

## ***Supporting Information***

# **On The Nature of Csp<sup>3</sup>)-C(sp<sup>2</sup>) Bond Formation in Nickel-Catalyzed Tertiary Radical Cross-Couplings: A Case Study of Ni/Photoredox Catalytic Cross-Coupling of Alkyl Radicals and Aryl Halides**

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## A. Computational Details and Details on Quasi-Classical Direct-Dynamics Simulations

All geometry optimizations of intermediates and transition states were achieved using spin-unrestricted UB3LYP<sup>1</sup>-D3<sup>2</sup>/def2-SVP<sup>3</sup> method, in THF solvent using the CPCM solvent model<sup>4</sup> with “opt=noeigen” and “guess=mix” keywords as implemented in Gaussian09<sup>5</sup>. Frequency calculations were also conducted at the same level of theory to obtain vibrational frequencies to determine the identity of stationary points as intermediates (no imaginary frequencies) or transition states (only one imaginary frequency), as well as obtaining the thermal corrections to enthalpy ( $H_{\text{correction}}$ ) and free energy ( $G_{\text{correction}}$ ) at the temperature of 298 K. Energies were refined by computing single point energies in implicit solvent with UB3LYP-D3/def2-TZVPP<sup>3</sup> method using larger basis set compared to optimization method. To explore the effect of DMA solvent in Ni-TMHD catalytic system, single point energy was also calculated in implicit DMA solvent as labeled as UB3LYP-D3/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF). For comparison, single point energies were also computed using UM06<sup>6</sup>/def2-TZVPP method using Gaussian 09<sup>5</sup>. Moreover, domain-based local pair-natural orbital coupled-cluster calculations using single and double excitations with perturbative triple excitations (DLPNO-CCSD(T))<sup>7</sup> were performed with def2-TZVPP as basis set and def2-TZVPP/C as auxiliary basis set using ORCA software<sup>8</sup>. This method was used to obtain energy with high accuracy at affordable computational cost compared to DFT calculations.<sup>9</sup> Free energies obtained with each method was calculated generally as  $G_{\text{method-solvent}} = E_{\text{method-solvent}} + G_{\text{correction}}$ . In part E, single point energies obtained directly with each method ( $E_{\text{method-solvent}}$  computed with UB3LYP-D3 and UM06 methods, and  $E^{\text{DLPNO-CCSD(T)-gas}}$  computed with DLPNO-CCSD(T) method) would be shown. All structural figures were generated with CYLview.<sup>10</sup> Distances in structural figures are shown in Å and energies are in kcal/mol.

Molecular dynamics (MD) simulations were performed in the gas phase on the truncated doublet transition states **<sup>2</sup>I'-TS** (shown in Figure S7) and **<sup>2</sup>K-TS** (shown in Figure S20). The truncated **<sup>2</sup>I'-TS** and **<sup>2</sup>K-TS** were located at the UB3LYP-D3/def2-SVP level of theory using Gaussian 09 with HOMO-LUMO mixing for the initial guess in both transition state optimization and dynamics simulation. Quasi-classical direct-dynamics simulations were then initialized within the region of the potential energy surface near **<sup>2</sup>I'-TS** and **<sup>2</sup>K-TS**, adding zero-point energy for each real normal mode in corresponding transition states, plus a Boltzmann sampling of thermal energy available at 298.15 K with a random phase. The trajectories were propagated backward on **<sup>2</sup>I'-TS** and forward on **<sup>2</sup>K-TS** (settings are made on “searchdir” keyword line as shown in Section B), with focus on observing possible radical dissociation or C(sp<sup>2</sup>)-C(sp<sup>3</sup>) bond formation within the lifetime of Ni(III) intermediate. Parameters of radical dissociation concomitant with generation of Ni(II) intermediate were set as **<sup>1</sup>G/<sup>1</sup>G'** ( $C_{\text{Ph}}-C_{t\text{Bu}} > 3.00 \text{ \AA}$  and  $\text{Ni}-C_{t\text{Bu}} > 2.70 \text{ \AA}$ ) for **<sup>2</sup>K-TS** and **<sup>1</sup>G/<sup>1</sup>G'** ( $C_{\text{Ph}}-C_{t\text{Bu}} > 3.00 \text{ \AA}$  and  $\text{Ni}-C_{t\text{Bu}} > 3.50 \text{ \AA}$ ) for **<sup>2</sup>I'-TS** or product formation as **<sup>2</sup>P<sub>B</sub> + tBu-Ph** ( $C_{\text{Ph}}-C_{t\text{Bu}} < 1.70 \text{ \AA}$  and  $\text{Ni}-C_{t\text{Bu}} > 5.00 \text{ \AA}$ ) based on corresponding distances at the transition states. The classical equations of motion were integrated with a velocity-Verlet algorithm using Singleton’s program Progdyn<sup>11</sup>, with the energies and derivatives computed using the UB3LYP-D3/def2-SVP method with Gaussian 09. The step length for integration was 1 fs.

