SUPPORTING INFORMATION

Sodium Diisopropylamide in Tetrahydrofuran: Selectivities, Rates, and Mechanisms of Alkene Isomerizations and Diene Metalations

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I. Rate Studies



Figure S-1. Representative plot of pentene concentration versus time (¹H NMR) for the isomerization of 0.76 M 1-pentene with 0.18 M NaDA and 0.59 M diisopropylamine in 8.83 M THF/hexane at 25 °C. Each interval represents 300 seconds.



Figure S-2. Plot of alkene concentration versus time (¹H NMR) for the isomerization of 0.76 M 1-pentene with 0.18 M NaDA and 0.59 M diisopropylamine in 8.83 M THF/hexane at 25 °C. The black trace represents 1-pentene, the blue trace represents *trans*-2-pentene, and the red trace represents *cis*-2-pentene.



Figure S-3. Plot of initial rate versus THF concentration for the isomerization of 0.76 M 1-pentene with 0.18 M NaDA and 0.59 M diisopropylamine in hexane cosolvent at 25 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax + b: $a = 0.168 \pm 0.006$; $b = 0.03 \pm 0.03$.

[THF] (M)	Initial Rate $\times 10^3$ (M s ⁻¹)	Standard deviation $\times 10^3$ (M s ⁻¹)
0.23	0.104	0.008
1.47	0.248	0.0007
3.93	0.653	0.007
6.40	1.15	0.07
8.86	1.51	0.03



Figure S-4. Plot of initial rate versus NaDA concentration for the isomerization of 0.76 M 1-pentene with 0.59 M diisopropylamine in 3.93 M THF/hexane at 25 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 19.2 \pm 0.8$; $b = 0.67 \pm 0.03$. The covariance represents measured titer of NaDA.

[NaDA] (M)	Initial Rate $\times 10^4$ (M s ⁻¹)	Standard deviation $\times 10^4$ (M s ⁻¹)
0.027	2.0	0.1
0.080	3.50	0.03
0.15	5.0	0.4
0.20	6.53	0.07
0.405	10.6	0.1



Figure S-5. Plot of initial rate versus concentration of 1,4-cyclohexadiene following product growth at 1558 cm⁻¹ with 0.10 M NaDA in 6.04 M THF/DMEA at -95 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 132 \pm 4$; $b = 0.968 \pm 0.009$.

[alkene] (M)	Initial Rate $\times 10^5$ (AU s ⁻¹)	Standard deviation $\times 10^5$ (AU s ⁻¹)
0.010	1.51	0.06
0.020	3.0	0.1
0.035	5.1	0.7
0.050	7.3	0.8



Figure S-6. Plot of initial rate versus concentration of THF following product growth at 1558 cm⁻¹ with 0.10 M NaDA and 0.020 M 1,4-cyclohexadiene in DMEA cosolvent at -95 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 0.17 \pm 0.05$; $b = 1.7 \pm 0.1$.

[THF] (M)	Initial Rate $\times 10^5$ (AU s ⁻¹)	Standard deviation $\times 10^5$ (AU s ⁻¹)
2.01	0.696	0.03
4.03	1.75	0.2
6.04	3.38	0.4
8.06	6.39	1
10.1	8.57	3



Figure S-7. Plot of initial rate versus concentration of NaDA following product growth at 1558 cm⁻¹ with 0.020 M 1,4-cyclohexadiene in 6.04 M THF/DMEA at -95 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 9.53 \pm 0.8$; $b = 0.52 \pm 0.05$.

[NaDA] (M)	Initial Rate $\times 10^5$ (M s ⁻¹)	Standard deviation $\times 10^5$ (M s ⁻¹)
0.025	1.3	0.3
0.05	1.87	0.8
0.10	2.97	0.2
0.15	3.84	0.9
0.20	4.14	0.8
0.25	4.44	0.8



Figure S-8. Plot of k_{obsd} versus concentration of THF following product growth with 0.10 M NaDA and 0.020 M 1,4-pentadiene at -116 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b + c$: $a = 0.15 \pm 0.02$; $b = 2.23 \pm 0.07$; $c = 0.64 \pm 0.08$. The THF order depicted here is consistent with and supportive of the elevated THF order observed in the metalation of 1,4-cyclohexadiene.

[THF] (M)	$k_{\rm obsd} \times 10^3 ({\rm s}^{-1})$	Standard deviation $\times 10^3$ (s ⁻¹)
0.81	0.7	0.1
1.79	1.28	0.05
2.80	2.2	0.1
3.81	3.6	0.3
5.83	8.4	0.9



Figure S-9. Plot of initial rate versus concentration of NaDA following product growth with 0.020 M 1,4-pentadiene in 2.80 M THF/DMEA at -116 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 2.5 \pm 0.5$; $b = 0.7 \pm 0.1$.

[NaDA] (M)	Initial Rate $\times 10^4$ (s ⁻¹)
0.025	0.167
0.05	0.339
0.1	0.433
0.175	0.617
0.25	0.944



Figure S-10. Plot of initial rate versus concentration of allyl methyl ether following product growth at 1674 cm⁻¹ (methyl enol ether) with 0.10 M NaDA in 5.5 M THF/DMEA at -116 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax: $a = 35 \pm 1$.

[allyl methyl ether] (M)	Initial Rate $\times 10^4$ (AU s ⁻¹)
0.010	0.299
0.030	1.18
0.050	1.65
0.070	2.62
0.10	3.37



Figure S-11. Plot of initial rate versus concentration of THF following product growth at 1674 cm⁻¹ (methyl enol ether) with 0.10 M NaDA and 0.030 M allyl methyl ether at -116 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 0.16 \pm 0.02$; $b = 1.10 \pm 0.05$.

[THF] (M)	Initial Rate $\times 10^4$ (AU s ⁻¹)	Standard deviation $\times 10^4$ (AU s ⁻¹)
2.46	0.43	0.03
4.18	0.72	0.04
6.06	1.17	0.03
8.12	1.7	0.6
10.1	2.0	0.5



Figure S-12. Plot of initial rate versus concentration of NaDA following product growth at 1674 cm⁻¹ (methyl enol ether) with 0.030 M allyl methyl ether in 5.5 M THF/DMEA at -116 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 4.0 \pm 0.3$; $b = 0.52 \pm 0.05$.

[NaDA] (M)	Initial Rate $\times 10^4$ (AU s ⁻¹)	Standard deviation $\times 10^4$ (AU s ⁻¹)
0.025	0.6	0.1
0.05	0.94	0.06
0.10	1.16	0.03
0.15	1.52	0.04
0.20	1.6	0.5
0.25	2.1	0.4



Figure S-13. Plot of k_{obsd} versus THF concentration following product growth at 1601 cm⁻¹ (trimethylsilyl enol ether) with 0.010 M allyloxytrimethylsilane in hexane cosolvent at -78 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax + b: $a = 0.23 \pm 0.04$; $b = 0.14 \pm 0.4$.

[THF] (M)	$k_{\rm obsd} \times 10^3 ({\rm s}^{-1})$	Standard deviation $\times 10^3$ (s ⁻¹)
4.16	1.07	0.05
6.20	1.5	0.2
9.20	2.6	0.2
12.05	2.8	0.3



Figure S-14. Plot of k_{obsd} versus NaDA concentration following product growth at 1601 cm⁻¹ (trimethylsilyl enol ether) with 0.010 M allyloxytrimethylsilane in 6.0 M THF/hexane at -78 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 6.0 \pm 0.6$; $b = 0.60 \pm 0.05$.

[NaDA] (M)	$k_{\rm obsd} imes 10^3 ({ m s}^{-1})$	Standard deviation $\times 10^3$ (s ⁻¹)
0.05	1.00	0.01
0.10	1.6	0.3
0.15	1.8	0.1
0.20	2.25	0.04
0.25	2.7	0.1



Figure S-15. ¹H NMR spectra of 0.78 M allyloxy-*tert*-butyldimethylsilane with 0.27 M NaDA in neat THF at -80 °C. Each spectrum corresponds to an interval of 881 seconds.



Figure S-16. Plot of k_{obsd} versus THF concentration following product growth at 1664 cm⁻¹ (*tert*-butyldimethylsilyl enol ether) with 0.010 M allyloxy-*tert*-butyldimethylsilane in 6.00 M THF/hexane at -78 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax + b: $a = 0.12 \pm 0.06$; $b = 2.9 \pm 0.5$.

[THF] (M)	$k_{\rm obsd} imes 10^4 ({ m s}^{-1})$	Standard deviation $\times 10^4$ (s ⁻¹)
3.53	3.1	0.3
5.61	3.8	0.9
7.69	3.51	0.01
9.77	4.5	1
11.9	3.9	1.4



Figure S-17. Plot of initial rate versus NaDA concentration following product growth at 1664 cm⁻¹ (*tert*-butyldimethylsilyl enol ether) with 0.010 M allyloxy-*tert*-butyldimethylsilane in 6.0 M THF/hexane at -78 °C. The curve depicts an unweighted least-squares fit to the function $f(x) = ax^b$: $a = 12 \pm 1$; $b = 0.56 \pm 0.06$.

[NaDA] (M)	$k_{\rm obsd} imes 10^4 ({ m s}^{-1})$	Standard deviation $\times 10^4$ (s ⁻¹)
0.025	1.5	0.7
0.050	2.4	0.7
0.090	2.6	0.3
0.15	4.1	0.7
0.25	5.3	0.7



Figure S-18. Plot of k_{obsd} versus THF concentration following product growth at 1661 cm⁻¹ (triisopropylsilyl enol ether) with 0.010 M allyloxytriisopropylsilane and 0.10 M NaDA with hexane cosolvent at 0 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax + b: $a = 0.68 \pm 0.04$; $b = 1.2 \pm 0.3$.

[THF] (M)	$k_{\rm obsd} \times 10^3 ({\rm s}^{-1})$	Standard deviation $\times 10^3$ (s ⁻¹)
0.11	0.8	0.1
3.09	3.66	0.07
5.28	5.2	0.7
7.47	6.3	0.6
9.66	7.76	0.07
11.9	8.9	0.8



Figure S-19. Plot of k_{obsd} versus NaDA concentration following product growth at 1661 cm⁻¹ (triisopropylsilyl enol ether) with 0.010 M allyloxytriisopropylsilane in 5.28 M THF/hexane at 0 °C. The curve depicts an unweighted least-squares fit to the function f(x) = ax + b: $a = 0.68 \pm 0.04$; $b = 1.2 \pm 0.3$.

[NaDA] (M)	$k_{\rm obsd} imes 10^3 ({ m s}^{-1})$
0.025	2.38
0.050	3.70
0.10	5.68
0.175	6.70
0.25	7.99



Figure S-20. ¹H NMR spectrum (CDCl₃) of myrcene recovered from reaction of NaDA/THF with trimethylsilyl ether of geraniol. Resonances at δ 0.85 ppm and δ 1.35 ppm correspond to residual pentane from chromatography.



Figure S-21. Representative concentration trace showing loss of 0.050 M geraniol TMS ether 1-trimethylsilyloxy-3-cyclohexyl-2-propene with 0.10 M NaDA in neat THF at -78 °C (monitored by ReactIR).



Figure S-22. ¹H NMR spectra of 0.090 M trimethylsilyl ether of crotyl alcohol with 0.32 M NaDA in neat THF at -60 °C. Each spectrum corresponds to an interval of 263 seconds. These spectra are consistent with stoichiometric consumption of NaDA to give 1,3-butadiene.



Figure S-23. Concentration of 0.090 M trimethylsilyl ether of crotyl alcohol with 0.32 M NaDA in neat THF at -60 °C.



Figure S-24. ¹H NMR spectra of 0.23 M trimethylsilyl ether of prenol with 0.30 M NaDA in neat THF at -60 °C. Each spectrum corresponds to an interval of 131 seconds. These spectra are consistent with stoichiometric consumption of NaDA to give isoprene.



Figure S-25. Concentration of 0.23 M trimethylsilyl ether of prenol with 0.30 M NaDA in neat THF at -60 °C.



Figure S-26. ¹H NMR spectra of 0.080 M 1-trimethylsilyloxy-3-cyclohexyl-2-propene with 0.19 M NaDA in neat THF at -40 °C. Each spectrum corresponds to an interval of 626 seconds. These spectra are consistent with stoichiometric consumption of NaDA to give several products.



Figure S-27. ¹H NMR spectrum of isomerization of 1.5 mL neat allyloxytrimethylsilane with 7 mol % NaDA at room temperature after one minute. The resonances at 2.9 ppm and 1.1 ppm are consistent with diisopropylamine.



Figure S-28. ¹H NMR spectrum (CDCl₃) of isolated material (distillation) from isomerization of 1.5 mL neat allyloxytrimethylsilane with 7 mol % NaDA at room temperature after one minute.



Figure S-29. ¹H NMR spectrum of partial isomerization of 0.47 M allyloxy-*tert*butyldimethylsilane with 0.094 M NaDA and 1.43 M DN*i*-Pr₂ in THF- d_8 . That the relative integration of the terminal methyl (~1.4 ppm) is significantly above 2 is consistent with predominant retention of protons from substrate without trapping of the allylsodium intermediate by DN*i*-Pr₂.

II. Computations

Geometries are optimized at the B3LYP level of theory using the 6–31G(d) basis set. Energies are defined as follows: G is the sum of electronic and thermal free energies calculated at the B3LYP level of theory (T = 195 K). G_{MP2} is derived from a MP2 single point calculation corresponding to the DFT-optimized geometry and includes a thermal correction from the DFT optimization.

Table S-1. Geometric coordinates and thermally corrected MP2 energies for 1-pentene.



G = -196.427062 Hartree $G_{MP2} = -195.6789852$ Hartree

Ate	om	Х	Y	Ζ
С	0.0	00000000	0.00000000	0.00000000
С	-1.	34505100	0.65407700	-0.16177800
С	-2.4	41055200	0.09604400	-0.73768800
Η	-2.	37937500	-0.91648900	-1.13602800
Η	-3.	35559300	0.62465400	-0.83136300
Η	-1.4	42477700	1.67371300	0.22098000
Η	-0.	04326100	-1.03222700	-0.37274100
Η	0.2	24798500	-0.06478100	1.07124600
С	1.1	13172800	0.76137100	-0.71658500
Η	1.	15496900	1.79890600	-0.35477500
Η	0.9	90064400	0.81671400	-1.78845700
С	2.5	50583800	0.11609400	-0.51279000
Η	2.5	52118500	-0.91250900	-0.89360800
Η	3.2	29061900	0.67579600	-1.03417400
Η	2.7	77298800	0.07870800	0.55060700

Table S-2. Geometric coordinates and thermally corrected MP2 energies for $[A(THF)_2(1-pentene)]^{\ddagger}$.



