557.49_{0.37}$	$14.958_{0.060}$	$0.4422_{0.0048}$	$25.331_{0.052}$	$0.87_{0.01}$
400	$568.93_{0.50}$	$12.058_{0.076}$	$0.3547_{0.0047}$	$26.137_{0.064}$	$0.886_{0.013}$
390	$580.08_{0.47}$	$9.622_{0.094}$	$0.2810_{0.0046}$	$26.908_{0.069}$	$0.902_{0.018}$
380	$590.67_{0.42}$	$7.59_{0.11}$	$0.2195_{0.0045}$	$27.631_{0.075}$	$0.917_{0.023}$
370	$601.07_{0.39}$	$5.91_{0.14}$	$0.1689_{0.0043}$	$28.333_{0.091}$	$0.931_{0.031}$
360	$611.44_{0.41}$	$4.53_{0.18}$	$0.1279_{0.0041}$	$29.02_{0.13}$	$0.945_{0.045}$
350	$620.82_{0.40}$	$3.42_{0.21}$	$0.0950_{0.0040}$	$29.64_{0.19}$	$0.956_{0.071}$

Table SI.LV: GCMC-MBAR results for 2,2-dimethylpropane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
410	$409.5_{2.6}$	$78.8_{2.6}$	$2.218_{0.029}$	$11.33_{0.11}$	$0.596_{0.013}$
400	$433.9_{2.3}$	$61.3_{1.8}$	$1.869_{0.018}$	$13.140_{0.083}$	$0.661_{0.014}$
390	$454.8_{1.5}$	$48.8_{1.1}$	$1.565_{0.010}$	$14.591_{0.051}$	$0.713_{0.013}$
380	$472.4_{1.1}$	$39.38_{0.72}$	$1.3004_{0.0056}$	$15.750_{0.045}$	$0.754_{0.011}$
370	$487.88_{0.98}$	$31.87_{0.48}$	$1.0704_{0.0035}$	$16.732_{0.045}$	$0.788_{0.011}$
360	$502.34_{0.89}$	$25.71_{0.33}$	$0.8720_{0.0036}$	$17.617_{0.044}$	$0.817_{0.012}$
350	$516.37_{0.86}$	$20.61_{0.21}$	$0.7019_{0.0043}$	$18.443_{0.046}$	$0.844_{0.011}$
340	$529.79_{0.93}$	$16.38_{0.18}$	$0.5574_{0.0045}$	$19.210_{0.057}$	$0.868_{0.011}$
330	$542.27_{0.96}$	$12.88_{0.23}$	$0.4363_{0.0041}$	$19.910_{0.078}$	$0.891_{0.016}$
320	$553.7_{1.0}$	$10.00_{0.28}$	$0.3360_{0.0037}$	$20.54_{0.11}$	$0.911_{0.029}$
310	$564.4_{1.4}$	$7.66_{0.32}$	$0.2543_{0.0044}$	$21.12_{0.18}$	$0.930_{0.052}$
300	$575.3_{1.8}$	$5.77_{0.33}$	$0.1887_{0.0061}$	$21.68_{0.28}$	$0.946_{0.087}$
290	$587.1_{1.3}$	$4.27_{0.32}$	$0.1368_{0.0084}$	$22.27_{0.38}$	$0.96_{0.14}$
280	$599.1_{1.1}$	$3.08_{0.29}$	$0.096_{0.011}$	$22.85_{0.55}$	$0.97_{0.22}$

Table SI.LVI: GCMC-MBAR results for 3,3-dimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
510	$433.8_{1.4}$	$83.3_{2.1}$	$2.128_{0.013}$	$14.92_{0.15}$	$0.604_{0.012}$
500	$456.2_{1.8}$	$68.2_{1.6}$	$1.8431_{0.0076}$	$16.79_{0.16}$	$0.651_{0.013}$
490	$475.3_{1.5}$	$56.40_{0.99}$	$1.5893_{0.0041}$	$18.38_{0.12}$	$0.693_{0.012}$
480	$492.10_{0.90}$	$47.00_{0.51}$	$1.3630_{0.0031}$	$19.752_{0.070}$	$0.73_{0.01}$
470	$507.53_{0.52}$	$39.28_{0.29}$	$1.1617_{0.0029}$	$20.968_{0.041}$	$0.76_{0.01}$
460	$521.91_{0.42}$	$32.81_{0.22}$	$0.9831_{0.0027}$	$22.072_{0.042}$	$0.78_{0.01}$
450	$535.23_{0.39}$	$27.34_{0.16}$	$0.8258_{0.0025}$	$23.077_{0.038}$	$0.81_{0.01}$
440	$547.46_{0.40}$	$22.676_{0.096}$	$0.6879_{0.0023}$	$23.990_{0.031}$	$0.83_{0.01}$
430	$559.01_{0.43}$	$18.706_{0.065}$	$0.5678_{0.0022}$	$24.839_{0.027}$	$0.85_{0.01}$
420	$570.36_{0.53}$	$15.325_{0.054}$	$0.4642_{0.0020}$	$25.651_{0.032}$	$0.87_{0.01}$
410	$581.43_{0.85}$	$12.456_{0.067}$	$0.3754_{0.0020}$	$26.425_{0.058}$	$0.89_{0.01}$
400	$591.91_{0.96}$	$10.03_{0.10}$	$0.3000_{0.0021}$	$27.146_{0.083}$	$0.901_{0.013}$
390	$601.87_{0.71}$	$8.00_{0.14}$	$0.2368_{0.0024}$	$27.82_{0.10}$	$0.915_{0.024}$
380	$611.41_{0.54}$	$6.30_{0.17}$	$0.1842_{0.0031}$	$28.46_{0.14}$	$0.927_{0.040}$
370	$620.56_{0.38}$	$4.90_{0.19}$	$0.1411_{0.0040}$	$29.06_{0.21}$	$0.938_{0.066}$
360	$629.62_{0.19}$	$3.76_{0.19}$	$0.1063_{0.0052}$	$29.64_{0.30}$	$0.95_{0.10}$