## B. Input Parameters on Progdyn

The configuration file progdyn.conf for trajectory initialization for the doublet reduction elimination transition state  **$^2\Psi^-$ -TS** is given below as an example.

```
/#This is the configuration file for PROGDYN. This file is read by progdynstarterHP and
# the awk programs proggenHP, prog1stpoint, prog2ndpoint, and progdynb.
#The programs won't read anything past the first blank line,
#and this file must end with a blank line.
#The program has a number of default values but they are unlikely to be what you want.
#Do not delete lines - rather, comment out lines for unwanted options.
#The values here are read repeatedly and most can be changed in the middle of running
jobs
#***The keywords are case sensitive. The following keywords should always be
defined:***
#***method, charge, multiplicity, memory, processors, title
#*** method --The following word is copied exactly to the gaussian input file.
method ub3lyp/def2svp
#To do a nonstandard route, make nonstandard 1. For normal calcs, use nonstandard 0 or
else leave it out.
#Then make a file called "nonstandard" containing the nonstandard route with no extra
lines.
nonstandard 0
# NMROptions As is NMR=1 will add a section for an NMR calc at every NMRevery intervals.
If you want to combine the two use nonstandard
#NMRtype 1
#NMRmethod B3LYP/cc-pvtz
#NMRevery 4
#geometry linear
rotationmode 1
#*** method2 --The options here are restricted, unrestricted, and read. restricted is the
default
#If the method is U..., put unrestricted here and the .com files will have in them guess=mix.
#If you put read here, the .com files will contain guess=tcheck, which sometimes makes
things faster, sometimes not.
#The use of read requires a specifically defined checkpoint file name using the keyword
checkpoint.
method2 unrestricted
charge 0
multiplicity 2
processors 32
#*** memory --The following "word" is copied exactly to the gaussian input file
after %mem=.
memory 64gb
#*** killcheck and checkpoint -- You can use a specifically defined checkpoint file name by
putting
```

```

#the name after the keyword checkpoint. This is necessary if you use the read option with
method2.

#Defined checkpoint names are an unnecessary modest hassle and if you do not want to
bother, use killcheck 1
killcheck 1
#checkpoint g09.chk
#*** diagnostics -- 0 prints out nothing extra, 1 (default) prints out extra stuff to a
#file "diagnostics", 2 adds more stuff, 3 adds velocities to a file "vellist"
#4 adds the apparent temperature to vellist, but this is meaningless with quasiclassical
calculations
diagnostics 1
#*** title -- the title keyword must be followed by exactly four words
title j j j Ni-trun-TS-RE-r2-g
#*** initialdis -- 0 (default) turns off displacement of the normal modes, so that all
trajectories start from the same place
# and only the energies and signs of the motion in the modes are randomized
# 1 gives a flat distribution of displacements where all of the possible values are equally
likely
# 2 (recommended) gives a QM-like gaussian distribution of displacements, so that
displacements in the middle are more likely than
# those at the end by 1/e
initialdis 2
#*** timestep -- this is the time between points in the trajectory. Typical values would be
1E-15 or 0.5E-15 or 0.25E-15
timestep 1E-15
#*** scaling -- this lets you scale the gaussian frequencies by a constant
scaling 1.0
temperature 298.15
#*** method3, method4, method5, and method6 -- These keywords let you add extra lines
to the gaussian input file.
#method3 and method4 add lines at the top of the input after the lines defining the method,
and
#this is useful to implement things like the iop for mPW1k
#method5 and method6 add lines after the geometry, after a blank line of course
#only a single term with no spaces can be added, one per method line. Here are some
examples to uncomment if needed
#add the line below with big structures to get it to put out the distance matrix and the
input orientation
#method3 iop(2/9=2000)
#method4 iop(3/124=3)
#method5 eps=32.63
#method6 rsolv=1.329
#*** methodfile -- This keyword lets you add more complicated endings to gaussian input
files
#such as a gen basis set. Put after the keyword the number of lines in a file you create
called