G = -1115.07229 Hartree $G_{MP2} = -1111.273063$ Hartree

Ato	om	Х	Y	Ζ
С	0.00	0000000	0.00000000	0.00000000
С	-1.4	9330000	-0.22641000	0.25981600
С	-2.4	0677500	0.46614700	-0.62462100
С	-3.6	4942800	0.08905100	-1.05336800
Η	-4.0	1271700	-0.93019300	-0.91481500
Η	-4.2	4898000	0.73199500	-1.69350600
Η	-2.0	9247400	1.47583700	-0.91588800
Η	-1.7	2487400	-1.30219700	0.31459300
Н	0.24	4201000	1.06639000	0.13815200
Η	0.2	7461300	-0.22877100	-1.04384000
С	0.89	9422700 -	-0.83882200	0.92092400
Н	0.7	1446600	-1.91050800	0.76443500
Н	1.9	5872200	-0.65191700	0.73183000
Н	0.6	9458900	-0.62359900	1.97672700
Н	-1.7	8085900	0.24358600	1.58894300
Ν	-2.2	5168400	0.66378500	2.80586900
Na	-4.2	1830800	0.49496900	1.51962400
0	-5.8	5479100	2.19753000	1.74938000
С	-6.0	1805400	3.02390700	2.92802600
Н	-5.0	2593800	3.19016200	3.36176500
Н	-6.6	3018400	2.48118900	3.65684300
С	-6.6	7163600	4.33138200	2.45416100
Η	-7.7	6093200	4.28119900	2.56598500
Н	-6.3	1618000	5.19870300	3.01775000
С	-6.2	9182500	4.37841600	0.96493000
Н	-5.2	7402000	4.76408800	0.83727000
Н	-6.9	7041400	4.99219200	0.36514200
Η	0.3	5213900	1.74377600	2.53967200
Η	0.0	7579100	1.43299900	4.25906900
С	-2.0	7406400	2.98533100	2.00322400
Η	-1.7	6727900	4.00428400	2.27034900

С	-6.34558600 2.89950500	0.58837300
Η	-7.37549900 2.57824100	0.37393100
Η	-5.70942600 2.62406400	-0.25743300
0	-5.82157600 -1.20240500	1.83235400
С	-6.97271100 -1.04118500	2.68635100
Η	-7.37053900 -0.03551100	2.52427700
Η	-6.65509500 -1.13129300	3.73473900
С	-7.93480900 -2.16557500	2.29965100
Η	-8.60789100 -2.44173000	3.11679600
Η	-8.54566300 -1.86864100	1.43872800
С	-6.96067800 -3.28710200	1.90582500
Η	-7.40802300 -4.04301300	1.25382100
Η	-6.57994700 -3.79235300	2.80104500
С	-5.83876200 -2.50851200	1.21253900
Η	-4.85121500 -2.96278400	1.33720600
Η	-6.03417600 -2.38252600	0.13995700
С	-1.99683400 -0.34531600	3.83782500
С	-2.60817600 0.00743100	5.20927900
Н	-2.43908900 -0.80100000	5.93253000
Η	-3.69272200 0.15937800	5.11186000
Η	-2.18035400 0.92108600	5.63524900
С	-2.54283500 -1.70881800	3.38625600
Н	-2.31227300 -2.48339900	4.12835800
Н	-2.10592500 -2.01535100	2.43155700
Η	-3.63492300 -1.67160000	3.26950400
Η	-0.91111100 -0.48991000	4.00758100
С	-1.73105100 1.99834000	3.13094700
С	-0.21320100 2.06801800	3.41475800
Η	0.08988900 3.09445200	3.65963100

Table S-3. Geometric coordinates and thermally corrected MP2 energies for pro-Z $[A(THF)_3(1-pentene)]^{\ddagger}$.



G = -1347.419378 Hartree $G_{MP2} = -1342.854422$ Hartree

Atc	m	Х	Y	Ζ
NT	0.0		0 0000000	0 00000000
Na	0.0	7192400	0.00000000	0.00000000
N	1.8	/182400	-1.56019900	-0.21990600
C	2.3	8464500	-1.81/6/100	-1.5/232200
C	1.9	4030900	-3.15395800	-2.19923800
Н	2.3	3031000	-3.24513100	-3.22116800
Н	0.8	4502300	-3.21152200	-2.25069900
Н	2.2	9694000	-4.02265600	-1.63629400
С	1.9	4411400	-0.67275300	-2.49522300
Η	2.4	1510600	-0.74997300	-3.48353000
Н	2.2	1365900	0.29907100	-2.07043800
Н	0.8	5423900	-0.69699500	-2.64047100
Η	3.4	9337700	-1.82565200	-1.58543700
С	2.0	8631200	-2.66248300	0.72787100
С	3.5	6408700	-3.06349400	0.94488300
Η	3.6	4265600	-3.90697900	1.64345900
Η	4.1	3960900	-2.22635800	1.35486500
Η	4.0	4307600	-3.36837900	0.00793600
С	1.4	4788000	-2.30399000	2.07935900
Η	1.5	4153200	-3.13282700	2.79247300
Н	0.3	7946800	-2.07888700	1.96286200
Η	1.9	3168300	-1.42551200	2.52122600
Н	1.5	6705200	-3.57354300	0.37397400
Ο	-1.8	6683700	-1.55864700	-0.66739100
С	-2.7	1532300	-1.36455200	-1.81001500
Н	-2.4	8722600	-0.37562200	-2.21149500
Н	-3.7	7347100	-1.39603600	-1.50304200
С	-2.3	8069900	-2.52794800	-2.74043300
H	-0.3	5340500	4.44574100	-2.97062800
Н	-1.8	7312800	3.65502000	-3.42274700
C	-0.4	0288300	2 28655400	-2 58043300
÷	÷. i			

Η	-3.16640300 -2.72577100 -3.47635100
Η	-1.44930400 -2.31920000 -3.27828700
С	-2.18443600 -3.68405800 -1.74192700
Η	-1.48822800 -4.44415700 -2.10781100
Η	-3.14071100 -4.17895600 -1.53894400
С	-1.65666400 -2.97750700 -0.47423200
Η	-2.19195800 -3.30211300 0.42778000
Η	-0.58478300 -3.11732600 -0.31946700
0	-1.52112100 0.48999600 1.94106300
С	-2.76331700 -0.19034000 2.17842600
Η	-2.91050400 -0.88258600 1.34961700
Η	-3.58204000 0.54669400 2.18532100
С	-2.61064000 -0.85061100 3.56294900
Η	-3.55260300 -0.83465400 4.11962100
Η	-2.30209500 -1.89564100 3.46487900
С	-1.49657200 -0.01102900 4.24944500
Η	-0.61773900 -0.62945400 4.45272300
Η	-1.82137200 0.42997100 5.19653700
С	-1.16340900 1.06826300 3.20463000
Η	-1.76157100 1.97807300 3.36969100
Η	-0.10948200 1.34285700 3.15850200
0	-1.12014300 1.56497500 -1.55027600
С	-2.08396000 2.43891600 -0.92259100
Η	-2.08556200 2.21656200 0.14721400
Η	-3.08269500 2.22556000 -1.33245100
С	-1.63769600 3.85820800 -1.26905800
Η	-2.45949700 4.57985900 -1.22909300
Η	-0.85347500 4.18664500 -0.57826800
С	-1.06468100 3.66633800 -2.68196200
Η	-0.45630800 1.71179500 -3.51093000
---	------------------------------------
Η	0.64617000 2.36170100 -2.27454100
С	3.28754300 0.60428000 0.83020200
С	2.20682600 1.41003500 1.36032900
С	1.47767800 2.40391400 0.77121100
Η	1.77322400 2.85359700 -0.17433700
Η	0.64437800 2.87408500 1.28822700
Η	1.85328300 1.08738100 2.34654800
С	4.19081100 1.22154300 -0.23653800
Η	3.58285900 1.62769000 -1.05889200
Η	4.80957600 0.43443400 -0.69157400
С	5.11364500 2.33989000 0.27930000
Η	5.78622700 1.95932500 1.05849600
Η	5.73655900 2.76209000 -0.52162800
Η	4.52761100 3.15479700 0.71936000
Η	3.88071700 0.15406800 1.63709700
Η	2.59317400 -0.51968400 0.27651100

Table S-4. Geometric coordinates and thermally corrected MP2 energies for pro-E $[A(THF)_3(1-pentene)]^{\ddagger}$.



G = -1347.422632 Hartree $G_{MP2} = -1342.856104$ Hartree

Ato	om X	Y	Ζ
С	0.00000000	0.00000000	0.00000000
С	1.44643100	-0.07906100	-0.48436200
С	2.31927200	-0.97962100	0.23351100
С	3.44352800	-1.62159500	-0.20050800
Η	3.69011300	-1.66968300	-1.26050000
Η	4.05432100	-2.22171000	0.46982200
Η	2.10078200	-1.06562800	1.30655400
Η	1.49086700	-0.23689900	-1.57266400
Η	-0.49291700	0.87367400	-0.45114900
Η	-0.01411200	0.18191300	1.08718100
С	-0.85467700	-1.24424400	-0.29814100
Η	-0.43292000	-2.13494200	0.18381400
Η	-1.89028200	-1.12768700	0.05123700
Η	-0.88682800	-1.44315200	-1.37714600
Η	2.12313000	1.17888900	-0.36441800
Ν	2.88750400	2.30276000	-0.34715300
Na	4.77452900	0.84811100	0.13801100
0	6.60449300	2.56645500	0.45898200
С	7.80598500	2.63770400	-0.32237100
Η	7.82841200	1.74575900	-0.95049200
Η	8.68796700	2.63854300	0.33932700
С	7.68873500	3.95956500	-1.07794500
Η	8.65115400	4.32963100	-1.44503900
Η	7.01666100	3.83791400	-1.93437000
С	7.05413100	4.87953900	-0.01526600
Η	6.41305700	5.64747800	-0.45709300
Η	7.83092900	5.38937900	0.56490000
С	6.25599900	3.90360400	0.88151800
С	3.37128100	4.27660900	-1.86634100
Η	3.30208100	4.60152600	-2.91255900
Η	4.43044800	4.29557500	-1.57644100
Η	2.84277000	5.01865400	-1.25894000

Η	6.51199100 4.02684700 1.94252000
Η	5.17456300 3.99972300 0.76653300
0	5.54954900 0.03549200 2.32611900
С	6.65885600 0.54211100 3.07937100
Η	7.07784900 1.37135500 2.50904400
Η	7.41947800 -0.24818700 3.18609900
С	6.07175400 0.92703600 4.44760600
Η	6.79921600 0.79409000 5.25416100
Η	5.76178500 1.97641200 4.44886600
С	4.83705000 -0.00676400 4.58972200
Η	3.92444900 0.57614100 4.74183800
Η	4.93168300 -0.69957400 5.43127100
С	4.78950000 -0.76280800 3.24567400
Η	5.25525000 -1.75630500 3.33313600
Η	3.78893300 -0.87926000 2.82819100
0	6.51917000 -0.29923800 -1.21604600
С	7.20783300 -1.34692200 -0.49814000
Η	6.72987000 -1.44960700 0.47919700
Η	8.25664900 -1.04881100 -0.35174200
С	7.10401200 -2.59023400 -1.38070800
Η	7.90955300 -3.30692600 -1.19360700
Η	6.14624200 - 3.09493700 - 1.21297500
С	7.14993100 -1.97642300 -2.78901800
Η	6.71592900 -2.62211500 -3.55803800
Η	8.18533400 -1.75679400 -3.07571000
С	6.35392500 -0.68013800 -2.60203100
Η	6.70905400 0.14122900 -3.23330700
Η	5.28525900 -0.82914900 -2.79396600
С	2.79517000 2.85658500 -1.70407700

С	3.52211400	1.91901900	-2.68024800
Η	3.36932400	2.23293900	-3.72083400
Η	3.16395400	0.88970500	-2.58176300
Η	4.60491600	1.92905200	-2.48628800
Η	1.74097100	2.90714500	-2.04524200
С	2.35185200	3.17784200	0.70352100
С	0.87948400	3.61473500	0.52582600
Η	0.57969100	4.30091200	1.32902100
Η	0.20641200	2.75137000	0.54828100
Η	0.72130700	4.13527300	-0.42494600
С	2.51901800	2.48796100	2.06668600
Η	2.15751600	3.12889000	2.88082400
Η	3.57242000	2.25255300	2.26598700
Η	1.95228500	1.55039000	2.09895800
Η	2.94127500	4.11389700	0.75472700

 Table S-5.
 Geometric coordinates and thermally corrected MP2 energies for 1,4pentadiene.



G = -195.196301 Hartree $G_{MP2} = -194.4806266$ Hartree

Ate	om	Х	Y	Ζ
С	0.0	00000000	0.00000000	0.00000000
Η	0.2	28469200	0.45255400	-0.96181500
Η	-0.	01737200	0.83380800	0.72058300
С	1.(04547400	-0.99523000	0.42429700
Η	0.	89593800	-1.44602700	1.40627700
С	2.	10802900	-1.34793500	-0.29944800
Η	2.2	29239800	-0.92933100	-1.28710800
Η	2.3	83764000	-2.06577900	0.06619400
С	-1.4	40497100	-0.55318800	-0.10755400
Η	-2.	18830900	0.19915300	-0.20887600
С	-1.	74339700	-1.84261800	-0.09728700
Η	-0.	99966800	-2.63043400	-0.00881500
Η	-2.	77995700	-2.15642400	-0.18562600

Table S-6. Geometric coordinates and thermally corrected MP2 energies for $[A(THF)_3 \cdot (1,4\text{-pentadiene})]^{\ddagger}$.