Table SI.LVII: GCMC-MBAR results for 2,2,3-trimethylbutane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
520	$427.6_{2.7}$	90.4 _{3.4}	$2.310_{0.041}$	$14.20_{0.20}$	$0.592_{0.014}$
510	$451.1_{1.5}$	$73.9_{3.6}$	$2.014_{0.030}$	$16.22_{0.27}$	$0.644_{0.025}$
500	$471.00_{0.92}$	$61.5_{2.9}$	$1.748_{0.020}$	$17.84_{0.29}$	$0.684_{0.029}$
490	$488.37_{0.82}$	$51.6_{1.9}$	$1.509_{0.014}$	$19.22_{0.23}$	$0.719_{0.023}$
480	$503.84_{0.76}$	$43.39_{0.99}$	$1.295_{0.011}$	$20.44_{0.15}$	$0.749_{0.015}$
470	$518.11_{0.61}$	$36.42_{0.54}$	$1.1046_{0.0087}$	$21.544_{0.083}$	$0.78_{0.01}$
460	$531.60_{0.50}$	$30.51_{0.45}$	$0.9358_{0.0068}$	$22.566_{0.064}$	$0.80_{0.01}$
450	$544.35_{0.55}$	$25.49_{0.41}$	$0.7870_{0.0049}$	$23.512_{0.070}$	$0.83_{0.01}$
440	$556.68_{0.54}$	$21.21_{0.32}$	$0.6564_{0.0032}$	$24.402_{0.069}$	$0.85_{0.01}$
430	$568.85_{0.50}$	$17.55_{0.23}$	$0.5424_{0.0020}$	$25.255_{0.062}$	$0.87_{0.01}$
420	$580.37_{0.55}$	$14.42_{0.15}$	$0.4436_{0.0014}$	$26.050_{0.058}$	$0.88_{0.01}$
410	$590.79_{0.86}$	$11.748_{0.088}$	$0.3590_{0.0014}$	$26.769_{0.064}$	$0.90_{0.01}$
400	$600.6_{1.0}$	$9.476_{0.069}$	$0.2870_{0.0014}$	$27.440_{0.077}$	$0.91_{0.01}$
390	$610.62_{0.83}$	$7.560_{0.096}$	$0.2267_{0.0013}$	$28.107_{0.086}$	$0.927_{0.013}$
380	$621.8_{1.5}$	$5.96_{0.14}$	$0.1765_{0.0014}$	$28.82_{0.16}$	$0.940_{0.025}$
370	$633.5_{2.1}$	$4.63_{0.17}$	$0.1353_{0.0019}$	$29.56_{0.26}$	$0.952_{0.044}$
360	$643.07_{0.71}$	$3.54_{0.20}$	$0.1019_{0.0029}$	$30.16_{0.26}$	$0.963_{0.071}$

Table SI.LVIII: GCMC-MBAR results for 2,2,3-trimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
550	$424.5_{1.7}$	$94.6_{5.8}$	$2.204_{0.085}$	$14.90_{0.26}$	$0.582_{0.014}$
540	$446.63_{0.90}$	$77.5_{4.4}$	$1.931_{0.070}$	$17.04_{0.27}$	$0.634_{0.015}$
530	$466.95_{0.61}$	$64.6_{2.9}$	$1.685_{0.058}$	$18.86_{0.22}$	$0.676_{0.010}$
520	$485.81_{0.76}$	$54.4_{1.9}$	$1.463_{0.050}$	$20.45_{0.13}$	$0.71_{0.01}$
510	$502.62_{0.63}$	$45.9_{1.6}$	$1.263_{0.044}$	$21.840_{0.082}$	$0.74_{0.01}$
500	$517.03_{0.59}$	$38.7_{1.7}$	$1.084_{0.037}$	$23.05_{0.11}$	$0.77_{0.01}$
490	$529.98_{0.76}$	$32.6_{1.9}$	$0.925_{0.029}$	$24.14_{0.18}$	$0.795_{0.024}$
480	$542.7_{1.0}$	$27.4_{1.8}$	$0.784_{0.021}$	$25.18_{0.25}$	$0.818_{0.037}$
470	$555.2_{1.2}$	$23.0_{1.6}$	$0.659_{0.013}$	$26.17_{0.29}$	$0.840_{0.045}$
460	$566.7_{1.3}$	$19.1_{1.2}$	$0.5500_{0.0068}$	$27.09_{0.30}$	$0.860_{0.047}$
450	$577.6_{1.2}$	$15.82_{0.81}$	$0.4550_{0.0027}$	$27.94_{0.29}$	$0.878_{0.045}$
440	$588.5_{3.8}$	$13.0_{1.5}$	$0.4_{1.3}$	$28.8_{7.3}$	$0.9_{2.0}$
430	$599.6_{2.4}$	$10.61_{0.31}$	$0.3023_{0.0035}$	$29.58_{0.30}$	$0.910_{0.038}$
420	$610.1_{2.1}$	$8.58_{0.19}$	$0.2423_{0.0044}$	$30.34_{0.26}$	$0.923_{0.036}$
410	$619.5_{1.4}$	$6.88_{0.12}$	$0.1918_{0.0049}$	$31.02_{0.22}$	$0.935_{0.038}$
400	$628.4_{1.5}$	$5.447_{0.098}$	$0.1498_{0.0051}$	$31.66_{0.24}$	$0.945_{0.044}$
380	$645.33_{0.68}$	$3.285_{0.097}$	$0.0874_{0.0051}$	$32.86_{0.25}$	$0.961_{0.067}$