```

```

#methodfile that contains the test you want to add to the end of the gaussian input
methodfile 0
#*** numimag --This tells the program the number of imaginary frequencies in the starting
structure.
#if 0, treats as ground state and direction of all modes is random
#if 1, motion along the reaction coordinate will start out in the direction defined by
searchdir
#if 2, only lowest freq will go direction of searchdir and other imag mode will go in random
direction
numimag 2
#*** searchdir -- This keyword says what direction to follow the mode associated with the
imaginary frequency.
#The choices are "negative" and "positive". Positive moves in the direction defined in the
gaussian frequency calculation
#for the imaginary frequency, while negative moves in the opposite direction. The correct
choice can be made either
#by a careful inspection of the normal modes and standard orientation geometry, or by
trial and error.
searchdir positive
#*** classical -- for quasiclassical dynamics, the default, use 0. for classical dynamics, use
1
#if there are no normal modes and the velocities are to be generated from scratch, use
classical 2
classical 0
#*** DRP, saddlepoint, and maxAtomMove --to run a DRP use 'DRP 1' in the line below,
otherwise leave it at 0 or comment it out
#the treatment of starting saddlepoints is not yet implemented so use saddlepoint no
#if DRP shows oscillations then decrease maxAtomMove
#DRP 1
#saddlepoint no
#maxAtomMove 0.01
#*** cannonball -- The program can "fire" a trajectory from a starting position toward a
particular target, such as toward
#a ts. To use this, make a file cannontraj with numAtom lines and three numbers per line
that defines the vector
#for firing the trajectory, relative to the starting geometry's standard orientation. The
number following cannonball sets
#the extra energy being put into the structure in kcal/mol
#cannonball 10
#*** keepevery --This tells the program how often to write the gaussian output file to file
dyn, after the first two points.
#Use 1 for most dynamics to start with, but use a higher number to save on disk space or
molden loading time.
keepevery 1
#*** highlevel --For ONIOM jobs, the following line states the number of highlevel atoms,