G = -1346.169909 Hartree $G_{MP2} = -1341.634158$ Hartree

Ato	om	Х	Y	Ζ
С	0.0	0000000	0.00000000	0.00000000
Ν	-0.7	71360800) -0.29172100	-1.25047400
Na	-2.	7962370	0 -1.42128000	-1.06212000
0	-3.9	95910900) -2.39810400	0.79412900
С	-3.5	50013300) -3.49286900	1.60556200
С	-3.3	35188000) -2.91951300	3.02878500
С	-4.2	25213900) -1.65216200	3.01719700
С	-4.8	38157500) -1.66707500	1.61363200
Η	-5.0)0588700	0 -0.68400100	1.15725000
Η	-5.8	35224000) -2.18818000	1.62280000
Η	-3.6	54908200	0-0.75009000	3.15762300
Η	-5.0)1455900) -1.66745200	3.80193100
Η	-2.3	31149100) -2.65367500	3.23356000
Η	-3.6	56419300) -3.64666000	3.78435800
Η	-4.2	25176700	0 -4.29699800	1.58015500
Η	-2.5	57316700) -3.85816600	1.15972700
0	-4.	15755600	0-2.56073100	-2.70558600
С	-4.8	35092500) -3.75686500	-2.27466800
С	-5.2	29573200) -4.47389900	-3.55323000
С	-4.2	22002800) -4.03675900	-4.55974600
С	-3.9	96972300) -2.58893300	-4.13775400
Η	-4.6	59203500) -1.90491000	-4.60606100
Η	-2.9	95984800	0 -2.23173800	-4.35513400
Η	-4.5	54036000	0 -4.12030800	-5.60248800
Η	-3.3	30971700	0 -4.63340200	-4.43158400
Η	-6.2	28320900	0 -4.11903700	-3.87204400
Η	-5.3	35196900	0 -5.55858700	-3.42290600
С	-0.4	41364300) -1.02589000	1.06534300
Η	0.1	9817000	0-0.92467600	1.97080300
Η	-0.3	30690500	0 -2.05063900	0.69562600
Η	-1.4	16392000	0 -0.87532300	1.35561700

Η	-4.15346100 -4.36565000 -1.68765200
Η	-5.68101000 -3.45961000 -1.62654200
0	-4.45544700 0.37498300 -1.05364700
С	-5.74716400 0.29936200 -1.69580600
С	-6.00106500 1.67337800 -2.35347800
С	-4.61951200 2.35226800 -2.30642900
С	-4.01514800 1.74567000 -1.04255900
Η	-2.92427100 1.73090200 -1.01800700
Η	-4.39083400 2.25098000 -0.13909800
Η	-4.68035600 3.44401800 -2.26679000
Η	-4.01651200 2.07357400 -3.17754400
Η	-6.72790600 2.24955400 -1.77026600
Η	-6.39424900 1.57944700 -3.37009100
Η	-5.69613500 -0.52299100 -2.41455500
Η	-6.51171400 0.05781000 -0.94578300
С	-0.45042500 0.66604000 -2.33033200
С	1.03851900 0.84479300 -2.70763100
Η	1.14730200 1.55401900 -3.53895000
Η	1.48793700 -0.10864000 -3.00569300
Η	1.62391500 1.23309800 -1.86768000
С	-1.24960100 0.25129800 -3.57626400
Η	-1.11686900 0.97132800 -4.39356000
Η	-2.32304300 0.19193300 -3.35190800
Η	-0.92236900 -0.73055700 -3.93963500
Η	-0.81811300 1.67470100 -2.05349300
С	-0.23400600 1.41646300 0.56225000
Η	0.10301600 2.19994200 -0.12489100
Η	0.31004500 1.55397900 1.50583000
Η	-1.30252600 1.57949700 0.76209300

H1.09289900 -0.11197600 -0.13234500C0.35180400 -2.73715300 -2.10809000H0.55386200 -2.52585500 -3.16624700C1.59616900 -2.97547600 -1.35348800H1.47644200 -3.34760500 -0.33310100C2.84909700 -2.72707700 -1.77112700H3.05614900 -2.36741700 -2.77788000H3.70866700 -2.87202100 -1.12196000C-0.76483500 -3.67615900 -1.95835700H-1.51666100 -3.59471800 -2.75028200C-1.05263600 -4.55017700 -0.96291500H-0.36266800 -4.75736700 -0.14845200H-1.95647400 -5.15415500 -0.99041700H-0.15960900 -1.53746400 -1.68933600

Table S-7. Geometric coordinates and thermally corrected MP2 energies for 1,4cyclohexadiene.



G = -233.310874 Hartree $G_{MP2} = -232.4838718$ Hartree

Ate	om	Х	Y	Ζ
С	0.0	000000	0 0.00000000	0.00000000
С	1.2	2551540	0 -0.83325500	-0.00004200
С	1.2	2551540	0 -2.16801300	-0.00004200
С	0.0	000000	0 -3.00126800	0.00000000
С	-1.2	2551540	0 -2.16801300	0-0.00004200
С	-1.2	2551540	0 -0.83325500	0-0.00004200
Η	-2.2	2011230	0 -0.29343500	0-0.00006700
Η	-2.2	2011230	0 -2.70783300	0-0.00006700
Η	0.0	000000	0 -3.67865300	0-0.87080500
Η	0.0	000000	0 -3.67858500	0.87085100
Η	2.2	2011230	0 -2.70783300	0-0.00006700
Η	2.2	2011230	0 -0.29343500	0-0.00006700
Η	0.0	000000	0 0.67731700	0.87085100
Η	0.0	000000	0 0.67738500	-0.87080500

Table S-8. Geometric coordinates and thermally corrected MP2 energies for $[A(THF)_3 \cdot (1,4-cyclohexadiene)]^{\ddagger}$.



G = -1384.281914 Hartree $G_{MP2} = -1379.639054$ Hartree

Ato	om X	Y	Ζ	Н	4.87
NT-	0 00000000	0 00000000	0 0000000	H	3.18
Na N	0.00000000	0.00000000000000000000000000000000000	0.00000000		3.73
N C	-1.19409600	2.03014300	-0.21291100	П	3.01 4.65'
C	-0.9155/800	3.00383900	-1.2/30/300	н С	4.03
	-0.043/9/00	4.20223100	-0.84/00000		2.33^{2}
п	0.14820600	4.80240200	-1./02/8800	П	$\frac{2.11}{1.62}$
п	0.92003300	3.83/40100	-0.40301300	П	0.104
П	-0.31093/00	4.80009000	-0.00703300	C C	0.100
	-0.2369/100	2.29300800	-2.455/4100		1.21
п	-0.09440/00	2.98003800	-3.30143100	П	1.994
п	-0.83090000	1.45050000	-2.80307700	н С	1.38
п	0.75258900	1.91019000	-2.139/0000		0.05
П	-1.83037000	3.43/13000	-1.0/339900	П	1.14.
C	-1./2003/00	2.02230200	1.023/9300	П	0.75^{2}
	-3.03234100	3.43101/00	0.803/0300		-0.80
п	-3.32812300	3.8/801300	1.82400300	П	-1.31
п	-3.8301/900	2.79083100	0.31831900	П	-1.14
П	-2.91/10900	4.23096200	0.14/21400		-1.01
	-1.93/82300	1.31108200	2.00484000	П	-0.9/
п	-2.34913100	1.91083800	2.99730100	П	-1.92
п	-0.98840100	1.01303300	2.312/9400	C C	0.04
п u	-2.03310/00	0.73208200	1.08/38400		1.010
	-0.9/9/4/00	0.52286000	1.43934200	п u	2.00
C	2.29093800	0.33380900	0.34039800	П	2.093
	2.02790100	0.31000/00	1 22560600		0.200
п	2.93/80100	-0.24233900	-1.23300000	П	0.720
П	4.13810000	1 71925000	0.00/89300	П	-0.77
С U	5.83032900	1./1823000	-0.73432000	C	0.32.
п u	1 20205400	2 02861000	-4.38800800		
П	1.29393400	-3.02801000	-4.10399300		
с u	0.10100900	0 72822000	2 66000100		
п	0.//820100	-0./3823900	-3.00808100		

Η	4.87135700 1.71775200 -1.16355400
Η	3.18736800 2.17743100 -1.49053600
С	3.75152100 2.44182400 0.59998500
Η	3.61736000 3.52217100 0.49511900
Η	4.65792100 2.27330400 1.19229900
С	2.53472200 1.76989300 1.26576000
Η	2.71377500 1.53834900 2.32275800
Η	1.62247700 2.36760400 1.18388400
0	0.10653400 -1.70991600 1.71684100
С	1.21174200 -1.84539600 2.62269800
Η	1.99416000 -1.16782100 2.27869600
Η	1.58623400 -2.88086800 2.58628400
С	0.63128700 -1.51903100 4.00785200
Η	1.14388400 -2.06852600 4.80327100
Η	0.73411300 -0.45064700 4.22184200
С	-0.86827500 -1.90704300 3.87492400
Η	-1.51293500 -1.05872300 4.11964000
Η	-1.14381700 -2.73490800 4.53504100
С	-1.01978100 -2.30269400 2.39210200
Η	-0.97192000 -3.39539600 2.26875100
Η	-1.92387100 -1.92940700 1.90834100
0	0.64801500 -1.51613600 -1.74493200
С	1.01032300 -2.86989900 -1.39542900
Η	0.72974000 -3.02267600 -0.35098600
Η	2.09963200 -2.99137000 -1.49424800
С	0.26611300 -3.75547400 -2.39179900
Η	0.72867100 -4.74025400 -2.50865400
Η	-0.77254900 -3.89747400 -2.07205400
С	0.32581400 -2.90034500 -3.66762600

Η	-0.86190800 -1.14283700 -3.11085500
С	-3.06961800 0.29917700 -1.36911700
С	-4.43507400 0.85112800 -1.26499100
С	-5.37791600 0.38340000 -0.42530600
С	-5.15023200 -0.80864500 0.48184400
С	-3.88146800 -1.54600000 0.10512500
С	-2.97232800 -1.03533400 -0.75551700
Η	-2.10324600 -1.65176400 -1.00785600
Η	-3.74107000 -2.54386800 0.52190500
Η	-5.12239000 -0.49255800 1.54410500
Η	-6.01412200 -1.49642200 0.43301600
Η	-6.35549500 0.86198400 -0.37874900
Η	-4.67562400 1.71155100 -1.89178500
Η	-2.69316100 0.31196000 -2.40282000
Η	-2.14702000 1.20469100 -0.74074700

Table S-9.
 Geometric coordinates and thermally corrected MP2 energies for allyl methyl ether.



G = -232.316236 Hartree $G_{MP2} = -231.5298026$ Hartree

Ato	om	Х	Y	Z
С	0.0	00000000	0.00000000	0.00000000
Η	-0.	76316100	-0.78262600	-0.00015300
Η	-0.	13659800	0.63226600	0.89297300
Η	-0.	13651100	0.63252900	-0.89279900
Ο	1.2	25529000	-0.64147500	-0.00003400
С	2.3	33503700	0.27428800	0.00015400
Η	2.2	27760500	0.93251000	-0.88578100
Η	2.2	27751100	0.93225000	0.88627700
С	3.0	52214700	-0.51204900	0.00010800
Η	3.4	48782200	-1.59248700	0.00022700
С	4.8	84346100	0.01906400	-0.00004400
Η	5.0	00828100	1.09534300	-0.00016900
Η	5.	73301500	-0.60403300	-0.00004800

Table S-10. Geometric coordinates and thermally corrected MP2 energies for pro-Z $[A(THF)_3 \cdot (allyl methyl ether)]^{\ddagger}$.



G = -1383.296497 Hartree $G_{MP2} = -1378.692455$ Hartree

Ato	om X	Y	Ζ	Н	2
				Η	1
Na	0.00000000	0.00000000	0.00000000	С	0
Ν	-2.02382300	1.08297800	-0.79721800	Н	0
С	-1.99530400	1.91826600	-2.00311500	Н	1
С	-2.03192500	3.43501300	-1.73958300	С	0
Η	-1.96738100	3.99116700	-2.68388500	Η	-(
Η	-1.18376600	3.73775300	-1.11080400	Η	-(
Η	-2.95216500	3.74992300	-1.23690700	0	0
С	-0.73456600	1.58132000	-2.81356400	С	0
Η	-0.72820100	2.10882300	-3.77607600	Η	1
Η	-0.67302700	0.50732900	-3.01588100	Η	1
Н	0.16784400	1.87988700	-2.26047500	С	-(
Η	-2.85786800	1.69610700	-2.66477900	Η	0
С	-3.14175700	1.36386600	0.11626400	Η	-(
С	-4.54818000	1.25252500	-0.51566300	С	-1
Η	-5.32523100	1.50002900	0.21931700	Η	-2
Η	-4.73178700	0.23314200	-0.87232800	Η	-(
Η	-4.67063500	1.93495500	-1.36405800	С	-(
С	-3.05372400	0.42261400	1.32736000	Η	0
Η	-3.83894700	0.65100200	2.05934900	Η	-(
Η	-2.08510100	0.52619300	1.83475700	0	2
Η	-3.17435000	-0.62248600	1.01804900	С	3
Η	-3.05301000	2.39339500	0.51172500	Η	2
0	1.05060400	1.89739700	1.24125900	Η	3
С	2.28182900	2.50749100	0.82197500	С	3
Η	2.71047600	1.86596500	0.04925800	Η	4
Η	2.98089500	2.56603700	1.67170200	Η	2
С	1.88436700	3.90205000	0.34325500	С	3
Η	3.61555800	-1.95307900	-3.52955900		
Η	4.54754200	-0.72735300	-2.65223900		
С	2.39568400 -	0.49714400	-2.42852200		
Η	2.48029800	0.49788700	-2.87670600		

Η	2.72425500 4.60387200 0.32938200
Η	1.46489200 3.84518400 -0.66752400
С	0.79473700 4.28178100 1.36133400
Η	0.07347200 5.00188600 0.96419900
Η	1.24841300 4.72218800 2.25654400
С	0.13650300 2.92755500 1.69272500
Η	-0.03828700 2.80426300 2.76882400
Η	-0.80410800 2.77175800 1.15857200
0	0.50340200 -1.29413300 1.97655100
С	0.91845400 -0.79194300 3.25342500
Η	1.20041300 0.25212700 3.11007800
Η	1.79503400 -1.36111600 3.60283600
С	-0.28730800 -1.01548500 4.17975900
Η	0.02075600 -1.18965000 5.21530500
Η	-0.94230900 -0.13897400 4.16939800
С	-1.00869200 -2.23996900 3.55044800
Η	-2.05559000 -2.00897500 3.33746500
Η	-0.98842800 -3.11542700 4.20663000
С	-0.22726700 -2.50370100 2.24588400
Η	0.48605000 -3.33263400 2.37296000
Η	-0.85244500 -2.69568000 1.37370700
0	2.21933700 -0.34408600 -1.00100800
С	3.14364300 -1.20203500 -0.29547800
Η	2.60538100 -1.65081800 0.54308200
Η	3.96779300 -0.59025100 0.09984000
С	3.65017300 -2.20548000 -1.33176700
Η	4.63417700 -2.61145600 -1.07821100
Η	2.94805900 - 3.04132400 - 1.42861100
С	3.65056200 -1.35685800 -2.61296700

Η	1.50334400 -0.98672900 -2.83552400
С	-2.14345400 -1.51230700 -1.71394200
С	-3.31350700 -2.31562300 -1.38329200
С	-3.52727700 -3.16644500 -0.35509000
Η	-2.74369100 -3.43179000 0.35028200
Η	-4.49743700 -3.62817700 -0.19631200
Η	-4.15667200 -2.12103800 -2.05314000
Η	-1.97080800 -1.46612600 -2.80176700
Ο	-0.91148300 -1.92950300 -1.05898500
С	-0.37654100 -3.14169600 -1.56625400
Η	-0.04046100 -3.01951900 -2.60888400
Η	0.48592000 -3.41849900 -0.94694800
Η	-1.11844100 -3.94810600 -1.53212500
Η	-2.20209500 -0.14485600 -1.25106400

Table S-11. Geometric coordinates and thermally corrected MP2 energies for pro- $[A(THF)_3 \cdot (allyl methyl ether)]^{\ddagger}$.