Table SI.LIX: GCMC-MBAR results for 2,2,4-trimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}\left(\mathbf{K}\right)$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
530	400.9 _{2.2}	97.9 _{4.8}	$2.120_{0.038}$	$13.58_{0.32}$	$0.561_{0.019}$
520	$427.5_{1.3}$	$79.4_{3.2}$	$1.844_{0.028}$	$15.89_{0.27}$	$0.614_{0.018}$
510	$448.78_{0.77}$	$65.1_{1.8}$	$1.597_{0.021}$	$17.79_{0.18}$	$0.661_{0.012}$
500	$466.83_{0.75}$	$53.9_{1.1}$	$1.378_{0.018}$	$19.40_{0.13}$	$0.70_{0.01}$
490	$483.78_{0.81}$	$44.87_{0.89}$	$1.182_{0.015}$	$20.85_{0.12}$	$0.74_{0.01}$
480	$499.84_{0.70}$	$37.41_{0.89}$	$1.009_{0.012}$	$22.18_{0.12}$	$0.772_{0.012}$
470	$514.41_{0.57}$	$31.21_{0.86}$	$0.8545_{0.0083}$	$23.38_{0.13}$	$0.800_{0.016}$
460	$527.62_{0.53}$	$26.00_{0.75}$	$0.7187_{0.0053}$	$24.45_{0.13}$	$0.826_{0.020}$
450	$539.75_{0.68}$	$21.58_{0.59}$	$0.5995_{0.0030}$	$25.42_{0.12}$	$0.848_{0.022}$
440	$551.12_{0.66}$	$17.82_{0.43}$	$0.4958_{0.0022}$	$26.32_{0.12}$	$0.869_{0.022}$
430	$562.67_{0.50}$	$14.62_{0.31}$	$0.4062_{0.0027}$	$27.21_{0.13}$	$0.888_{0.023}$
420	$574.59_{0.49}$	$11.89_{0.25}$	$0.3292_{0.0035}$	$28.09_{0.13}$	$0.906_{0.027}$
410	$585.04_{0.41}$	$9.58_{0.25}$	$0.2637_{0.0042}$	$28.86_{0.15}$	$0.922_{0.037}$
400	$594.23_{0.30}$	$7.66_{0.27}$	$0.2087_{0.0051}$	$29.54_{0.20}$	$0.936_{0.055}$
390	$604.33_{0.26}$	$6.05_{0.29}$	$0.1630_{0.0063}$	$30.26_{0.28}$	$0.949_{0.082}$
380	$615.43_{0.23}$	$4.72_{0.30}$	$0.1252_{0.0077}$	$31.01_{0.41}$	$0.96_{0.12}$
370	$625.27_{0.23}$	$3.63_{0.29}$	$0.0946_{0.0092}$	$31.68_{0.58}$	$0.97_{0.18}$

Table SI.LX: GCMC-MBAR results for 2,3,3-trimethylpentane with the MiPPE-SL force field. Subscripts correspond to the 95% confidence interval computed with bootstrap resampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
560	$426.5_{9.5}$	$98.7_{1.6}$	$2.304_{0.024}$	$14.76_{0.51}$	$0.57_{0.01}$
550	$452.4_{5.9}$	$80.7_{1.4}$	$2.023_{0.020}$	$17.10_{0.29}$	$0.63_{0.01}$
540	$473.3_{2.0}$	$67.2_{1.1}$	$1.769_{0.018}$	$18.98_{0.12}$	$0.67_{0.01}$
530	$490.60_{0.59}$	$56.53_{0.74}$	$1.540_{0.016}$	$20.523_{0.065}$	$0.71_{0.01}$
520	$505.89_{0.92}$	$47.80_{0.63}$	$1.335_{0.014}$	$21.863_{0.043}$	$0.74_{0.01}$
510	$520.2_{1.1}$	$40.46_{0.65}$	$1.151_{0.012}$	$23.086_{0.038}$	$0.77_{0.01}$
500	$534.1_{1.0}$	$34.21_{0.66}$	$0.9866_{0.0088}$	$24.232_{0.057}$	$0.79_{0.01}$
490	$546.97_{0.93}$	$28.87_{0.60}$	$0.8402_{0.0063}$	$25.283_{0.071}$	$0.816_{0.012}$
480	$558.82_{0.80}$	$24.29_{0.47}$	$0.7108_{0.0044}$	$26.241_{0.071}$	$0.837_{0.013}$
470	$570.25_{0.87}$	$20.36_{0.30}$	$0.5967_{0.0036}$	$27.143_{0.070}$	$0.857_{0.013}$
460	$581.5_{1.1}$	$16.96_{0.17}$	$0.4969_{0.0037}$	$28.007_{0.066}$	$0.875_{0.013}$
450	$592.0_{1.0}$	$14.04_{0.13}$	$0.4101_{0.0041}$	$28.816_{0.071}$	$0.892_{0.016}$
440	$602.0_{1.1}$	$11.53_{0.15}$	$0.3353_{0.0046}$	$29.578_{0.085}$	$0.908_{0.022}$
430	$612.3_{1.6}$	$9.39_{0.16}$	$0.2714_{0.0052}$	$30.34_{0.10}$	$0.924_{0.031}$
420	$622.8_{1.6}$	$7.58_{0.16}$	$0.2171_{0.0058}$	$31.10_{0.13}$	$0.937_{0.042}$
410	$632.87_{0.75}$	$6.05_{0.16}$	$0.1716_{0.0064}$	$31.82_{0.20}$	$0.950_{0.057}$
400	$642.39_{0.29}$	$4.78_{0.17}$	$0.1338_{0.0070}$	$32.50_{0.28}$	$0.961_{0.077}$
390	$651.59_{0.31}$	$3.73_{0.18}$	$0.1028_{0.0075}$	$33.14_{0.37}$	$0.97_{0.11}$

SI.VI.2 Alkynes

Table SI.LXI: GCMC-MBAR results for ethyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
290	$421.3_{3.5}$	$69.7_{3.3}$	$4.083_{0.020}$	$8.80_{0.21}$	$0.632_{0.033}$
280	$450.7_{1.9}$	$50.69_{0.91}$	$3.196_{0.026}$	$10.27_{0.12}$	$0.705_{0.017}$
270	$474.9_{3.0}$	$37.66_{0.39}$	$2.465_{0.024}$	$11.390_{0.086}$	$0.76_{0.01}$
260	$497.3_{1.1}$	$27.95_{0.53}$	$1.865_{0.017}$	$12.361_{0.034}$	$0.804_{0.011}$
250	$517.91_{0.59}$	$20.53_{0.34}$	$1.378_{0.012}$	$13.214_{0.042}$	$0.841_{0.013}$
240	$536.87_{0.67}$	$14.82_{0.12}$	$0.992_{0.010}$	$13.975_{0.028}$	$0.87_{0.01}$
230	$554.12_{0.54}$	$10.447_{0.095}$	$0.6925_{0.0089}$	$14.649_{0.037}$	$0.903_{0.012}$
220	$570.90_{0.53}$	$7.155_{0.097}$	$0.4667_{0.0082}$	$15.283_{0.049}$	$0.929_{0.021}$