```

```

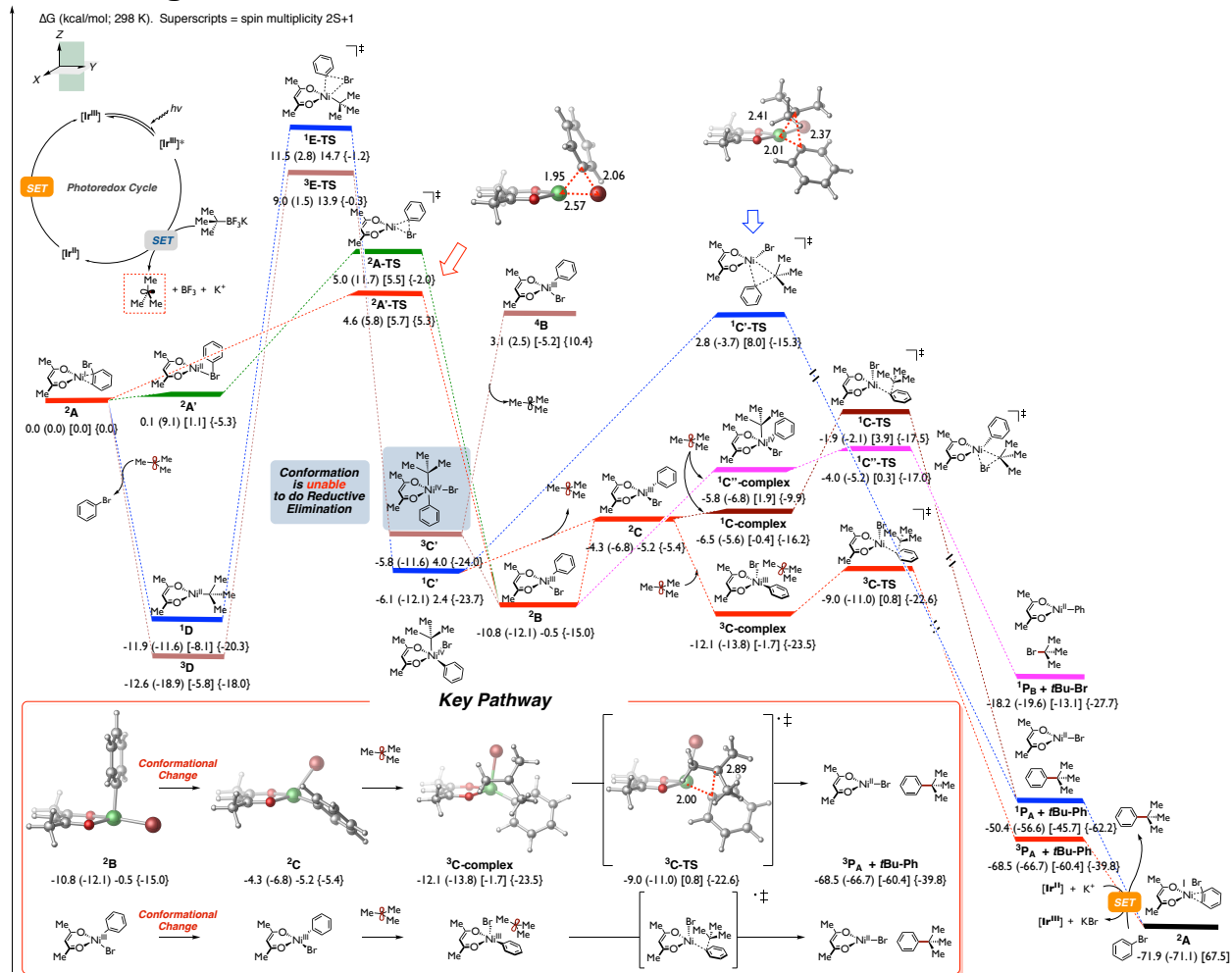
#which must come before the medium level atoms. Use some high value such as 999 if not
using ONIOM
highlevel 999
#*** fixedatom1, fixedatom2, fixedatom3, and fixedatom4 - These fix atoms in space.
#Fixing one atom serves no useful purpose and messes things up, while fixing two atoms
#fixes one distance and fixing three has the effect of fixing three distances, not just two
#in current form fixed atoms only are meant to work with no displacements, that is,
initialdis=0
#fixedatom1 2
#fixedatom2 3
#fixedatom3 19
#*** boxon and boxsize - With boxon 1, a cubic box is set such that atoms that reach the
edge
#are reflected back toward the middle. Useful for dynamics with solvent molecules. This is
a crude
#implementation that is ok for a few thousand femtoseconds but will not conserve energy
long term.
#Set the box size so as to fit the entire initial molecule but not have too much extra room.
#The dimensions of the box are two times the boxsize, e.g. boxsize 7.5 leads to a box that is
15 x 15 x 15 angstroms
boxon 0
boxsize 7.5
#*** displacements -- This keyword lets you set the initialdis of particular modes by using a
series of lines of the format
# displacements NumberOfMode InitialDisForThatMode, as in the example below. You
should be able to do as many of these as you like
# you might consider this for rotations where a straight-line displacement goes wrong at
large displacements
# The choices for InitialDisForThatMode are 0, 1, 2, and 10, where 10 does the same thing
as 0 but is maintained for now because
# a previous version of the program had a bug that made 0 not work.
#displacements 2 0
#displacements 3 0
#*** etolerance --This sets the allowable difference between the desired energy in a
trajectory and the actual
#energy, known after point 1 from the potential energy + the kinetic energy in the initial
velocities.
#The unit is kcal/mol and 1 is a normal value for mid-sized organic systems. For very large
and floppy molecules, a larger value
#may be needed, but the value must stay way below the average thermal energy in the
molecule (not counting zpe).
#If initialdis is not 0 and few trajectories are being rejected, decrease the value.
etolerance 0.5
#*** controlphase --It is sometimes useful to set the phase of particular modes in the
initialization of trajectories.