G = -1383.294895 Hartree $G_{MP2} = -1378.689974$ Hartree

Ato	om	Х	Y	Ζ
Na	0.	00000000	0.00000000	0.00000000
Ν	-2.	02751600	1.17882600	-0.67558400
С	-2.	06054100	1.82638800	-1.99389000
С	-1.9	97603600	3.36283500	-1.95711600
Η	-1.	95721900	3.77144500	-2.97573600
Η	-1.	06023300	3.68802100	-1.44514900
Η	-2.	82859100	3.81464900	-1.43971600
С	-0.	90742000	1.27579400	-2.84545300
Η	-0.	95469100	1.65756700	-3.87336700
Η	-0.	94048100	0.18239700	-2.88870500
Η	0.0	06143300	1.57611000	-2.42012600
Η	-2.	99412000	1.57759500	-2.53752800
С	-3.	04262400	1.66970900	0.27015900
С	-4.	50627100	1.51083700	-0.20114800
Η	-5.	20032100	1.91094300	0.54998900
Η	-4.	75160100	0.45557600	-0.36800200
Η	-4.	69191400	2.05181000	-1.13602900
С	-2.	85554300	0.95477300	1.61710300
Н	-3.	57436500	1.31905200	2.36175800
Н	-1.	84547700	1.12261400	2.01292400
Н	-3.	00625000	-0.12496700	1.50576800
Η	-2.	88496600	2.74648500	0.46239200
0	1.	32912800	1.92015000	0.89319700
С	2.5	55784300	2.35453800	0.28731100
Η	2.3	84502300	1.59082000	-0.43821700
Η	3.	34430000	2.43982600	1.05420500
С	2.2	23591900	3.71761900 -	-0.32109400
Η	3.0	02981700 -	-2.70137200	-3.56366700
Η	4.	14238300 -	-1.43382700	-3.02154700
С	2.0)5789600 -	-1.00676500	-2.56221100
Η	2.	14524700	-0.09363600	-3.15959300

Η	3.12784200 4.32694600 -0.49785200
Η	1.71145400 3.58785800 -1.27452400
С	1.29413600 4.31592800 0.73830900
Н	0.60565800 5.05927900 0.32600800
Н	1.87496800 4.80141400 1.53093000
С	0.55207200 3.08042600 1.28405100
Н	0.46317600 3.09961500 2.37741400
Н	-0.44201500 2.95536300 0.84715400
0	0.59603300 -1.14638500 2.04867200
С	1.19045500 -0.54729400 3.20807100
Н	1.54175900 0.44208700 2.91148600
Н	2.04935100 -1.15635500 3.53223900
С	0.08274900 -0.54580800 4.27626900
Η	0.49137100 -0.68108300 5.28234800
Η	-0.46195000 0.40287400 4.26353400
С	-0.84807500 -1.71377300 3.84330400
Н	-1.86314800 -1.35072500 3.66189100
Н	-0.90631900 -2.50422200 4.59756300
С	-0.21193600 -2.22999900 2.53735000
Η	0.43261200 -3.10189700 2.72933500
Н	-0.92413600 -2.47591600 1.74903200
0	2.07427200 -0.64354400 -1.16147300
С	3.04555100 -1.44497700 -0.45560300
Н	2.60635100 -1.72797300 0.50429400
Η	3.94436000 -0.83848900 -0.26999000
С	3.35840200 -2.62377100 -1.37763700
Н	4.34753800 -3.05294500 -1.19080600
Η	2.61172200 -3.41609800 -1.25091500
С	3.22571100 -1.97865000 -2.76612100

H 1.09191500 -1.47472400 -2.78262400
O -1.11997200 -1.93248700 -0.76983900
C -2.44806000 -1.49777200 -1.21181500
C -3.52891300 -2.19405500 -0.51899500
C -4.76026700 -2.49294200 -0.98509300
H -5.03202400 -2.33046400 -2.02676600
H -5.54095300 -2.87604000 -0.33364600
H -3.33045700 -2.39853700 0.53978900
H -2.50521200 -1.60677200 -2.30736900
C -0.79172900 -3.25856900 -1.14239600
H -0.75711500 -3.36741900 -2.23857400
H 0.19673900 -3.49530800 -0.72817600
H -1.52770600 -3.97699400 -0.75571700
H -2.32845200 -0.05600000 -0.91412600

Table S-12. Geometric coordinates and thermally corrected MP2 energies for allyloxytrimethylsilane.



G = -601.664955 Hartree $G_{MP2} = -600.0511662$ Hartree

At	om	Х	Y	Ζ
С	0.0	0000000	0.00000000	0.00000000
Η	0.0)5666200	1.04776400	-0.33103700
Η	-0.	15535000	0.00852600	1.09195800
С	1.2	28718400	-0.70631600	-0.31609100
Η	1.2	28161000	-1.77998100	-0.13002000
С	2.3	38488800	-0.10519900	-0.77249900
Η	2.4	40903900	0.96361000	-0.97682600
Η	3.3	30402600	-0.65426200	-0.95821300
0	-1.0	07830300	-0.67941500	-0.63431700
Si	-2.6	59405000	-0.24700300	-0.49355100
С	-2.9	97013000	1.49386900	-1.17508900
Η	-4.0	03086100	1.77048400	-1.12537300
Η	-2.4	41060500	2.25231400	-0.61427900
Η	-2.0	65873100	1.55930100	-2.22428000
С	-3.6	52283100	-1.51752900	-1.52155400
Η	-3.4	45599400	-2.53136000	-1.14032600
Η	-4.′	70315600	-1.32795500	-1.50866500
Η	-3.2	29129100	-1.49641900	-2.56587300
С	-3.2	24426000	-0.31922600	1.31351500
Η	-4.	31154500	-0.08073300	1.40362300
Η	-3.0	09181500	-1.32037100	1.73386600
Η	-2.0	69876000	0.39301700	1.94438800

Table S-13. Geometric coordinates and thermally corrected MP2 energies for pro-Z $[A(THF)_3 \cdot (allyloxytrimethylsilane)]^{\ddagger}$.



G = -1752.641904 Hartree $G_{MP2} = -1747.217566$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0	0000000	0.00000000	0.00000000
Ν	-0.2	1848100	-2.08640500	-1.19752800
С	-0.4	4656800	-1.99564700	-2.64421300
С	-1.8	9021300	-2.29360100	-3.08850400
Н	-1.9	9768700	-2.14949600	-4.17146300
Η	-2.5	9340400	-1.61644000	-2.58468800
Η	-2.1	9502700	-3.32096700	-2.86402700
С	-0.0	5865800	-0.58798700	-3.12112900
Н	-0.0	9749900	-0.51280900	-4.21538000
Η	0.9	5414400	-0.32605300	-2.79923800
Η	-0.7	5541400	0.15751400	-2.71034200
Η	0.2	0445300	-2.70241200	-3.19956900
С	-0.6	4887000	-3.35248200	-0.58749000
С	-0.0	1158300	-4.62284000	-1.19698500
Η	-0.4	0095100	-5.52549800	-0.70839700
Η	1.0	7610800	-4.61211800	-1.06840000
Η	-0.2	2243800	-4.71286600	-2.26842700
С	-0.3	6727600	-3.31162400	0.92232500
Η	-0.7	2939800	-4.22434400	1.41281600
Η	-0.8	7455800	-2.45727900	1.38931700
Η	0.7	0717900	-3.22310400	1.12063900
Н	-1.7	4330400	-3.46829700	-0.69786100
0	-2.5	7029500	0.31521400	0.46885000
С	-3.3	4507700	1.33569900	-0.18478900
Η	-2.6	7696100	1.87672700	-0.85913900
Н	-3.7	3241200	2.04066000	0.56741200
С	-4.4	8956000	0.60489000	-0.88711900
Η	0.3	5566600	5.25096000	-2.20283600
Η	-1.2	8682500	4.77044200	-1.74121700
С	0.1	0825600	3.10413200 -	-1.81841800
Η	-0.5	0412300	2.64604600	-2.60109000

Η	-5.35713600 1.24844600 -1.06323500
Η	-4.15263300 0.20894700 -1.85192800
С	-4.76940900 -0.54208800 0.09594200
Η	-5.26700100 -1.39863100 -0.36826200
Η	-5.40148000 -0.18961300 0.91979400
С	-3.36278200 -0.89088000 0.59935000
Η	-3.35188400 -1.20441900 1.64939900
Η	-2.88833800 -1.66919200 -0.00448000
0	0.23957200 0.55653400 2.33641500
С	-0.79836300 0.54276100 3.32582800
Η	-1.74900100 0.51888300 2.79124700
Η	-0.74285900 1.46779500 3.92211700
С	-0.50802300 -0.69112100 4.19932000
Н	-0.79595300 -0.52372600 5.24181400
Н	-1.06459000 -1.56011900 3.83576900
С	1.02129400 -0.91238400 4.02568400
Н	1.22892900 -1.90198500 3.61030900
Η	1.56639300 -0.82510800 4.97045000
С	1.44266500 0.18529900 3.03058300
Η	1.84135300 1.06788400 3.55450600
Η	2.15707500 -0.14234700 2.27571800
0	-0.14999600 2.41576400 -0.57451100
С	-0.36183400 3.37531600 0.48704500
Η	0.19111100 3.03001700 1.36331000
Н	-1.43175600 3.39649800 0.73509900
С	0.10522000 4.72188800 -0.06838300
Η	-0.39973900 5.56831100 0.40697400
Η	1.18489100 4.84270400 0.07672200
С	-0.22208400 4.57729200 -1.56315300

Η	1.16454000 2.96934400 -2.08126700
0	2.47415000 -0.38807900 -0.34402400
С	2.50051200 -1.74984100 -0.91190000
С	3.18188700 -2.73678300 -0.09138800
С	3.53476700 -2.71277900 1.21413300
Η	3.43825200 -1.81945500 1.82394800
Η	3.96291100 - 3.58638100 1.69700000
Η	3.34196400 - 3.67569500 - 0.63104100
Η	2.89783200 -1.71590100 -1.93667100
Si	3.76682900 0.64453100 -0.65792700
С	3.82003400 1.08537900 -2.50119500
Η	4.61091900 1.81961200 -2.70195600
Η	4.02820000 0.20434200 -3.11932100
Η	2.87152300 1.51204300 -2.84902200
С	3.41715000 2.20455400 0.35128400
Η	3.54203700 2.02129200 1.42493300
Η	4.09581300 3.02034700 0.07242000
Η	2.38842300 2.54155400 0.18632600
С	5.44881400 -0.06279800 -0.17585000
Η	6.24343200 0.65145900 -0.43062100
Η	5.50946000 -0.27452100 0.89631900
Η	5.65492300 -1.00107400 -0.70045100
Η	1.09010100 -2.02982600 -1.06298600

Table S-14. Geometric coordinates and thermally corrected MP2 energies for pro-
E $[A(THF)_3 \cdot (allyloxytrimethylsilane)]^{\ddagger}$.



G = -1752.64151 Hartree $G_{MP2} = -1747.216122$ Hartree

Ate	om	Х	Y	Ζ
Na	0.	00000000	0.00000000	0.00000000
Ν	0.0	03283800	-2.19033500	-1.04755400
С	-0.	18941100	-2.21652900	-2.49952200
С	-1.:	56252100	-2.76623500	-2.92427100
Η	-1.	68715900	-2.69397200	-4.01249400
Η	-2.	36814600	-2.18933600	-2.45040300
Η	-1.	69525200	-3.81742200	-2.64916800
С	-0.0	02390700	-0.79314900	-3.05362000
Η	-0.	05982000	-0.78689100	-4.15058400
Η	0.9	93208600	-0.35672000	-2.74521100
Η	-0.	83502600	-0.14560400	-2.68940300
Η	0.:	57190500	-2.84040500	-3.00976600
С	-0.	18874000	-3.47760100	-0.37152500
С	0.6	57302700	-4.65067700	-0.89119000
Η	0.4	43272000	-5.57497200	-0.34912400
Η	1.'	74050200	-4.44176900	-0.75445000
Η	0.4	49819700	-4.84313200	-1.95585100
С	0.0)4979600	-3.29617100	1.13529400
Η	-0.	15684700	-4.22453500	1.68246600
Η	-0.	59799700	-2.50996100	1.54312800
Η	1.(09153400	-3.01740700	1.32851000
Η	-1.	24453100	-3.78342000	-0.49054700
0	0.	15104100	0.60796200	2.31585700
С	-0.9	91526400	0.67930900	3.27178400
Η	-1.	84999900	0.61457300	2.71286100
Η	-0.	87014200	1.64794400	3.79494100
С	-0.0	66281600	-0.48401000	4.24735300
Si	3.6	57851500	1.08443600 -	-0.42573500
С	4.(04913800	1.39002900	-2.25725300
Η	4.′	79387200	2.18683000	-2.37942300
Η	4.4	44554800	0.49135900	-2.74276200

Η	-0.98068600 -0.23461400 5.26435700
Η	-1.21697700 -1.37427400 3.93516600
С	0.86858700 -0.72974900 4.13939900
Η	1.07379900 -1.75541600 3.82144200
Η	1.38651500 -0.56565400 5.08911400
С	1.33034400 0.27744200 3.06735000
Η	1.72974400 1.19285900 3.53062400
Η	2.06140700 -0.11304900 2.35861500
0	-0.48165200 2.30363300 -0.78021700
С	-0.76910400 3.31470500 0.21421500
Η	-0.16479400 3.10057500 1.09942900
Η	-1.82966100 3.24105900 0.48888300
С	-0.45441100 4.66065200 -0.44511300
Η	-1.06485300 5.47469000 -0.04263800
Η	0.59971100 4.92220500 -0.29930300
С	-0.73153400 4.36616700 -1.92777700
Η	-0.21793800 5.05013200 -2.60991600
Η	-1.80682800 4.42232800 -2.13595500
С	-0.23314600 2.92515200 -2.05998300
Η	-0.75368900 2.34602300 -2.82798500
Η	0.84374600 2.89156100 -2.27032100
Ο	2.44310500 -0.04677600 -0.25869500
С	2.66708900 -1.42242600 -0.79469100
С	3.52737700 -2.23219500 0.05696400
С	4.34724400 - 3.24242100 - 0.31258900
Η	4.53498500 -3.47314400 -1.35981800
Η	4.82454100 - 3.88677800 0.42033200
Η	3.40472200 - 2.05213100 1.13044700
Η	3.03601400 -1.34539600 -1.82676800

Η	3.14663800 1.68940700 -2.80494300
С	3.03073200 2.68030100 0.35647000
Η	2.87644100 2.55997000 1.43535900
Η	3.73807800 3.50642100 0.21128400
Η	2.07245900 2.96908600 -0.08629600
С	5.26719300 0.56997300 0.45736000
Η	6.05711000 1.31470100 0.29283000
Η	5.11157100 0.48543400 1.54001600
Η	5.62631900 -0.40093800 0.10247000
0	-2.61588800 0.04972200 0.43434400
С	-3.51430000 0.91264200 -0.28276900
Η	-2.90739300 1.55235300 -0.92719700
Η	-4.06620800 1.54670500 0.42933800
С	-4.46588700 -0.01634500 -1.03484400
Η	-5.41434100 0.46607800 -1.29120200
Η	-3.99799600 -0.37020900 -1.96046700
С	-4.62165800 -1.17180400 -0.03337500
Η	-4.91835500 -2.11357300 -0.50421700
Η	-5.37646100 -0.91962800 0.72090900
С	-3.22431300 -1.25544800 0.59974000
Η	-3.26026200 -1.49482800 1.66917800
Η	-2.58063700 -1.98143500 0.09623100
Η	1.28652300 -1.92152800 -0.91004100

 Table S-15. Geometric coordinates and thermally corrected MP2 energies for allyloxy-tertbutyldimethylsilane.