Table SI.LXII: GCMC-MBAR results for propyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
380	$441.0_{7.9}$	$82.2_{3.1}$	$3.839_{0.090}$	$10.96_{0.34}$	$0.59_{0.01}$
370	$472.7_{5.0}$	$62.7_{2.7}$	$3.164_{0.073}$	$12.83_{0.31}$	$0.657_{0.015}$
360	$498.3_{2.8}$	$48.6_{1.9}$	$2.584_{0.057}$	$14.34_{0.24}$	$0.711_{0.015}$
350	$520.1_{2.4}$	$38.1_{1.3}$	$2.088_{0.045}$	$15.58_{0.19}$	$0.754_{0.011}$
340	$539.4_{2.1}$	$29.88_{0.90}$	$1.667_{0.035}$	$16.65_{0.15}$	$0.79_{0.01}$
330	$556.6_{1.4}$	$23.31_{0.72}$	$1.313_{0.027}$	$17.58_{0.11}$	$0.822_{0.010}$
320	$572.1_{1.3}$	$18.02_{0.60}$	$1.018_{0.019}$	$18.41_{0.11}$	$0.850_{0.014}$
310	$587.6_{1.3}$	$13.76_{0.47}$	$0.776_{0.012}$	$19.20_{0.12}$	$0.876_{0.017}$
300	$603.3_{1.5}$	$10.35_{0.32}$	$0.5799_{0.0068}$	$19.97_{0.13}$	$0.900_{0.019}$
290	$617.9_{2.2}$	$7.65_{0.20}$	$0.4240_{0.0040}$	$20.67_{0.15}$	$0.921_{0.020}$

Table SI.LXIII: GCMC-MBAR results for 1-butyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
410	$445.1_{5.3}$	76_{12}	$2.95_{0.10}$	$12.82_{0.72}$	$0.620_{0.056}$
400	$472.3_{3.4}$	$59.2_{6.8}$	$2.464_{0.050}$	$14.61_{0.60}$	$0.677_{0.054}$
390	$495.0_{2.6}$	$47.1_{2.3}$	$2.038_{0.030}$	$16.08_{0.32}$	$0.722_{0.027}$
380	$515.5_{2.0}$	$37.54_{0.86}$	$1.669_{0.023}$	$17.37_{0.16}$	$0.76_{0.01}$
370	$534.5_{1.3}$	$29.90_{0.70}$	$1.351_{0.018}$	$18.53_{0.13}$	$0.79_{0.01}$
360	$552.0_{1.6}$	$23.69_{0.62}$	$1.081_{0.013}$	$19.57_{0.16}$	$0.824_{0.013}$
350	$567.8_{1.9}$	$18.62_{0.51}$	$0.8524_{0.0086}$	$20.50_{0.18}$	$0.851_{0.017}$
340	$582.2_{1.1}$	$14.49_{0.38}$	$0.6623_{0.0060}$	$21.34_{0.13}$	$0.874_{0.019}$
330	$595.8_{1.1}$	$11.15_{0.24}$	$0.5061_{0.0049}$	$22.112_{0.082}$	$0.895_{0.019}$
320	$610.6_{2.8}$	$8.45_{0.14}$	$0.3795_{0.0049}$	$22.905_{0.095}$	$0.913_{0.019}$
310	$624.6_{1.8}$	$6.284_{0.087}$	$0.2782_{0.0051}$	$23.650_{0.059}$	$0.929_{0.022}$

Table SI.LXIV: GCMC-MBAR results for 2-butyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$\rho_{\mathrm{vap}}^{\mathrm{sat}}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
450	$431.8_{7.4}$	$93.5_{4.3}$	$3.624_{0.050}$	$11.93_{0.26}$	$0.561_{0.018}$
440	$466.0_{5.5}$	$74.1_{3.5}$	$3.073_{0.028}$	$14.00_{0.24}$	$0.613_{0.024}$
430	$491.6_{2.0}$	$59.1_{1.9}$	$2.587_{0.016}$	$15.71_{0.17}$	$0.662_{0.020}$
420	$512.10_{0.77}$	$47.58_{0.69}$	$2.163_{0.014}$	$17.114_{0.094}$	$0.704_{0.010}$
410	$530.5_{1.1}$	$38.42_{0.30}$	$1.795_{0.013}$	$18.346_{0.054}$	$0.74_{0.01}$
400	$548.28_{0.94}$	$30.99_{0.31}$	$1.476_{0.011}$	$19.487_{0.051}$	$0.77_{0.01}$
390	$565.05_{0.94}$	$24.89_{0.27}$	$1.2013_{0.0089}$	$20.533_{0.061}$	$0.81_{0.01}$
380	$580.6_{1.1}$	$19.89_{0.25}$	$0.9674_{0.0068}$	$21.483_{0.073}$	$0.83_{0.01}$
370	$594.97_{0.83}$	$15.78_{0.25}$	$0.7697_{0.0050}$	$22.349_{0.074}$	$0.858_{0.011}$
360	$608.65_{0.59}$	$12.41_{0.22}$	$0.6041_{0.0038}$	$23.157_{0.070}$	$0.880_{0.017}$
350	$621.69_{0.66}$	$9.66_{0.19}$	$0.4672_{0.0041}$	$23.912_{0.091}$	$0.899_{0.024}$
340	$633.85_{0.59}$	$7.42_{0.16}$	$0.3552_{0.0052}$	$24.60_{0.12}$	$0.916_{0.033}$
330	$646.13_{0.73}$	$5.62_{0.13}$	$0.2651_{0.0064}$	$25.26_{0.16}$	$0.930_{0.044}$