```

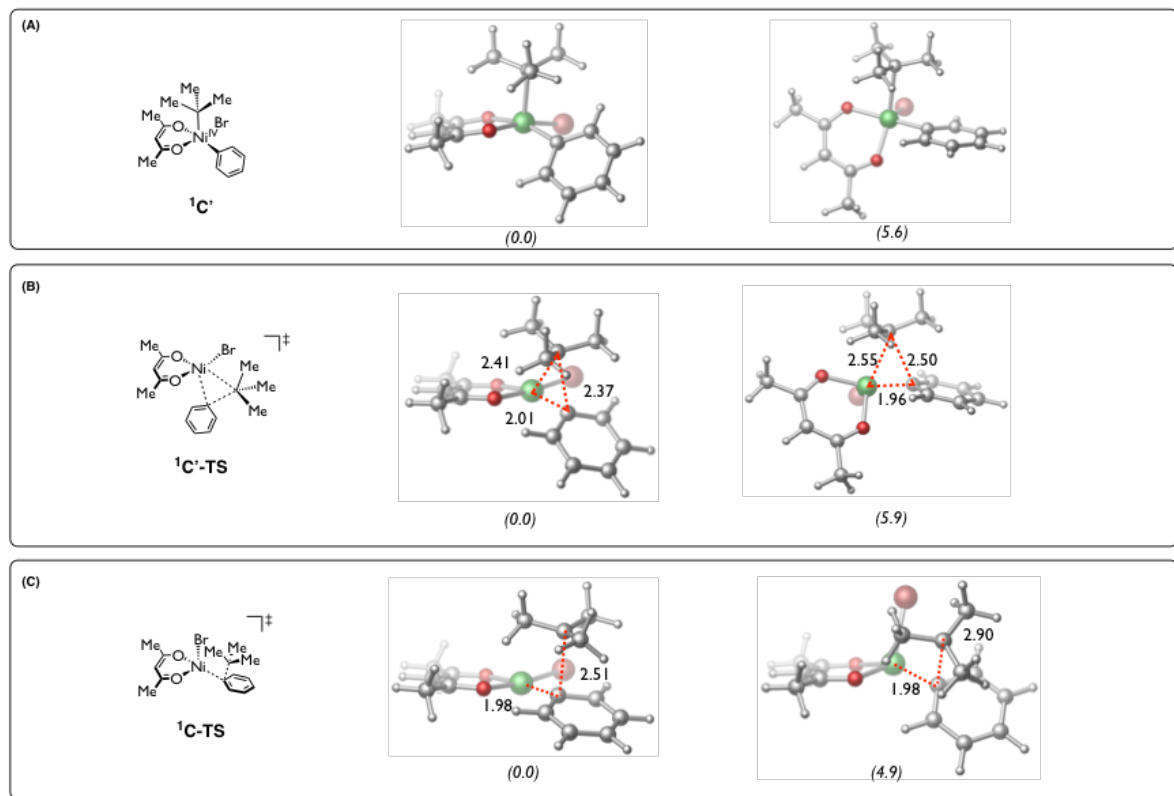
```
#The format is controlphase numberOfModeToControl positive or controlphase
numberOfModeToControl negative.
#controlphase 2 positive
#*** damping -- The damping keyword lets you add or subtract energy from the system at
each point, by multiplying the velocities
#by the damping factor. A damping of 1 has no effect, and since you mostly want to change
the energy slowly, normal values range
#from 0.95 to 1.05. The use of damping lets one do simulated annealing - you add energy
until the structure is moving enough
#to sample the kinds of possibilities you are interested in, then you take away the energy
slowly.
damping 1
#*** reversetraj --This keyword sets the trajectories so that both directions from a
transition state are explored.
reversetraj true
```

```
#updated Aug 9, 2007 to include the possibility of classical dynamics by the keyword
classical
#updated Jan 2008 to include fixed atoms, ONIOM jobs, keepevery, and box size
#update Feb 2008 to include methodfile parameter
# updated Nov 2008 to allow for start without an initial freq calc using classical = 2
# update Aug 2010 to include etolerance, damping controlphase and reversetraj
```