G = -719.514504 Hartree $G_{MP2} = -717.4643974$ Hartree

Ato	om	Х	Y	Ζ
С	0.0	00000000	0.00000000	0.00000000
Η	-0.2	20510000	0.55381700	0.92820000
Η	-0.0	04023500	0.72417200	-0.83075400
С	-1.(04586300	-1.05787000	-0.20712300
Η	-0.3	86340500	-1.72931000	-1.04577900
С	-2.1	14006100	-1.19236800	0.54070100
Η	-2	33583700	-0.54006100	1.38968600
Η	-2.8	88534400	-1.95580200	0.33454900
0	1.2	28544400	-0.61037600	0.04382700
Si	2.7	4073000	0.21570700	0.20868600
С	2.7	74290500	1.20389000	1.82242600
Η	3.6	69978600	1.71835700	1.97275400
Η	1.9	96104900	1.97338200	1.82008700
Η	2.5	56957600	0.55946300	2.69164300
С	4.()6537700	-1.16209700	0.21556300
С	3.9	99581400	-1.95820200	-1.10558600
Η	4.7	74338600	-2.76498400	-1.10508900
Η	3.0	01051300	-2.41517400	-1.24820300
Η	4.2	20239400	-1.32461800	-1.97704500
С	5.4	47389400	-0.54650200	0.36223800
Η	5.5	58435800	0.01514400	1.29837200
Η	6.2	23689000	-1.33834600	0.36795700
Η	5.7	71754100	0.13079400	-0.46593200
С	3.8	80720500	-2.12536200	1.39428900
Η	4.5	54724600	-2.93912100	1.39181500
Η	3.8	88672700	-1.61686400	2.36313000
Η	2.8	81132900	-2.57744900	1.33306200

- C 2.97624400 1.41000600 -1.24063000
- Н 3.94066400 1.92818900 -1.17462800
- Н 2.94247400 0.88865400 -2.20389400
- Н 2.19706000 2.18197400 -1.25497200

Table S-16. Geometric coordinates and thermally corrected MP2 energies for pro-Z $[A(THF)_2 \cdot (allyloxy-tert-butyldimethylsilane)]^{\ddagger}$.



G = -1638.147241 Hartree $G_{MP2} = -1633.055742$ Hartree

Ato	m X		Y	Ζ
Na	0.000	00000	0.00000000	0.00000000
Ν	0.9339	2300	2.17820300	-0.00532400
С	1.0455	9600	2.98232500	-1.22862900
С	2.4878	4300	3.21856200	-1.71380400
Н	2.4872	3500	3.76248700	-2.66708600
Н	3.0057	8400	2.26226800	-1.86867400
Η	3.0786	8100	3.80606800	-1.00359800
С	0.2418	5000	2.30265200	-2.34750700
Η	0.2190	1400	2.92219600	-3.25298700
Η	-0.7902	26400	2.11719100	-2.03618500
Η	0.6971	6100	1.33686700	-2.60831600
Η	0.5971	9100	3.98828200	-1.08994500
С	1.7586	7800	2.64938900	1.11564200
С	1.4211	5000	4.07324100	1.61410600
Η	2.0974	1400	4.37744900	2.42366000
Η	0.3948	3900	4.11956300	1.99603700
Η	1.5124	5000	4.81510800	0.81285600
С	1.6572	6300	1.65360700	2.28134600
Η	2.2889	7200	1.96548600	3.12259200
Η	1.9909	8100	0.65447700	1.96748000
Η	0.6256	8100	1.57687800	2.64692000
Η	2.8208	5400	2.66226700	0.81479300
Ο	1.3793	9500	-1.29348900	-1.49961700
С	1.1743	7600	-1.81637700	-2.83138000
Η	0.6165	6200	-1.07549200	-3.41845200
Η	0.5706	7600	-2.72622700	-2.75498000
С	2.5717	8600	-2.05411100	-3.41007200
Si	-3.6793	1500	0.66915200	-0.69128500
С	-3.8727	4100	1.76482200	-2.22528200
Η	-4.8573	57300	1.63345700	-2.69066900
Η	-3.7804	5500	2.82405200	-1.95717500

Η	2.93922700 -3.04891500 -3.13128100
Η	2.58843100 -1.97982400 -4.50140400
С	3.39631600 -0.96208400 -2.71056500
Н	3.25417600 0.00390700 -3.20845900
Н	4.46857700 -1.17769600 -2.68475100
С	2.77261000 -0.93683900 -1.31490200
Н	3.23881500 -1.68035300 -0.65311400
Η	2.80192200 0.04421500 -0.83222000
0	0.02523500 -1.83441000 1.48399700
С	0.53971800 -3.14192700 1.18923400
Η	1.03775100 -3.07932400 0.22061200
Η	-0.29933200 -3.85007600 1.11714600
С	1.46673200 -3.50078000 2.37088300
Η	1.35398400 -4.55118100 2.65584200
Η	2.51773100 -3.34164700 2.11161600
С	1.02107800 -2.52952700 3.49981500
Η	1.80895900 -1.80173500 3.71583500
Н	0.77928900 -3.04652700 4.43309200
С	-0.20463700 -1.82253800 2.90373400
Н	-1.13071100 -2.37544500 3.12219100
Н	-0.33562800 -0.78391600 3.21075400
0	-2.13998400 0.89443400 -0.03770200
С	-1.77236700 2.17500800 0.59899200
С	-2.06169100 2.22058700 2.02365000
С	-2.34251800 1.21455800 2.88297500
Η	-2.51443900 0.19925000 2.53764500
Η	-2.45655800 1.39825300 3.94755200
Η	-1.93463500 3.22249200 2.44468800
Η	-2.22033300 3.01087800 0.04493400

Η	-3.11103500 1.55232700 -2.98400800
С	-3.78690600 -1.18952000 -1.17656800
С	-2.74058800 -1.53750300 -2.25540200
Η	-1.71634100 -1.35961000 -1.90555600
Η	-2.81114000 -2.60052500 -2.53273600
Η	-2.88311600 -0.95105100 -3.17116200
С	-5.19328200 -1.48999800 -1.74083400
Η	-5.27339000 -2.54958700 -2.02571800
Η	-5.98102500 -1.29090700 -1.00467400
Η	-5.41515300 -0.89571600 -2.63600600
С	-3.54647600 -2.07982900 0.06162300
Η	-4.28098300 -1.88333900 0.85199900
Η	-3.63503700 -3.14348500 -0.20865200
Η	-2.54897100 -1.92585100 0.48958400
С	-5.04893200 1.12083900 0.52851500
Η	-6.03171800 1.08621100 0.04175500
Η	-5.06939600 0.45529100 1.39687900
Η	-4.89790600 2.13639000 0.90919600
Η	-0.35182400 2.27426800 0.35090600

Table S-17. Geometric coordinates and thermally corrected MP2 energies for IRC of pro-Z $[A(THF)_2 \cdot (allyloxy-tert-butyldimethylsilane)]^{\ddagger}$.



G = -1638.162218 Hartree $G_{MP2} = -1633.063992$ Hartree

Ato	m X	Y	Ζ
Na	0.0000000	0.00000000	0.00000000
Ν	3.36272100	-0.93377000	-0.53662500
С	3.86728400	-2.23563900	-0.06200000
С	5.24250600	-2.09800200	0.60249700
Н	5.54401600	-3.05370800	1.04611200
Η	5.20712000	-1.34420900	1.39883000
Η	6.02404100	-1.80570100	-0.10659400
С	2.85584100	-2.82462400	0.92866400
Η	3.14749800	-3.84137800	1.21750800
Н	1.84868200	-2.86739300	0.49993200
Н	2.80896700	-2.21186600	1.83728500
Η	3.97103300	-2.95667400	-0.89362700
С	4.19014300	-0.21770700	-1.52835300
С	4.42928600	-0.99399500	-2.83712200
Н	5.05146600	-0.41163300	-3.52790600
Η	3.47533900	-1.20578900	-3.33551800
Η	4.93810800	-1.94804800	-2.66118200
С	3.52916800	1.13494500	-1.81881000
Η	4.12404800	1.71783600	-2.53217300
Η	3.42205100	1.71691500	-0.89635600
Η	2.53057800	0.97964500	-2.24527800
Η	5.16429100	-0.02169000	-1.06239300
0	0.71294700	0.49318500	2.16314200
С	0.10805900	-0.06388100	3.35165300
Η	0.02129200	-1.15145100	3.22948900
Н	-0.89772000	0.35546900	3.45325200
C	1.04605900	0.30062700	4.50385100
Si ·	-3.58/15300	-1.52646400	-0.39027000
C	-4.23676300	-3.07/017200	0.49119700
H	-5.30478900	-3.22565300	0.29745800
Η	-3.70554600	-3.96110800	0.13816100

Η	0.81408400 1.30226500 4.88539500
Η	0.97677900 -0.40471100 5.33734500
С	2.41705600 0.29401400 3.80954500
Η	2.78973100 -0.73220800 3.71233400
Η	3.17409100 0.88051600 4.33879300
С	2.08753700 0.87507200 2.43263800
Η	2.14468700 1.97273900 2.44015600
Н	2.71140500 0.48431600 1.62240300
0	-0.42207000 2.24632700 -0.47570100
С	-0.48686100 3.36964600 0.41340900
Η	-0.14553700 3.02434200 1.39156900
Η	-1.53134500 3.70344700 0.49855300
С	0.39726500 4.46385500 -0.22599400
Η	-0.07877700 5.44628400 -0.15219100
Η	1.36866100 4.53092600 0.27255600
С	0.56170700 3.99763600 -1.69962900
Η	1.59286100 3.68733300 -1.89150100
Η	0.30637200 4.77876400 -2.42179800
С	-0.38071500 2.78975500 -1.80412400
Η	-1.39473600 3.09468800 -2.10356700
Η	-0.03456000 1.99455900 -2.46732900
0	-1.94331000 -1.33467200 -0.03660300
С	-1.08554900 -2.43441200 -0.38043000
С	-0.15051800 -2.29414200 -1.36899800
С	0.16882300 -1.16930600 -2.18959500
Н	-0.65200700 -0.48332200 -2.42271700
Η	0.79522100 -1.37322800 -3.05688300
Η	0.50119500 -3.16793000 -1.46055400
Η	-1.16778300 -3.28938800 0.28148000

Η	-4.09822300 -3.00992500 1.57713000
С	-4.45029500 0.04875000 0.29253600
С	-4.09139600 0.24805500 1.78019600
Η	-3.01089500 0.37139900 1.91907100
Η	-4.58667700 1.14600100 2.18000400
Η	-4.41016600 -0.60141100 2.39682500
С	-5.98201200 -0.10123700 0.16287900
Η	-6.48826400 0.80277400 0.53310500
Η	-6.29445900 -0.24281700 -0.87945200
Η	-6.36403700 -0.94867500 0.74469000
С	-4.00840700 1.29761200 -0.50027200
Η	-4.28191400 1.22426700 -1.55996200
Η	-4.50126900 2.19708700 -0.09975300
Η	-2.92566800 1.45889300 -0.44427000
С	-3.83991100 -1.71803600 -2.24982000
Η	-4.88045200 -1.95912000 -2.49977200
Η	-3.55801500 -0.81325200 -2.79870800
Η	-3.20511500 -2.53270500 -2.61500000
Η	2.44081500 - 1.08079400 - 0.96824800

Table S-18. Geometric coordinates and thermally corrected MP2 energies for pro- $[A(THF)_2 \cdot (allyloxy-tert-butyldimethylsilane)]^{\ddagger}$.