Table SI.LXV: GCMC-MBAR results for 1-pentyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
450	$433.7_{2.3}$	$85.6_{2.5}$	$2.732_{0.022}$	$13.15_{0.18}$	$0.581_{0.014}$
440	$461.7_{1.7}$	$66.9_{2.0}$	$2.304_{0.015}$	$15.29_{0.18}$	$0.641_{0.018}$
430	$485.2_{1.5}$	$53.1_{1.2}$	$1.932_{0.012}$	$17.06_{0.16}$	$0.693_{0.016}$
420	$505.4_{1.6}$	$42.65_{0.51}$	$1.608_{0.012}$	$18.54_{0.11}$	$0.735_{0.011}$
410	$523.3_{1.4}$	$34.37_{0.30}$	$1.327_{0.010}$	$19.810_{0.078}$	$0.77_{0.01}$
400	$539.7_{1.2}$	$27.67_{0.31}$	$1.0844_{0.0077}$	$20.949_{0.055}$	$0.80_{0.01}$
390	$555.3_{1.1}$	$22.17_{0.30}$	$0.8770_{0.0052}$	$21.998_{0.041}$	$0.83_{0.01}$
380	$570.2_{1.0}$	$17.64_{0.27}$	$0.7009_{0.0033}$	$22.977_{0.033}$	$0.857_{0.012}$
370	$584.03_{0.90}$	$13.91_{0.25}$	$0.5528_{0.0028}$	$23.874_{0.050}$	$0.880_{0.018}$
360	$596.86_{0.83}$	$10.86_{0.22}$	$0.4300_{0.0040}$	$24.697_{0.083}$	$0.901_{0.026}$
350	$609.71_{0.69}$	$8.37_{0.19}$	$0.3291_{0.0055}$	$25.49_{0.11}$	$0.920_{0.037}$
340	$622.58_{0.99}$	$6.37_{0.15}$	$0.2474_{0.0070}$	$26.27_{0.13}$	$0.936_{0.050}$

Table SI.LXVI: GCMC-MBAR results for 2-pentyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
470	$445.8_{3.1}$	$83.4_{6.2}$	$2.77_{0.12}$	$14.09_{0.22}$	$0.578_{0.020}$
460	$473.8_{1.9}$	$65.6_{5.4}$	$2.343_{0.088}$	$16.27_{0.30}$	$0.636_{0.028}$
450	$496.8_{1.0}$	$52.5_{3.7}$	$1.973_{0.064}$	$18.06_{0.28}$	$0.685_{0.026}$
440	$516.64_{0.73}$	$42.4_{2.3}$	$1.650_{0.048}$	$19.56_{0.22}$	$0.725_{0.020}$
430	$534.83_{0.74}$	$34.3_{1.7}$	$1.369_{0.036}$	$20.89_{0.19}$	$0.761_{0.017}$
420	$551.57_{0.81}$	$27.7_{1.4}$	$1.126_{0.025}$	$22.08_{0.19}$	$0.792_{0.022}$
410	$566.89_{0.86}$	$22.3_{1.2}$	$0.917_{0.016}$	$23.16_{0.22}$	$0.821_{0.030}$
400	$581.3_{1.1}$	$17.9_{1.0}$	$0.7394_{0.0080}$	$24.15_{0.26}$	$0.847_{0.038}$
390	$595.3_{1.6}$	$14.24_{0.71}$	$0.5890_{0.0039}$	$25.09_{0.28}$	$0.869_{0.040}$
380	$608.6_{1.7}$	$11.23_{0.40}$	$0.4631_{0.0052}$	$25.95_{0.25}$	$0.889_{0.038}$
370	$621.0_{1.3}$	$8.76_{0.19}$	$0.3587_{0.0066}$	$26.75_{0.19}$	$0.907_{0.032}$
360	$632.99_{0.96}$	$6.75_{0.16}$	$0.2735_{0.0069}$	$27.52_{0.12}$	$0.922_{0.031}$
350	$644.7_{1.0}$	$5.12_{0.20}$	$0.2049_{0.0065}$	$28.249_{0.10}$	$0.937_{0.044}$

Table SI.LXVII: GCMC-MBAR results for 1-hexyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
490	$423.3_{3.5}$	$87.3_{2.1}$	$2.538_{0.068}$	$14.25_{0.14}$	$0.586_{0.010}$
480	$453.3_{2.0}$	$69.9_{1.5}$	$2.170_{0.060}$	$16.54_{0.12}$	$0.639_{0.015}$
470	$477.4_{1.3}$	$56.51_{0.96}$	$1.844_{0.056}$	$18.448_{0.089}$	$0.686_{0.016}$
460	$497.3_{1.4}$	$46.06_{0.95}$	$1.556_{0.051}$	$20.041_{0.074}$	$0.726_{0.011}$
450	$514.8_{1.3}$	$37.6_{1.3}$	$1.304_{0.044}$	$21.428_{0.10}$	$0.76_{0.01}$
440	$531.5_{1.2}$	$30.7_{1.7}$	$1.084_{0.035}$	$22.71_{0.16}$	$0.792_{0.017}$
430	$547.4_{1.2}$	$25.0_{1.7}$	$0.893_{0.024}$	$23.91_{0.22}$	$0.820_{0.032}$
420	$561.77_{0.87}$	$20.3_{1.5}$	$0.728_{0.013}$	$24.98_{0.23}$	$0.845_{0.042}$
410	$575.17_{0.62}$	$16.34_{0.99}$	$0.5875_{0.0050}$	$25.97_{0.20}$	$0.867_{0.043}$
400	$588.08_{0.75}$	$13.04_{0.55}$	$0.4682_{0.0027}$	$26.90_{0.15}$	$0.887_{0.038}$
390	$599.80_{0.63}$	$10.31_{0.28}$	$0.3684_{0.0047}$	$27.75_{0.11}$	$0.905_{0.033}$
380	$610.32_{0.79}$	$8.06_{0.17}$	$0.2858_{0.0061}$	$28.507_{0.080}$	$0.922_{0.036}$
370	$621.3_{2.9}$	$6.23_{0.13}$	$0.2185_{0.0072}$	$29.267_{0.086}$	$0.937_{0.048}$
360	$634.3_{3.1}$	$4.74_{0.11}$	$0.1640_{0.0083}$	$30.118_{0.078}$	$0.950_{0.067}$