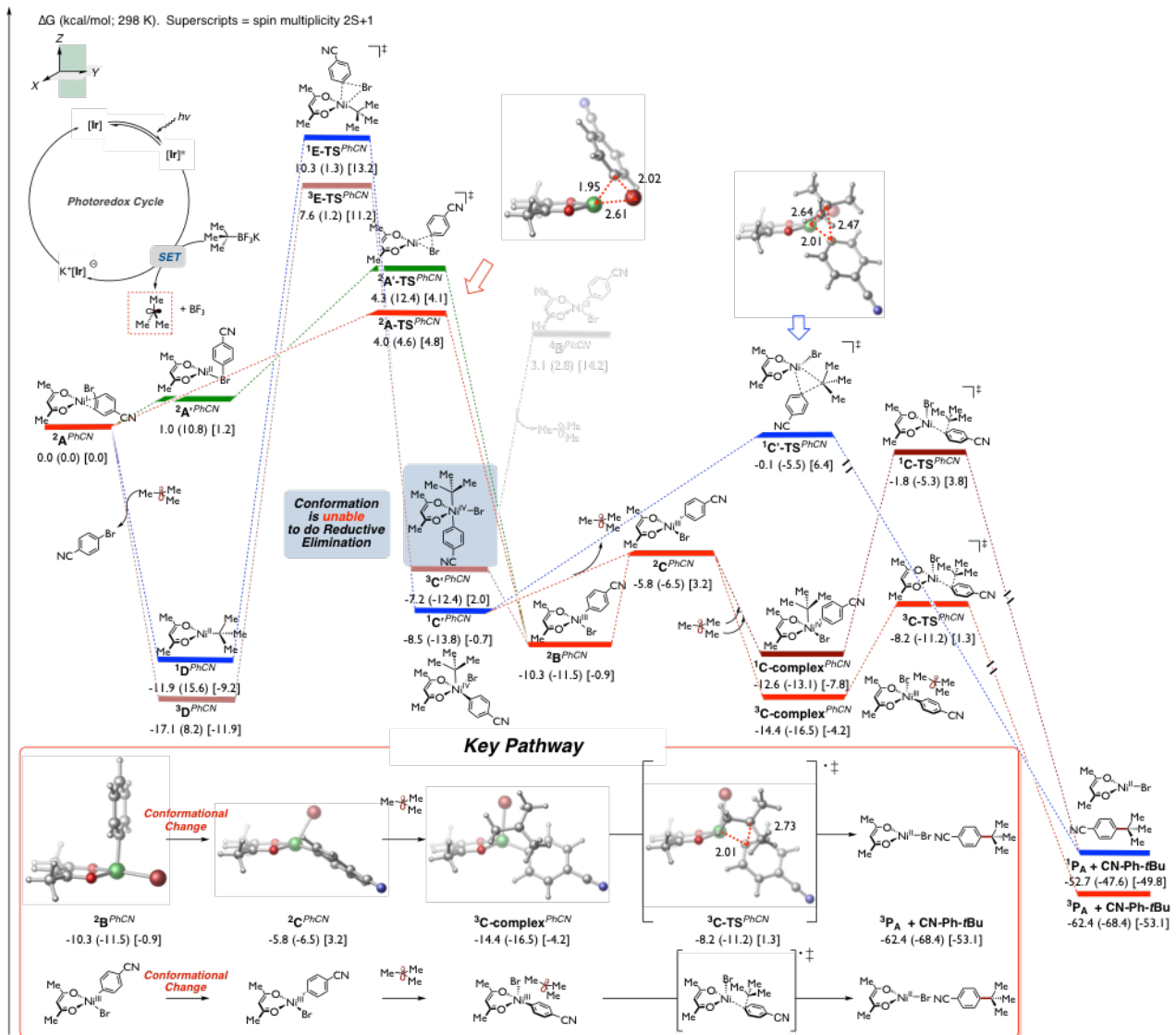
### C. Energetics with Different Methods and Conformational Search