G = -1638.147473 Hartree $G_{MP2} = -1633.054489$ Hartree

Ato	m	Х	Y	Ζ
Na	0.0	0000000	0.00000000	0.00000000
Ν	1.15	685700	2.04165700	-0.21493200
С	1.58	614600	2.49530000	-1.54376200
С	3.11	026600	2.47569400	-1.76085800
Η	3.35	615300	2.75926200	-2.79228600
Η	3.51	217000	1.46941900	-1.57910600
Η	3.63	618400	3.16955100	-1.09724100
С	0.91	334500	1.61967500	-2.61296200
Η	1.12	2964700	1.99083800	-3.62269700
Η	-0.17	7380800	1.60078500	-2.48602600
Η	1.28	3451900	0.58644900	-2.54858400
Н	1.25	535700	3.53546800	-1.73872900
С	1.79	085500	2.74548900	0.90978600
С	1.53	011200	4.26777600	0.95165800
Н	2.06	5105200	4.73088600	1.79388600
Η	0.45	5951100	4.47563700	1.06739000
Н	1.87	/515000	4.76268200	0.03668800
С	1.33	002500	2.10954500	2.23026500
Η	1.82	2722100	2.57953000	3.08796200
Η	1.56	5000600	1.03605700	2.25255600
Н	0.24	899400	2.23601100	2.36414800
Η	2.88	3509200	2.60783500	0.86228600
0	1.37	/557100	-1.78082600	-0.78033300
С	1.27	897500 -	-2.63562300	-1.94270700
Η	0.91	236000	-2.03793700	-2.78689900
Η	0.54	989700	-3.42514100	-1.73372100
С	2.69	365600 -	-3.16324700	-2.20056600
Si	-3.58	669300	0.65335700	-1.06617200
С	-3.94	1583700	1.73079200	-2.57956900
Н	-4.91	1988800	1.48617800	-3.01998900
Η	-3.96	5773900	2.79094900	-2.30428000

Η	2.87101300 -4.08356700 -1.63155900
Η	2.87007600 -3.37974600 -3.25826100
С	3.56659900 -2.02154700 -1.65674700
Η	3.63616900 -1.20796100 -2.38778800
Η	4.58183500 -2.33695500 -1.39858000
С	2.76694100 -1.56891600 -0.43546900
Η	3.00755600 -2.17571100 0.44879900
Η	2.89017700 -0.51043800 -0.18902200
0	-0.44470600 -1.19871100 1.96518000
С	0.13880900 -2.44964300 2.36753400
Η	1.08012700 -2.26547900 2.90785000
Η	0.36051000 - 3.01380600 1.45873200
С	-0.90231200 -3.08931400 3.28231300
Η	-1.69642500 -3.55045100 2.68382800
Η	-0.47795100 -3.85335100 3.94070900
С	-1.43529600 -1.86199700 4.04118000
Η	-0.77897000 -1.62602700 4.88641400
Η	-2.44639100 -2.00414100 4.43329100
С	-1.37333100 -0.74667400 2.98206000
Н	-2.33715800 -0.57783800 2.49164700
Н	-1.01761800 0.20525500 3.38933900
0	-2.03177400 0.96590200 -0.49066900
С	-1.59707700 2.37416600 -0.22308700
С	-2.20406600 2.93327700 0.98088500
С	-2.52109200 4.22518900 1.21761500
Η	-2.45682900 4.98178000 0.43766500
Η	-2.81807800 4.56875900 2.20459200
Η	-2.31045900 2.22731800 1.81119800
Η	-1.80203800 2.97377500 -1.11934500

Η	-3.18402100 1.60787200 -3.35832400
С	-3.60986000 -1.21005400 -1.55623800
С	-2.53309400 -1.49758400 -2.62294400
Η	-1.52547600 -1.25720400 -2.26354800
Η	-2.54080500 -2.56177700 -2.90448300
Η	-2.69936300 -0.91688200 -3.53833000
С	-4.99173700 -1.57734000 -2.14123500
Η	-5.02048900 -2.64183600 -2.41815300
Η	-5.80183800 -1.40827500 -1.42135900
Η	-5.22275600 -1.00118300 -3.04526100
С	-3.35002500 -2.10076600 -0.32240300
Η	-4.12843500 -1.97174500 0.43979000
Η	-3.35047200 -3.16392800 -0.60913600
Η	-2.38465800 -1.88637800 0.15189400
С	-4.89784700 0.99862000 0.25320600
Η	-5.91031600 0.84313700 -0.13997700
Η	-4.77929200 0.35474300 1.13290000
Η	-4.82149900 2.03692600 0.59270100
Η	-0.12961200 2.28374600 -0.16058100

Table S-19. Geometric coordinates and thermally corrected MP2 energies for pro-Z $[A(THF)_3 \cdot (allyloxy-tert-butyldimethylsilane)]^{\ddagger}$.



G = -1870.485032 Hartree $G_{MP2} = -1864.62615$ Hartree

Ato	om	Х	Y	Ζ
Na	0.	00000000	0.00000000	0.00000000
Ν	-0.2	22669800	-2.06796000	-1.20570800
С	-0.4	12094400	-1.98669100	-2.65846700
С	-1.8	35971700	-2.25647200	-3.13636600
Η	-1.9	93771600	-2.10788600	-4.22118600
Η	-2.5	56197400	-1.56711300	-2.64817500
Η	-2.1	18924800	-3.27833600	-2.92276000
С	0.0	0918200	-0.59318000	-3.13703300
Η	-0.0	00555700	-0.52312900	-4.23219000
Η	1.0	02182100	-0.35769900	-2.79697600
Η	-0.6	67668900	0.17125700	-2.74284300
Η	0.2	22687800	-2.71316800	-3.19187900
С	-0.7	73031100	-3.30679200	-0.59455100
С	-0.1	2348800	-4.61189600	-1.16012900
Η	-0.5	57837400	-5.49001000	-0.68349300
Η	0.9	95540000	-4.65085800	-0.97819600
Η	-0.2	28594000	-4.70341500	-2.23984400
С	-0.5	51812100	-3.25861000	0.92603700
Η	-0.9	94126800	-4.15008100	1.40678900
Η	-1.(01084500	-2.37950700	1.36226500
Η	0.5	54846500	-3.21456400	1.17473700
Η	-1.8	82240400	-3.37894900	-0.75254100
0	0.2	2166700	0.54629600	2.32668200
С	-0.8	84231100	0.53522000	3.28938700
Η	-1.7	77809800	0.47845100	2.73133100
Η	-0.8	82138000	1.47680100	3.86116500
С	-0.5	55188800	-0.66935700	4.20390200
Si	3.8	1981900	0.58642700	-0.67693100
С	3.7	0754700	0.98177200	-2.53171100
Η	4.3	39896900	1.78489100	-2.81314400
Η	3.9	94303200	0.11211100	-3.15554300

Η	-0.85056900 -0.46894200 5.23748000
Η	-1.09959200 -1.55350300 3.86478900
С	0.98066400 -0.88582800 4.05320600
Η	1.19515900 -1.87591000 3.64244500
Η	1.51296400 -0.79515200 5.00487400
С	1.41150700 0.20887700 3.06038800
Η	1.77356200 1.10541300 3.58739700
Н	2.15814000 -0.10857100 2.33235600
0	-0.10688200 2.43050300 -0.54664300
С	-0.28779900 3.39142900 0.52078300
Η	0.27294700 3.03635500 1.38790100
Η	-1.35341700 3.42644300 0.78459400
С	0.19051100 4.73243000 -0.03868100
Η	-0.29851500 5.58448400 0.44313400
Η	1.27306100 4.83879100 0.09490600
С	-0.15427900 4.59453600 -1.53021100
Н	0.42506700 5.26216400 -2.17474300
Н	-1.21844700 4.80040000 -1.69731800
С	0.15643900 3.11856300 -1.79040600
Н	-0.46440100 2.66893100 -2.57126300
Н	1.21019200 2.97178600 -2.05639500
0	2.58521300 -0.49011300 -0.26933800
С	2.50901300 -1.84018200 -0.85300000
С	3.05587000 -2.89236900 -0.01641400
С	3.29923300 -2.92384900 1.31497200
Η	3.19239300 -2.04270800 1.93964600
Η	3.62180400 - 3.83612200 1.80821100
Η	3.19037400 - 3.82796300 - 0.56809800
Η	2.95050900 -1.83650800 -1.85872400

Η	2.69507300 1.30609500 -2.79840800
С	3.39395700 2.14630000 0.30842500
Η	3.56298700 2.00807200 1.38231500
Η	3.98828800 3.00905800 -0.01490200
Η	2.33558100 2.38673200 0.16583400
С	5.63463400 0.05620200 -0.27866900
С	5.83215800 -0.20379400 1.22952400
Η	5.23316600 -1.05346300 1.57012300
Η	6.88767300 -0.43532700 1.43778900
Η	5.56935000 0.67170900 1.83741400
С	6.56551200 1.22123700 -0.69600900
Η	7.60979200 0.96242300 -0.46766400
Η	6.51418800 1.43235800 -1.77088400
Η	6.33938100 2.15080800 -0.15876300
С	6.05747400 -1.20600400 -1.06058500
Η	5.47576300 -2.08006800 -0.75451100
Η	5.94198400 -1.07771400 -2.14419100
Η	7.11921000 -1.42720100 -0.87225400
Ο	-2.60192100 0.37804500 0.38193300
С	-3.33298500 1.45353500 -0.22547700
Η	-2.61320900 2.06812500 -0.76975300
Η	-3.81120900 2.07038700 0.55272100
С	-4.38269800 0.78105700 -1.10735900
Η	-5.22884900 1.43627900 -1.33708700
Η	-3.93026800 0.45729500 -2.05149900
С	-4.77106500 -0.43347600 -0.24791700
Η	-5.16791400 -1.26599500 -0.83600000
Η	-5.53345800 -0.14879400 0.48649600
С	-3.44790300 -0.79633100 0.45082300
Η	-3.59410600 -1.06976900 1.50294400
Η	-2.91727300 -1.60733800 -0.05252800
Η	1.08721900 -2.04639600 -1.04236700

Table S-20. Geometric coordinates and thermally corrected MP2 energies for pro- $[A(THF)_3 \cdot (allyloxy-tert-butyldimethylsilane)]^{\ddagger}$.



G = -1870.484508 Hartree $G_{MP2} = -1864.62501$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0	00000000	0.00000000	0.00000000
Ν	-0.2	20162000	-2.14573100	-1.07431100
С	-0.4	0993100	-2.15307700	-2.52796400
С	-1.8	35352700	-2.46050700	-2.96474400
Η	-1.9	95092000	-2.38139600	-4.05522100
Η	-2.5	55133400	-1.74516500	-2.50822300
Η	-2.1	7299100	-3.46879400	-2.68177000
С	0.0	0580800	-0.78896600	-3.09744500
Η	-0.0)1943700	-0.79146400	-4.19456800
Н	1.0	1959400	-0.52199300	-2.78323000
Η	-0.6	58303600	-0.00473600	-2.75030700
Н	0.2	3611600	-2.90719300	-3.02080200
С	-0.6	64064500	-3.37265800	-0.39118400
С	0.0	2167500	-4.67753000	-0.88802200
Η	-0.3	38114800	-5.54388900	-0.34672100
Η	1.1	0610000	-4.65039200	-0.72933200
Η	-0.1	6281300	-4.84660800	-1.95492700
С	-0.4	0404900	-3.22388200	1.11983500
Η	-0.7	78223500	-4.09833000	1.66427600
Η	-0.9)1258600	-2.33487600	1.51374800
Η	0.6	6532500	-3.13226900	1.33690600
Η	-1.7	/2991400	-3.50194100	-0.52739100
0	0.2	7416200	0.61823700	2.29480900
С	-0.7	'5736800	0.77402200	3.27944900
Η	-1.7	71140100	0.72995200	2.75209700
Η	-0.6	5396400	1.76115900	3.75729900
С	-0.5	52529600	-0.35637200	4.30182800
Si	3.7	6950400	0.69024500 -	0.64303800
С	3.8	6500700	0.92204500 ·	-2.52354400
Η	4.6	64224100	1.64358300	-2.80358000

Η	-0.74408900 -0.02360400 5.32113800
Η	-1.17031200 -1.21378200 4.08838900
С	0.97052200 -0.73344300 4.10449000
Η	1.06388000 -1.76484500 3.75337000
Η	1.55390100 -0.64040800 5.02536000
С	1.45624600 0.24863700 3.02414700
Η	1.89589900 1.15094400 3.47718700
Η	2.16033600 -0.17123200 2.30540100
Ο	-0.26186300 2.36685400 -0.73986800
С	-0.46205400 3.38603000 0.26709600
Η	0.10445200 3.09312700 1.15422800
Η	-1.52871000 3.42102900 0.52686300
С	-0.00144200 4.69806000 -0.36958900
Η	-0.50076900 5.57044500 0.06278000
Η	1.07958600 4.82459700 -0.24388800
С	-0.34692800 4.47015200 -1.84965600
Η	0.22703700 5.10321600 -2.53265600
Η	-1.41245900 4.65929400 -2.02678200
С	-0.02726200 2.98277800 -2.02711400
Η	-0.65872200 2.48454300 -2.76893400
Η	1.02262400 2.82564400 -2.30312100
Ο	2.51881500 -0.37616200 -0.27180000
С	2.51822000 -1.78499500 -0.75428700
С	3.15118700 -2.70199000 0.18067200
С	3.77807400 - 3.86980100 - 0.09360200
Η	3.98293700 -4.18254000 -1.11597800
Η	4.07106800 -4.55881200 0.69349600
Η	2.99060300 -2.45529400 1.23528700
Η	2.95474500 -1.82350600 -1.76144300

Η	4.08563600 -0.01748000 -3.04183800
Η	2.90967700 1.29383100 -2.91440400
С	3.21183100 2.32080700 0.14442600
Η	3.33046900 2.31543800 1.23440400
Η	3.77406500 3.17661300 -0.24749800
Η	2.15105500 2.47656100 -0.07368800
С	5.51735800 0.23450000 0.03445500
С	5.44168900 -0.18888000 1.51624000
Η	4.88454600 -1.12247400 1.63617100
Η	6.45281400 -0.35140600 1.91884200
Η	4.96489100 0.57841300 2.14072600
С	6.40713700 1.49732300 -0.06787000
Η	7.42473600 1.26577200 0.27962100
Η	6.49530700 1.86328900 -1.09878000
Η	6.03306900 2.32249400 0.54989500
С	6.18764700 -0.89537700 -0.77735300
Η	5.63392000 -1.83685800 -0.70471500
Η	6.28563600 -0.63249400 -1.83800500
Η	7.20367200 -1.07953000 -0.39581200
Ο	-2.61978600 0.30906400 0.50089300
С	-3.39371800 1.27980000 -0.22765900
Η	-2.72322600 1.77413500 -0.93426600
Η	-3.78631700 2.03367400 0.47248300
С	-4.53342200 0.50075700 -0.88411300
Η	-5.40299400 1.12825900 -1.10299100
Η	-4.19292000 0.04468400 -1.82078900
С	-4.81030800 -0.58248900 0.16933300
Η	-5.30953100 -1.46654700 -0.23803600
Η	-5.43799200 -0.17893200 0.97295100
С	-3.40107300 -0.89713000 0.68529400
Н	-3.38407700 -1.16400600 1.74792400
Η	-2.92418900 -1.69869000 0.11391000
Η	1.07936400 -2.06226400 -0.91498600

Table S-21. Geometric coordinates and thermally corrected MP2 energies for
allyloxytriisopropylsilane.