Table SI.LXVIII: GCMC-MBAR results for 2-hexyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{ m vap}^{ m sat}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
500	$438.2_{2.9}$	$85.9_{4.0}$	$2.495_{0.026}$	$14.93_{0.36}$	$0.574_{0.024}$
490	$464.9_{1.8}$	$67.7_{2.5}$	$2.131_{0.016}$	$17.31_{0.29}$	$0.635_{0.021}$
480	$486.7_{1.5}$	$54.7_{1.3}$	$1.811_{0.011}$	$19.18_{0.19}$	$0.682_{0.016}$
470	$505.2_{1.7}$	$44.67_{0.74}$	$1.5302_{0.0087}$	$20.71_{0.14}$	$0.720_{0.012}$
460	$522.5_{2.3}$	$36.61_{0.52}$	$1.2839_{0.0074}$	$22.09_{0.14}$	$0.753_{0.011}$
450	$539.3_{2.5}$	$29.96_{0.40}$	$1.0686_{0.0062}$	$23.39_{0.15}$	$0.783_{0.010}$
440	$555.2_{1.8}$	$24.43_{0.31}$	$0.8815_{0.0050}$	$24.60_{0.11}$	$0.81_{0.01}$
430	$569.62_{0.68}$	$19.82_{0.28}$	$0.7201_{0.0040}$	$25.678_{0.066}$	$0.835_{0.010}$
420	$583.3_{1.4}$	$15.98_{0.26}$	$0.5821_{0.0034}$	$26.69_{0.11}$	$0.857_{0.014}$
410	$597.0_{1.4}$	$12.77_{0.22}$	$0.4651_{0.0035}$	$27.67_{0.12}$	$0.877_{0.017}$
400	$609.9_{1.1}$	$10.11_{0.16}$	$0.3669_{0.0039}$	$28.59_{0.11}$	$0.896_{0.020}$
390	$621.7_{1.4}$	$7.92_{0.14}$	$0.2857_{0.0045}$	$29.42_{0.14}$	$0.914_{0.026}$
380	$633.2_{1.2}$	$6.12_{0.16}$	$0.2190_{0.0050}$	$30.23_{0.16}$	$0.930_{0.040}$
370	$644.01_{0.40}$	$4.67_{0.18}$	$0.1653_{0.0059}$	$30.98_{0.19}$	$0.944_{0.063}$

Table SI.LXIX: GCMC-MBAR results for 1-heptyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
520	$427.5_{6.0}$	$82.9_{6.4}$	$2.218_{0.071}$	$16.05_{0.64}$	$0.595_{0.026}$
510	$454.3_{2.5}$	$66.9_{4.8}$	$1.906_{0.055}$	$18.37_{0.49}$	$0.646_{0.027}$
500	$476.5_{1.4}$	$54.5_{3.3}$	$1.629_{0.041}$	$20.34_{0.35}$	$0.691_{0.024}$
490	$495.7_{1.4}$	$44.8_{2.4}$	$1.384_{0.030}$	$22.02_{0.25}$	$0.730_{0.023}$
480	$512.9_{1.6}$	$36.9_{1.8}$	$1.168_{0.020}$	$23.51_{0.18}$	$0.764_{0.024}$
470	$528.7_{2.0}$	$30.3_{1.4}$	$0.979_{0.012}$	$24.85_{0.13}$	$0.794_{0.026}$
460	$543.2_{1.9}$	$24.91_{0.95}$	$0.8138_{0.0072}$	$26.066_{0.092}$	$0.821_{0.027}$
450	$556.9_{1.5}$	$20.37_{0.57}$	$0.6705_{0.0058}$	$27.199_{0.070}$	$0.846_{0.025}$
440	$570.2_{1.0}$	$16.57_{0.25}$	$0.5472_{0.0063}$	$28.279_{0.062}$	$0.868_{0.019}$
430	$583.08_{0.58}$	$13.38_{0.11}$	$0.4419_{0.0062}$	$29.301_{0.062}$	$0.888_{0.012}$
420	$595.19_{0.32}$	$10.71_{0.23}$	$0.3526_{0.0055}$	$30.253_{0.097}$	$0.907_{0.017}$
410	$606.63_{0.36}$	$8.50_{0.30}$	$0.2779_{0.0046}$	$31.14_{0.16}$	$0.923_{0.035}$
400	$617.58_{0.39}$	$6.66_{0.33}$	$0.2160_{0.0048}$	$31.98_{0.23}$	$0.938_{0.058}$
390	$628.63_{0.36}$	$5.16_{0.31}$	$0.1654_{0.0061}$	$32.81_{0.31}$	$0.950_{0.088}$

Table SI.LXX: GCMC-MBAR results for 1-octyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