G = -837.366923 Hartree $G_{MP2} = -834.8750592$ Hartree

Ate	om	Х	Y	Ζ
С	0.0	0000000	0 00000000	0 00000000
Н	-0	14497500	0.40850000	1 00933000
Н	0.0)4571700	0.85558200	-0.69123900
С	-1.1	6098100	-0.88232700	-0.36627100
Η	-1.(03415900	-1.44960500	-1.28801500
С	-2.2	28862300	-0.97906500	0.33789900
Η	-2.4	43453400	-0.42527300	1.26363400
Η	-3.	11425200	-1.60765600	0.01543100
0	1.2	21846000	-0.72650100	-0.11490300
Si	2.5	2198800	-0.81371000	0.94471300
С	3.6	52804500	-2.15119100	0.14728700
Η	3.9	98183300	-1.67897200	-0.78182000
С	4.8	86992400	-2.49269500	0.99494500
Η	4.5	59042300	-2.99908800	1.92765800
Η	5.5	54093200	-3.17105600	0.45092700
Η	5.4	45487600	-1.60510300	1.26636000
С	2.8	8166200	-3.43391900	-0.26585200
Η	1.9	98192200	-3.20865500	-0.84633400
Η	3.5	52866800	-4.07851700	-0.87689600
Η	2.5	57772300	-4.02411800	0.60700000
С	1.9	8258100	-1.31814600	2.71929200
Η	2.9	0159600	-1.74615800	3.15382300
С	1.5	5871300	-0.16201700	3.64706300
Η	0.6	53521300	0.32047400	3.30269000
Η	1.3	36279200	-0.53448100	4.66198600
Η	2.3	32858200	0.61268700	3.72530300
С	0.9	01059600	-2.42756800	2.73202300

Η	1.23077200 -3.32090700 2.18684600
Η	0.68332100 -2.73392900 3.76259500
Η	-0.02621600 -2.09044200 2.27403800
С	3.47109300 0.85823300 0.98917700
Η	4.08028200 0.83759000 1.90788800
С	4.43402100 1.03913600 -0.20167300
Н	4.94281300 2.01129600 -0.14498300
Η	5.20863000 0.26629100 -0.23529400
Η	3.89611300 1.01110600 -1.15785400
С	2.54178000 2.08773600 1.07786700
Η	3.12878800 3.00900300 1.19499000
Η	1.95119500 2.19907500 0.16063100
Н	1.84228000 2.03787100 1.91864600

Table S-22. Geometric coordinates and thermally corrected MP2 energies for pro-Z μ -allyl $[A(THF)_3 \cdot (allyloxytriisopropylsilane)]^{\ddagger}$.



G = -1988.359903 Hartree $G_{MP2} = -1982.055175$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0	0000000	0.00000000	0.00000000
Ν	-1.0	6450500	2.18843700	-0.07206800
С	-1.6	5303500	2.67678500	-1.32825800
С	-0.8	0782200	3.71822400	-2.08673100
Η	-1.2	9845800	3.99002300	-3.03004100
Н	0.1	8347200	3.31266400	-2.32865000
Н	-0.6	6282700	4.64296900	-1.51842000
С	-1.9	2227200	1.48061000	-2.25011600
Н	-2.4	5682400	1.79215000	-3.15711100
Η	-2.5	3238200	0.73415200	-1.73349500
Η	-0.9	7554900	1.01641900	-2.56388600
Н	-2.6	3692400	3.15525400	-1.14603200
С	-0.6	3768300	3.25622500	0.84362800
С	-1.7	6423500	4.20514300	1.31608000
Η	-1.3	6478800	4.99794300	1.96238000
Н	-2.5	2445600	3.65683500	1.88451500
Η	-2.2	6671800	4.69056500	0.47250100
С	0.0	5795600	2.64172800	2.06832800
Н	0.4	3418900	3.42310300	2.74073600
Η	0.9	0864600	2.01692100	1.76703400
Н	-0.6	3525300	2.01546200	2.64198500
Н	0.1	1993900	3.89403000	0.35205800
0	2.1	1918100	0.44757800	-1.24252000
С	2.4	2403900 -	-0.12520600	-2.52471100
Н	1.6	7044900 -	-0.89095400	-2.71793300
Н	3.4	1989900 -	-0.59649600	-2.49833900
С	2.4	0340200	1.05691900 -	-3.49061000
Η	-2.4	8094000	-4.09027300	-2.47448200
Η	-0.8	8197700	-4.05599200	-3.23953500
С	-1.3	8499500	-2.21103700	-2.20254800
Η	-1.3	0417300	-1.67547200	-3.15323500

Η	2.94910200 0.85932800 -4.41856600
Η	1.36900600 1.31231100 -3.74517000
С	3.04318500 2.16922400 -2.63912700
Η	2.70494600 3.16897100 -2.92613100
Η	4.13422600 2.14524400 -2.73835700
С	2.61911800 1.80513300 -1.20039100
Η	3.46086000 1.85773000 -0.49772900
Η	1.80429400 2.42941200 -0.82496400
0	1.52787900 -0.92269000 1.68086600
С	2.93640900 -1.13898900 1.51649400
Н	3.20898000 -0.73148800 0.54305000
Η	3.14170400 -2.22140200 1.52912600
С	3.59788600 -0.44256500 2.72031800
Η	4.48781200 -0.98215200 3.05819400
Η	3.90704200 0.57365300 2.45762800
С	2.47156200 -0.41348400 3.79170500
Η	2.22941500 0.61579200 4.07115600
Η	2.74582100 -0.94961100 4.70516400
С	1.27647400 -1.08059400 3.08410800
Η	1.22088600 -2.15326500 3.32485900
Η	0.31076800 -0.62238800 3.29971700
0	-0.18652800 -1.94592500 -1.43218200
С	0.32567900 -3.17790300 -0.87629000
Η	0.57976000 -2.99525400 0.17145600
Η	1.23997600 -3.45538500 -1.42026800
С	-0.77791200 -4.21982500 -1.07066100
Η	-0.38340600 -5.23790800 -1.14472300
Η	-1.48360600 -4.18425400 -0.23384500
С	-1.45360400 -3.73174900 -2.36148000

H -2.24660400 -1.81855800 -1.65130500 C -3.07793300 0.70374900 1.16163000 -2.36632900 -0.30758700 1.91810500 С С -2.09337600 -1.59453100 1.57490800 H -2.56586900 -2.05034300 0.71006800 H -1.47938600 -2.23336500 2.20365500 H -1.90498700 0.08265500 2.83078800 H -3.64359200 1.38633400 1.80897700 O -3.89622400 0.15563600 0.10571500 Si -5.57171300 0.13076900 0.01826300 C -6.28957100 -0.25932600 1.75429800 H -5.74943700 0.43135700 2.42093200 C -7.79390500 0.02226800 1.93814200 Н -8.10232200 -0.19142000 2.97158800 H -8.05138500 1.06691900 1.73287900 H -8.41445200 -0.60374600 1.28520100 С -5.94409200 -1.68743000 2.22455000 H -4.86751600 -1.88092900 2.18225400 H -6.27186400 -1.84323400 3.26248700 H -6.44837500 -2.44678000 1.61299000 С -5.90830300 -1.27014100 -1.24897400 H -5.27956000 -2.09393600 -0.87530300 С -7.35529700 -1.79563100 -1.30795800 H -8.05855800 -1.02859900 -1.65290700 H -7.43375700 -2.63933100 -2.00872600 H -7.70742500 -2.14897700 -0.33311700 С -5.40431200 -0.92084300 -2.66328200 H -4.35881700 -0.59505200 -2.65668600 H -5.48278700 -1.78856700 -3.33463500 H -5.99575200 -0.11423300 -3.11462500 С -6.13911400 1.84135500 -0.64762400 H -5.44429100 2.01907600 -1.48394500 C -7.56713200 1.90703600 -1.22363700 H -7.77440200 2.90706900 -1.63086100 H -7.71895300 1.18845200 -2.03639700 H -8.33107000 1.70813500 -0.46249300 C -5.91367600 2.98085800 0.36662000 H -4.87399600 3.02531400 0.70692700 H -6.15441000 3.95570600 -0.08096700 H -6.55096900 2.86922500 1.25318100 H -2.06055800 1.50717800 0.55105000

Table S-23. Geometric coordinates and thermally corrected MP2 energies for pro- $E \mu$ -allyl $[A(THF)_3 \cdot (allyloxytriisopropylsilane)]^{\ddagger}$.



G = -1988.358474 Hartree $G_{MP2} = -1982.051409$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0)0000000	0.00000000	0.00000000
Ν	1.4	6349800	-1.93113800	-0.13205900
С	1.4	2315300 -	-2.72814500	-1.36366800
С	0.5	3683900 -	-3.98571700	-1.28854100
Η	0.5	3001300	-4.51191600	-2.25187800
Η	-0.4	9851400	-3.71041100	-1.04575500
Η	0.8	8350300	-4.69520400	-0.53034000
С	0.9	3251500 -	-1.84174700	-2.51780800
Η	1.0	0023200	-2.36568600	-3.47977100
Н	1.5	2410200	-0.92314700	-2.59099100
Н	-0.1	2049700	-1.56238100	-2.36547500
Η	2.4	3689100	-3.07699800	-1.64808100
С	1.8	3554000 -	-2.69092600	1.07194800
С	3.1	8672300 -	-3.43497000	0.98727600
Н	3.3	9235000	-3.97229500	1.92261300
Η	3.9	9783800	-2.72018700	0.81157700
Н	3.1	9648500	-4.17468400	0.17870500
С	1.8	6070900 ·	-1.73644900	2.27580900
Н	2.0	7401700	-2.27990000	3.20497900
Η	0.8	9613200	-1.22570900	2.39930500
Η	2.6	4162700	-0.97885400	2.14224600
Η	1.0	6640500	-3.45712400	1.28938000
0	-2.1	3747300	-1.10162300	0.71465300
С	-3.3	7143600	-1.03697700	-0.01502400
Η	-3.2	3895700	-0.28563100	-0.79544200
Н	-4.1	9066600	-0.72645100	0.65416900
С	-3.5	8921500	-2.46143500	-0.52098800
Η	-1.3	6978600	2.95459500	-4.37553100
Η	-2.9	4462800	2.62559800	-3.63242400
С	-1.3	8104600	1.25613500	-2.98578700
Н	-1.9	7908100	0.42955700	-3.38382400

Η	-4.62972000 -2.66462700 -0.79305600
Η	-2.96138700 -2.64283400 -1.40042000
С	-3.10341500 -3.30254600 0.67641700
Η	-2.67935200 -4.26240900 0.36864500
Η	-3.93305700 -3.51016500 1.36097000
С	-2.05041500 -2.39576400 1.35480500
Η	-2.24437000 -2.28132500 2.42950000
Η	-1.02626300 -2.74595300 1.21489300
0	-0.33149800 1.47722300 1.91415300
С	-1.52419300 1.47838400 2.72204900
Η	-2.09728100 0.59024200 2.45015300
Η	-2.12322800 2.37431700 2.49462600
С	-1.02523300 1.51100700 4.16451300
Η	-1.77971400 1.87787500 4.86724900
Η	-0.71740700 0.50760300 4.48100800
С	0.19425000 2.43963700 4.04286100
Η	0.93155400 2.29207300 4.83722500
Η	-0.12486900 3.48798600 4.07240300
С	0.75629300 2.08158000 2.65709400
Η	1.12096500 2.95529100 2.10641000
Η	1.56532500 1.34722600 2.71761600
0	-1.49850700 1.25429200 -1.54391200
С	-1.83269100 2.58014100 -1.07684300
Η	-1.24202100 2.77491100 -0.17806800
Η	-2.90022700 2.60577800 -0.81313700
С	-1.52753300 3.51966200 -2.24311200
Η	-2.11467500 4.44227000 -2.20446600
Η	-0.46491400 3.78621100 -2.24719100
С	-1.86284900 2.63397300 -3.45318200

Н -0.33072700 1.08433400 -3.24713200 С 3.46652300 -0.07120600 -0.58408700 С 2.81721500 1.19144400 -0.31039600 С 1.92838300 1.87677800 -1.08582300 Η 1.76806000 1.62433300 -2.13283600 Η 1.43272800 2.77160900 -0.71860900 Η 2.97260400 1.55367200 0.70969500 0 4.58947000 -0.35755500 0.27616500 Si 6.20723300 0.00106300 0.01338400 С 6.69989700 -0.69149800 -1.70642000 Η 5.97090500 -0.23336600 -2.39468400 С 8.10079700 -0.30239800 -2.21811000 Η 8.26939400 -0.70147800 -3.22865700 Η 8.23862000 0.78319800 -2.27063400 Η 8.89720600 -0.70360000 -1.58003700 С 6.49741100 -2.21787400 -1.78632800 Η 5.48489500 -2.51219800 -1.48991200 Η 6.66776600 -2.58353800 -2.80901200 Η 7.20010200 -2.75083700 -1.13285000 С 7.02042800 -0.95474700 1.46007000 Η 6.56084100 -1.95172100 1.37483400 С 8.54646500 -1.15174400 1.39851200 Η 9.09041000 -0.20079500 1.44049800 8.89312600 -1.75670500 2.24865400 Η Η 8.86001900 - 1.66872800 0.48478100 С 6.59239800 -0.38458100 2.82685600 Η 5.50272300 -0.30863300 2.90318100 Η 6.94332900 -1.02584400 3.64805900 Η 7.01120000 0.61550600 2.99885600 С 6.48900400 1.90201800 0.12111200 Η 5.84301000 2.20924600 0.95915700 С 7.92482800 2.33778700 0.47623000 Η 7.98854100 3.43306300 0.55004000 Η 8.25932800 1.92718000 1.43474400 Η 8.65018900 2.02732200 -0.28554400 С 5.99627900 2.65906600 -1.12941000 4.94434600 2.45096500 -1.34870500 Η Η 6.09666200 3.74542300 -0.98965300 Η 6.58441000 2.39650600 -2.01778000 Η 2.46580800 -1.04442400 -0.32571400 Η 3.71884200 -0.23236700 -1.64075300

Table S-24. Geometric coordinates and thermally corrected MP2 energies for pro-Z κ -O $[A(THF)_2 \cdot (allyloxytriisopropylsilane)]^{\ddagger}$.



G = -1756.012449 Hartree $G_{MP2} = -1750.479449$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0	0000000	0.00000000	0.00000000
Ν	0.8'	7448200	-2.13517900	-0.55844800
С	1.08	3164000	-3.22808500	0.39771000
С	2.55	5508900	-3.47444300	0.77677400
Η	2.62	2940600	-4.25271200	1.54726300
Η	3.00	0784300	-2.55592700	1.17469000
Н	3.1.	5908500	-3.80190200	-0.07552700
С	0.27	7704400	-2.94149900	1.67234400
Н	0.3	5523300	-3.77020600	2.38734100
Η	-0.7	8129600	-2.79065600	1.44242100
Н	0.65	5261300	-2.03294600	2.16365300
Н	0.70	0015800	-4.19130000	0.00012100
С	1.6	1956400	-2.29133100	-1.81652000
С	1.22	2438000	-3.52889300	-2.65364100
Н	1.84	4647600	-3.60820200	-3.55469800
Η	0.1	7711800	-3.46781300	-2.97026300
Н	1.34	4941300	-4.45775700	-2.08627600
С	1.47	7800600	-1.02030300	-2.66619600
Н	2.04	4646000	-1.10581300	-3.60100400
Н	1.80	6690800	-0.14853100	-2.12178400
Η	0.42	2969300	-0.83110600	-2.92955600
Н	2.6	9698200	-2.38965200	-1.59784300
0	1.20	6476600	0.92781200	1.79440800
С	0.98	3433300	1.09877300	3.20225400
Η	0.4	1744500	0.22826000	3.55386900
Η	0.30	6302500	1.99205500	3.32662600
С	2.34	4590000	1.20429200	3.89694400
Si	-3.68	3351700	-0.38099500	0.60023800
С	-4.74	4355300	-1.96894100	0.78487700
Η	-4.0	2325300	-2.74072000	1.09970700

Η	2.68691900 2.24615200 3.92065200
Η	2.31694700 0.83429300 4.92603500
С	3.24465600 0.36792300 2.97264000
Η	3.11957500 -0.70115600 3.17893600
Η	4.30722700 0.61192000 3.06381800
С	2.68168900 0.70947100 1.59343700
Η	3.12894700 1.63041800 1.19310200
Η	2.79328800 -0.09242500 0.85782200
0	0.06173700 2.12957600 -1.03265500
С	0.10521300 3.36367700 -0.28055600
Η	0.87586500 3.27008700 0.48975100
Η	-0.86639900 3.50065800 0.21197400
С	0.37575400 4.47530600 -1.30016200
Η	-0.06466800 5.43082400 -1.00009400
Η	1.45414100 4.62525800 -1.43084900
С	-0.24096200 3.89332800 -2.58196600
Η	0.15827700 4.34019100 -3.49735000
Η	-1.32877700 4.02827600 -2.58012200
С	0.11255300 2.41282400 -2.45123600
Η	-0.58239000 1.73533700 -2.95305100
Η	1.12733400 2.21076400 -2.81948900
0	-2.17910200 -0.77974200 -0.08128600
С	-1.87590500 -1.92324600 -0.95870500
С	-2.21742700 -1.73112400 -2.35939300
С	-2.45545100 -0.59690300 -3.05794100
Η	-2.52628000 0.37084100 -2.57184200
Η	-2.60644200 -0.61911000 -4.13330000
Η	-2.16972900 -2.66576800 -2.92627900
Η	-2.32318100 -2.83174300 -0.53807700
С	-5.83052200 -1.89106600 1.87665100
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Η	-6.37708700 -2.84231200 1.94155400
Η	-5.41656300 -1.68641600 2.86931000
Η	-6.57086500 -1.11168500 1.66094600
С	-5.36919100 -2.44293900 -0.54474100
Η	-4.63322200 -2.51626900 -1.35036300
Η	-5.83567300 -3.43014900 -0.41977700
Η	-6.15621700 -1.75763100 -0.88239200
С	-3.15741500 0.36986800 2.29087900
Η	-2.33138300 1.04865800 2.01552900
С	-4.21388400 1.22619800 3.01606300
Η	-3.80586500 1.63371800 3.95219500
Η	-4.54672600 2.07471600 2.40988600
Η	-5.10345100 0.64365500 3.28287600
С	-2.56999700 -0.68243600 3.25148500
Η	-1.77460800 -1.26759400 2.78068700
Η	-2.15167500 -0.20446200 4.14930600
Η	-3.33589400 -1.38721600 3.59621400
С	-4.54817300 0.93153300 -0.50222600
Η	-4.54939500 0.46394200 -1.49676600
С	-6.01353100 1.26177700 -0.15321700
Η	-6.42981000 1.97160000 -0.88210100
Η	-6.65438700 0.37403200 -0.16619400
Η	-6.11305100 1.72509100 0.83529000
С	-3.71849900 2.22693300 -0.61012500
Η	-3.71187000 2.77989900 0.33860800
Η	-2.67648000 2.02540700 -0.88299900
Η	-4.13985300 2.90088700 -1.36987400
Η	-0.44923900 -2.11564600 -0.82782600

Table S-25. Geometric coordinates and thermally corrected MP2 energies for pro- $E \kappa$ -O $[A(THF)_2 \cdot (allyloxytriisopropylsilane)]^{\ddagger}$.



G = -1756.011312 Hartree $G_{MP2} = -1750.477393$ Hartree

Ato	om	Х	Y	Ζ
Na	0.0	0000000	0.00000000	0.00000000
Ν	0.7'	7602100	2.23156000	0.21075300
С	0.90	0250800	3.13123400	-0.94285000
С	2.3	5338700	3.49068200	-1.31325700
Н	2.3	7757300	4.09809000	-2.22711500
Н	2.9	4215600	2.58058900	-1.49307700
Η	2.8	5664400	4.06411800	-0.52813200
С	0.2	1455200	2.48964000	-2.15696600
Н	0.2	1181400	3.17088500	-3.01715400
Η	-0.8	2242800	2.22507500	-1.92938500
Η	0.74	4347500	1.57282600	-2.45424500
Η	0.3	7991600	4.09225400	-0.76055400
С	1.44	4822100	2.70651900	1.42938100
С	0.8	9768400	4.03189800	2.00106400
Н	1.4	7580000	4.34702000	2.87986100
Н	-0.1	5009500	3.91943300	2.30467100
Н	0.9	5365600	4.84242500	1.26565500
С	1.3	7588300	1.61183800	2.50370600
Н	1.8	8979200	1.92154100	3.42223100
Н	1.8	4981200	0.68682800	2.14791400
Η	0.3	3338100	1.39411000	2.76593200
Η	2.5	1964000	2.86729900	1.22067300
0	1.4	6155100	-1.18541400	-1.45923900
С	1.2	7603700	-1.65545800	-2.81438500
Н	0.6	5325800	-0.93134700	-3.35340900
Н	0.74	4546000	-2.61283900	-2.77923500
С	2.6	7772200	-1.75853100	-3.42371800
Si	-3.70	0627800	0.09489600	-0.66598000
С	-4.7	3968800	1.54519000	-1.38533400
Η	-3.9	8944600	2.16450000	-1.90324100

Η	3.11430100 -2.74400100 -3.22248100
Η	2.67154500 -1.60589200 -4.50695500
С	3.44220200 -0.66709700 -2.65820000
Η	3.23102500 0.32064700 -3.08341200
Η	4.52616900 -0.81501700 -2.65932900
С	2.83858000 -0.78006400 -1.25859000
Η	3.34870000 -1.54856900 -0.66107600
Η	2.82954100 0.16214600 -0.70288300
0	0.02070100 -1.82838700 1.51237200
С	0.17495900 -3.17655800 1.02084800
Η	0.84924200 -3.13771100 0.16175300
Η	-0.80344800 -3.55333900 0.68963900
С	0.70246100 - 3.98333100 2.20607500
Η	0.48326900 -5.05177400 2.11916600
Η	1.78842100 -3.86207500 2.29854800
С	-0.01907800 -3.30723400 3.38259200
Η	0.47915400 -3.46065600 4.34427500
Η	-1.04353800 -3.68757100 3.46738700
С	-0.02575600 -1.83307300 2.96047100
Η	-0.92309600 -1.29599700 3.28242900
Η	0.85427000 -1.29638600 3.33231400
0	-2.19388300 0.67042500 -0.16534600
С	-1.96244700 1.94027000 0.57554500
С	-2.43635000 1.89158400 1.95487900
С	-2.89067000 2.91796300 2.70834300
Η	-3.05880300 3.90685900 2.28575500
Η	-3.07384000 2.80746100 3.77345800
Η	-2.30422300 0.92392800 2.45090900
Η	-2.40043800 2.76500900 -0.00053100

-5.79055200 1.12829800 -2.43566600
-6.31170400 2.01286100 -2.82782200
-5.35007900 0.60557900 -3.29091100
-6.55781700 0.47142900 -2.00705100
-5.40583900 2.43447200 -0.31335100
-4.70904200 2.75956300 0.46374700
-5.83988200 3.33121200 -0.77749800
-6.22676400 1.90684700 0.18798700
-3.18445400 -1.18956900 -1.99681900
-2.39421100 -1.76699100 -1.48644500
-4.26306600 -2.19877600 -2.43427500
-3.85513500 -2.91331800 -3.16371500
-4.65100100 -2.78044800 -1.59163200
-5.11660800 -1.70543300 -2.91312300
-2.53017000 -0.52870800 -3.22683500
-1.72547300 0.15794300 -2.94298200
-2.10703200 -1.28667800 -3.90202500
-3.25751900 0.04717700 -3.81140600
-4.56866500 -0.73875000 0.83083200
-4.54825500 0.04579900 1.60223400
-6.04261900 -1.14188100 0.62855500
-6.46977500 -1.52942000 1.56426800
-6.66441500 -0.29733600 0.31401300
-6.15513500 -1.93128700 -0.12433600
-3.75327800 -1.93119900 1.36905000
-3.78293700 -2.78133100 0.67433100
-2.70057700 -1.67024900 1.52500500
-4.15956700 -2.28696000 2.32678900
-0.51369500 2.16019200 0.46404600

Table S-26. Geometric coordinates and thermally corrected MP2 energies for pro- $E \kappa$ -O $[A(THF)_3 \cdot (allyloxytriisopropylsilane)]^{\ddagger}$. The pro-Z isomer does not converge without THF extrusion.



G = -1988.34947 Hartree $G_{MP2} = -1982.049537$ Hartree

Ato	m	Х	Y	Ζ
Na	0.00	000000	0.00000000	0.00000000
Ν	0.60	164300	1.96298100	-1.26448300
С	0.75	387600	1.78999700	-2.71469700
С	2.20	956600	1.83613200	-3.21457900
Η	2.25	381600	1.62528600	-4.29093600
Η	2.81	369700	1.08026400	-2.69437700
Η	2.67	987000	2.81127000	-3.05309300
С	0.13	508100	0.44658400	-3.12343700
Η	0.10	098000	0.33683000	-4.21493000
Н	-0.88	748100	0.34970700	-2.74493200
Н	0.73	267300	-0.38417500	-2.72254600
Н	0.20	382700	2.57393600	-3.27328400
С	1.26	298300	3.16452900	-0.73173600
С	0.83	005700	4.50074400	-1.37588100
Н	1.38	382600	5.33838500	-0.93118800
Н	-0.24	082100	4.67830300	-1.22225500
Н	1.02	670000	4.51724500	-2.45375700
С	1.04	599900	3.23779700	0.78720600
Н	1.60	714700	4.07487000	1.22170400
Η	1.37	647500	2.31436200	1.27770400
Η	-0.01	318000	3.39074200	1.01938300
Η	2.35	466600	3.08185100	-0.88793400
Ο	2.53	022500	-0.85141900	0.43673400
С	3.14	160200 -	-1.94637500	-0.26779400
Η	2.39	845200	-2.36024000	-0.95354900
Η	3.42	733300	-2.72848200	0.45258100
С	4.37	415200 -	-1.36681200	-0.96361200
Η	-1.38	031600	-4.92137000	-2.39916200
Η	0.30	861600	-5.10200300	-1.90963800

Η	5.14295200 -2.12124800 -1.15767500
Η	4.09251800 -0.90900700 -1.91878200
С	4.81376000 -0.28682700 0.03636700
Η	5.42802500 0.49783600 -0.41512300
Η	5.38714700 -0.73719400 0.85572000
С	3.46870000 0.24694100 0.54136400
Η	3.50185200 0.57558700 1.58600900
Η	3.09323000 1.06895500 -0.07526900
0	-0.02665900 -0.02363700 2.38561100
С	0.83629000 -0.71239500 3.30119200
Η	1.65897600 -1.12275000 2.71420500
Η	0.27836900 -1.53571700 3.77429900
С	1.25384800 0.33959000 4.34980100
Η	1.29936800 -0.09568700 5.35272400
Η	2.24345500 0.74762000 4.12328600
С	0.16270800 1.44034400 4.22747400
Η	0.59848200 2.37719000 3.86945000
Η	-0.33682700 1.64629200 5.17885500
С	-0.81472800 0.87188300 3.18599800
Η	-1.62651500 0.30670100 3.66716100
Η	-1.25190300 1.61167200 2.51477100
0	-0.20014400 -2.42384600 -0.65109500
С	-0.16490100 - 3.46383100 0.34170700
Η	-0.56011400 -3.04756700 1.27039900
Η	0.87405000 -3.78509800 0.51113600
С	-1.00086300 -4.59656300 -0.24720600
Η	-0.78553900 - 5.56678200 0.21082800
Η	-2.06732900 -4.38232700 -0.11716300
С	-0.60772200 -4.52749900 -1.73281600

С -0.35715600 -3.02371700 -1.96011100 Η 0.54581500 -2.83736800 -2.55289300 H -1.19847600 -2.52640300 -2.45043900 С -2.14335900 2.05074900 -0.82906700 -2.55723400 3.24688600 -0.10535900 С С -2.93586300 4.43759400 -0.62037000 H -3.09485000 4.57434000 -1.68866100 H -3.07134500 5.31431300 0.00678300 H -2.43453900 3.18525200 0.98129100 H -2.58419600 1.99750100 -1.83431400 O -2.45321300 0.79266800 -0.09259800 Si -3.98416500 0.06033000 -0.06852900 C -5.38128400 1.34303700 -0.38364100 H -4.97196700 2.00791900 -1.15727900 C -6.69271300 0.74765700 -0.93757800 H -7.42284800 1.54699200 -1.12685600 H -6.54713400 0.21314900 -1.88187600 H -7.16213700 0.04933000 -0.23384900 С -5.68629100 2.23172400 0.84000000 H -4.79214900 2.74025900 1.20810400 H -6.41247900 3.01140000 0.57183200 Н -6.12109000 1.65752900 1.66716400 С -4.16463700 -0.70707700 1.68633800 H -3.97161600 0.14850300 2.35318200 C -5.56885800 -1.24169800 2.03518500 H -5.83792800 -2.11404500 1.42883600 H -5.60877200 -1.55951500 3.08696000 H -6.35198500 -0.49102900 1.89307300 C -3.10471400 -1.77709200 2.00591900 H -2.08777000 -1.39238200 1.88431400 H -3.20549800 -2.13051800 3.04329900 H -3.21628400 -2.65769300 1.35985600 C -3.91303200 -1.30311500 -1.42539200 H -2.93133400 -1.76331700 -1.23117200 C -4.96356600 -2.42759000 -1.34542300 H -4.80533900 -3.16445600 -2.14688600 H -4.92181100 -2.96967400 -0.39452400 H -5.98491200 -2.04771400 -1.46100900 C -3.86262200 -0.72387100 -2.85419600 H -3.09104600 0.04448400 -2.96364100 H -3.65098100 -1.51287900 -3.59108200 H -4.81827000 -0.27011800 -3.14218600 H -0.68028000 2.06834300 -1.04886700

III. Full reference (Gaussian):

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