T ^{sat} (K)	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
550	$419.4_{3.6}$	$89.5_{1.2}$	$2.075_{0.015}$	$16.27_{0.24}$	$0.56_{0.01}$
540	$445.2_{3.0}$	$70.95_{0.88}$	$1.788_{0.015}$	$18.98_{0.20}$	$0.62_{0.01}$
530	$467.6_{2.6}$	$57.19_{0.55}$	$1.535_{0.015}$	$21.30_{0.17}$	$0.67_{0.01}$
520	$487.1_{2.4}$	$46.85_{0.42}$	$1.311_{0.014}$	$23.22_{0.15}$	$0.71_{0.01}$
510	$504.3_{2.0}$	$38.69_{0.48}$	$1.114_{0.012}$	$24.87_{0.13}$	$0.75_{0.01}$
500	$519.8_{1.5}$	$32.01_{0.56}$	$0.9395_{0.0099}$	$26.33_{0.12}$	$0.78_{0.01}$
490	$534.3_{1.2}$	$26.45_{0.57}$	$0.7869_{0.0075}$	$27.66_{0.13}$	$0.805_{0.010}$
480	$548.2_{1.2}$	$21.77_{0.52}$	$0.6539_{0.0053}$	$28.92_{0.15}$	$0.829_{0.014}$
470	$561.6_{1.2}$	$17.83_{0.40}$	$0.5385_{0.0038}$	$30.10_{0.15}$	$0.852_{0.016}$
460	$574.15_{0.95}$	$14.52_{0.26}$	$0.4392_{0.0033}$	$31.21_{0.13}$	$0.872_{0.016}$
450	$586.02_{0.57}$	$11.73_{0.12}$	$0.3546_{0.0033}$	$32.245_{0.084}$	$0.890_{0.012}$
440	$597.51_{0.35}$	$9.404_{0.077}$	$0.2830_{0.0032}$	$33.229_{0.046}$	$0.91_{0.01}$
430	$608.64_{0.40}$	$7.46_{0.15}$	$0.2231_{0.0029}$	$34.171_{0.089}$	$0.921_{0.019}$
420	$619.31_{0.42}$	$5.86_{0.20}$	$0.1736_{0.0027}$	$35.06_{0.16}$	$0.935_{0.039}$
410	$630.00_{0.38}$	$4.55_{0.22}$	$0.1330_{0.0032}$	$35.94_{0.24}$	$0.946_{0.067}$

Table SI.LXXI: GCMC-MBAR results for 1-nonyne with the MiPPE force field. Subscripts correspond to the 95% confidence interval computed with bootstrap re-sampling.

$T^{\mathrm{sat}}(\mathbf{K})$	$ ho_{ m liq}^{ m sat}$ (kg/m ³)	$ ho_{ m vap}^{ m sat}$ (kg/m ³)	$P_{\mathrm{vap}}^{\mathrm{sat}}$ (MPa)	$\Delta H_{\rm v}$ (kJ/mol)	$Z_{ m vap}^{ m sat}$
570	$427.1_{1.2}$	$80.5_{1.0}$	$1.776_{0.015}$	$18.58_{0.17}$	$0.58_{0.01}$
560	$450.9_{1.3}$	$64.43_{0.86}$	$1.531_{0.013}$	$21.28_{0.20}$	$0.63_{0.01}$
550	$471.7_{1.6}$	$52.31_{0.73}$	$1.315_{0.010}$	$23.58_{0.20}$	$0.68_{0.01}$
540	$489.9_{1.5}$	$43.02_{0.62}$	$1.1239_{0.0080}$	$25.52_{0.17}$	$0.72_{0.01}$
530	$506.3_{1.1}$	$35.61_{0.52}$	$0.9552_{0.0059}$	$27.21_{0.13}$	$0.76_{0.01}$
520	$521.47_{0.66}$	$29.51_{0.42}$	$0.8067_{0.0041}$	$28.739_{0.093}$	$0.79_{0.01}$
510	$535.75_{0.67}$	$24.42_{0.33}$	$0.6762_{0.0026}$	$30.149_{0.083}$	$0.81_{0.01}$
500	$549.10_{0.85}$	$20.14_{0.26}$	$0.5626_{0.0015}$	$31.452_{0.094}$	$0.83_{0.01}$
490	$561.65_{0.81}$	$16.54_{0.20}$	$0.46410_{0.00097}$	$32.666_{0.094}$	$0.856_{0.010}$
480	$573.68_{0.62}$	$13.50_{0.15}$	$0.3792_{0.0010}$	$33.811_{0.080}$	$0.875_{0.011}$
470	$585.26_{0.49}$	$10.94_{0.11}$	$0.3068_{0.0014}$	$34.896_{0.067}$	$0.892_{0.012}$
460	$596.35_{0.58}$	$8.794_{0.096}$	$0.2455_{0.0017}$	$35.924_{0.069}$	$0.907_{0.015}$
450	$607.27_{0.71}$	$7.003_{0.091}$	$0.1942_{0.0020}$	$36.924_{0.081}$	$0.921_{0.019}$
440	$618.21_{0.61}$	$5.516_{0.091}$	$0.1516_{0.0023}$	$37.91_{0.10}$	$0.933_{0.027}$
430	$628.74_{0.51}$	$4.292_{0.090}$	$0.1166_{0.0026}$	$38.85_{0.14}$	$0.944_{0.037}$
420	$638.50_{0.50}$	$3.296_{0.086}$	$0.0885_{0.0030}$	$39.72_{0.18}$	$0.955_{0.052}$

SI.VII Simulation state points

SI.VII.1 Cyclohexane

Table SI.LXXII: State points simulated for cyclohexane with the MiPPE force field (second iteration, $\theta^{\langle 2 \rangle}$ $\lambda_{\rm CH_2}=16$).

T(K)	μ (K)	L (nm)
450	-4367	3.0
500	-4367	3.0
550	-4367	3.0
500	-4149	3.0
460	-4024	3.0
410	-3893	3.0
360	-3792	3.0

Table SI.LXXIII: State points simulated for cyclohexane with the TraPPE force field (zeroth iteration, $\theta^{(0)}$).

T (K)	μ (K)	L (nm)
450	-4350	3.0
500	-4350	3.0
550	-4350	3.0
500	-4120	3.0
460	-3977	3.0
410	-3790	3.0
350	-3562	3.0

Table SI.LXXIV: State points simulated for cyclohexane with the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\text{CH}_2} = 14$ force field.

T (K)	μ (K)	L (nm)
450	-4389	3.0
500	-4389	3.0
550	-4389	3.0
500	-4164	3.0
460	-4033	3.0
410	-3891	3.0
360	-3780	3.0

Table SI.LXXV: State points simulated for cyclohexane with the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{CH_2} = 16$ force field.

T (K)	μ (K)	L (nm)
450	-4367	3.0
500	-4367	3.0
550	-4367	3.0
500	-4149	3.0
460	-4024	3.0
410	-3893	3.0
360	-3792	3.0

Table SI.LXXVI: State points simulated for cyclohexane with the first iteration $(\theta^{(1)})$ $\lambda_{\text{CH}_2} = 18$ force field.

T (K)	μ (K)	L (nm)
450	-4370	3.0
500	-4370	3.0
550	-4370	3.0
500	-4158	3.0
460	-4037	3.0
410	-3912	3.0
360	-3825	3.0

Table SI.LXXVII: State points simulated for cyclohexane with the first iteration $(\theta^{\langle 1 \rangle})$ $\lambda_{\rm CH_2}=20$ force field.

T(K)	μ (K)	L (nm)
450	-4386	3.0
500	-4386	3.0
550	-4386	3.0
500	-4178	3.0
460	-4062	3.0
410	-3946	3.0
360	-3866	3.0

SI.VII.2 Branched alkanes

 $Table \ SI.LXXVIII: State \ points \ simulated \ for \ 2-methyl propone \ with \ the \ TraPPE \ force \ field.$

T(K)	μ (K)	L (nm)
350	-3120	3.0
380	-3120	3.0
405	-3117	3.0
380	-2980	3.0
350	-2880	3.0
320	-2790	3.0
290	-2705	3.0
260	-2645	3.0
230	-2600	3.0
200	-2570	3.0

Table SI.LXXIX: State points simulated for 2,2-dimethylpropane with the TraPPE force field.

T (K)	μ (K)	L (nm)
380	-3405	3.0
410	-3405	3.0
440	-3405	3.0
410	-3250	3.0
380	-3140	3.0
350	-3037	3.0
330	-2970	3.0
300	-2900	3.0
270	-2820	3.0

Table SI.LXXX: State points simulated for 2,2-dimethylbutane with the TraPPE force field.

T (K)	μ (K)	L (nm)
420	-3860	3.5
450	-3860	3.5
480	-3860	3.5
450	-3719	3.5
420	-3600	3.5
400	-3524	3.5
380	-3450	3.5
360	-3368	3.5
340	-3288	3.5
310	-3280	3.5

 $Table \ SI.LXXXI: State \ points \ simulated \ for \ 2,3-dimethylbutane \ with \ the \ TraPPE \ force \ field.$

T (K)	μ (K)	L (nm)
440	-4015	3.0
470	-4015	3.0
500	-4011	3.0
470	-3845	3.0
440	-3735	3.0
410	-3635	3.0
380	-3555	3.0
350	-3480	3.0
320	-3415	3.0

Table SI.LXXXII: State points simulated for 3,3-dimethylhexane with the TraPPE force field.

и (K)	L (nm)
, , ,	` /
- 4670	3.5
-4670	3.5
-4670	3.5
-4476	3.5
-4370	3.5
-4268	3.5
-4164	3.5
-4039	3.5
-3925	3.5
	-4670 -4476 -4370 -4268 -4164 -4039

Table SI.LXXXIII: State points simulated for 3-methyl-3-ethylpentane with the TraPPE force field.

T (K)	μ (K)	L (nm)
500	-4785	4.0
550	-4785	4.0
580	-4785	4.0
550	-4636	4.0
520	-4520	4.0
490	-4400	4.0
460	-4280	4.0
430	-4160	4.0
410	-4080	4.0
390	-3990	4.0

Table SI.LXXXIV: State points simulated for 2,3,4-trimethylpentane with the TraPPE force field.

T(K)	μ (K)	L (nm)
480	-4740	3.5
520	-4740	3.5
565	-4735	3.5
530	-4549	3.5
500	-4436	3.5
470	-4337	3.5
440	-4241	3.5
410	-4182	3.5
380	-4090	3.5
350	-4020	3.5

Table SI.LXXXV: State points simulated for 2,2,4-trimethylpentane with the TraPPE force field.

T(K)	μ (K)	L (nm)
480	-4600	4.0
530	-4600	4.0
560	-4600	4.0
530	-4450	4.0
500	-4330	4.0
470	-4210	4.0
440	-4090	4.0
410	-3960	4.0
380	-3840	4.0

Table SI.LXXXVI: State points simulated for 2-methylpropane with the MiPPE-gen force field.

T(K)	μ (K)	L (nm)
350	-3150	3.0
380	-3150	3.0
410	-3145	3.0
380	-3010	3.0
350	-2910	3.0
320	-2830	3.0
290	-2760	3.0
260	-2700	3.0
230	-2670	3.0
200	-2640	3.0

Table SI.LXXXVII: State points simulated for 2,2-dimethylpropane with the MiPPE-gen force field.

T(K)	μ (K)	L (nm)
368	-3344	3.0
398	-3344	3.0
430	-3400	3.0
398	-3216	3.0
372	-3124	3.0
346	-3032	3.0
326	-2961	3.0
299	-2865	3.0
270	-2759	3.0

Table SI.LXXXVIII: State points simulated for 2,2-dimethylbutane with the MiPPE-gen force field.

T(K)	μ (K)	L (nm)
415	-3873	3.5
445	-3873	3.5
480	-3895	3.5
450	-3756	3.5
420	-3654	3.5
400	-3588	3.5
380	-3521	3.5
360	-3454	3.5
340	-3384	3.5
310	-3380	3.5

Table SI.LXXXIX: State points simulated for 2,3-dimethylbutane with the MiPPE-gen force field.

T (K)	μ (K)	L (nm)
440	-4010	3.0
470	-4010	3.0
500	-4009	3.0
470	-3860	3.0
440	-3760	3.0
410	-3670	3.0
380	-3600	3.0
350	-3530	3.0
320	-3480	3.0

Table SI.XC: State points simulated for 2,3,4-trimethylpentane with the MiPPE-gen force field.

T (K)	μ (K)	L (nm)
480	-4720	3.5
520	-4720	3.5
565	-4713	3.5
530	-4540	3.5
500	-4360	3.5
470	-4355	3.5
440	-4275	3.5
410	-4205	3.5
380	-4165	3.5
350	-4115	3.5

Table SI.XCI: State points simulated for 2,2,4-trimethylpentane with the MiPPE-gen force field.

T (K)	μ (K)	L (nm)
470	-4570	4.0
520	-4570	4.0
550	-4570	4.0
520	-4420	4.0
490	-4300	4.0
460	-4170	4.0
430	-4050	4.0
400	-3920	4.0
370	-3790	4.0