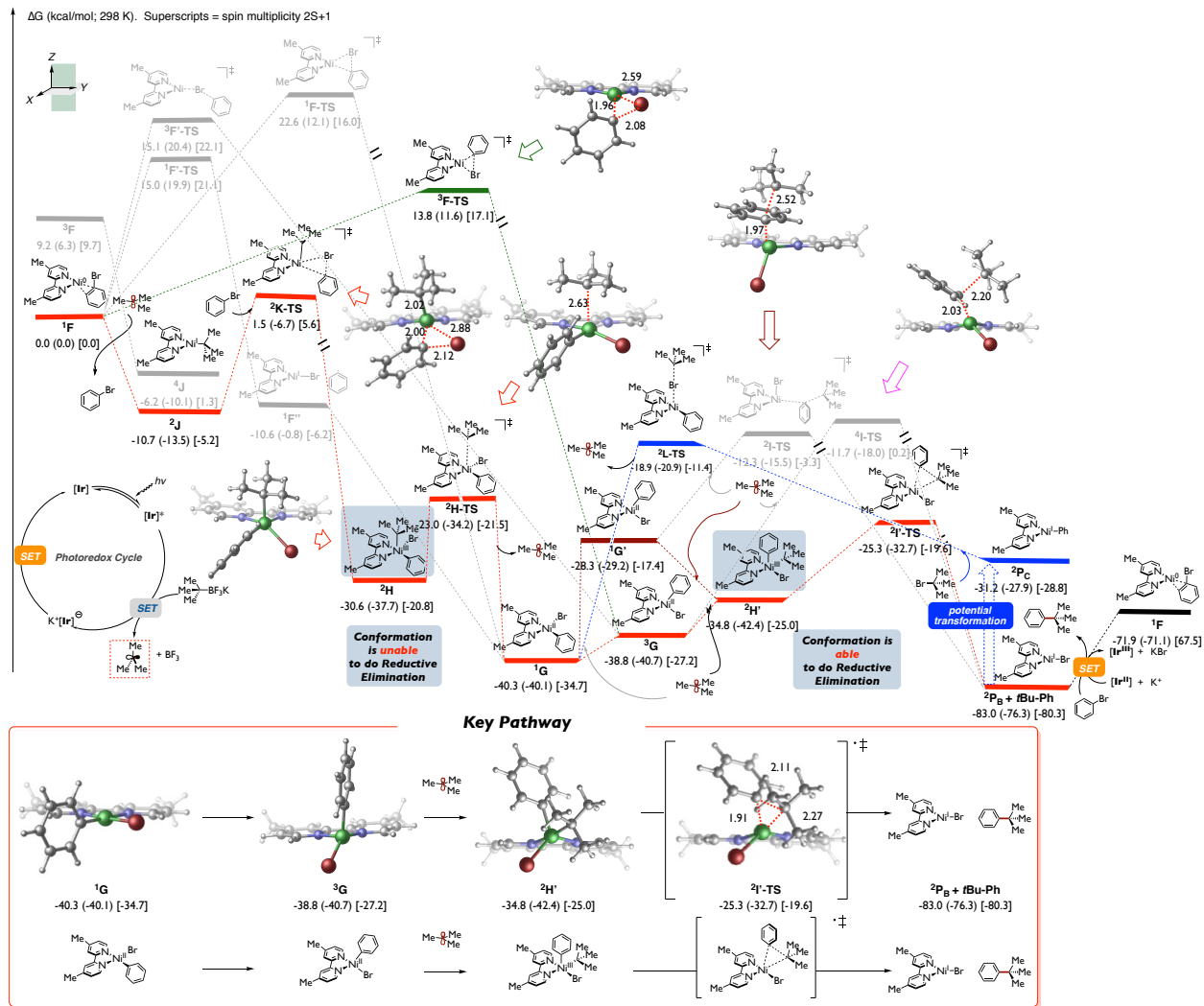
**Figure S1.** Calculated energetics of the Ni-catalyzed cross-coupling between *tert*-butyl radical and phenyl bromide using anionic TMHD as ligand. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis), UM06/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket), and DLPNO-CCSD(T)/def2-TZVPP-gas//UB3LYP-D3/def2-SVP-CPCM(THF) (in brace) levels of theory. Energetics of the last SET step was calculated in UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis), UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory for better comparison with the Ni-bipyridine system.



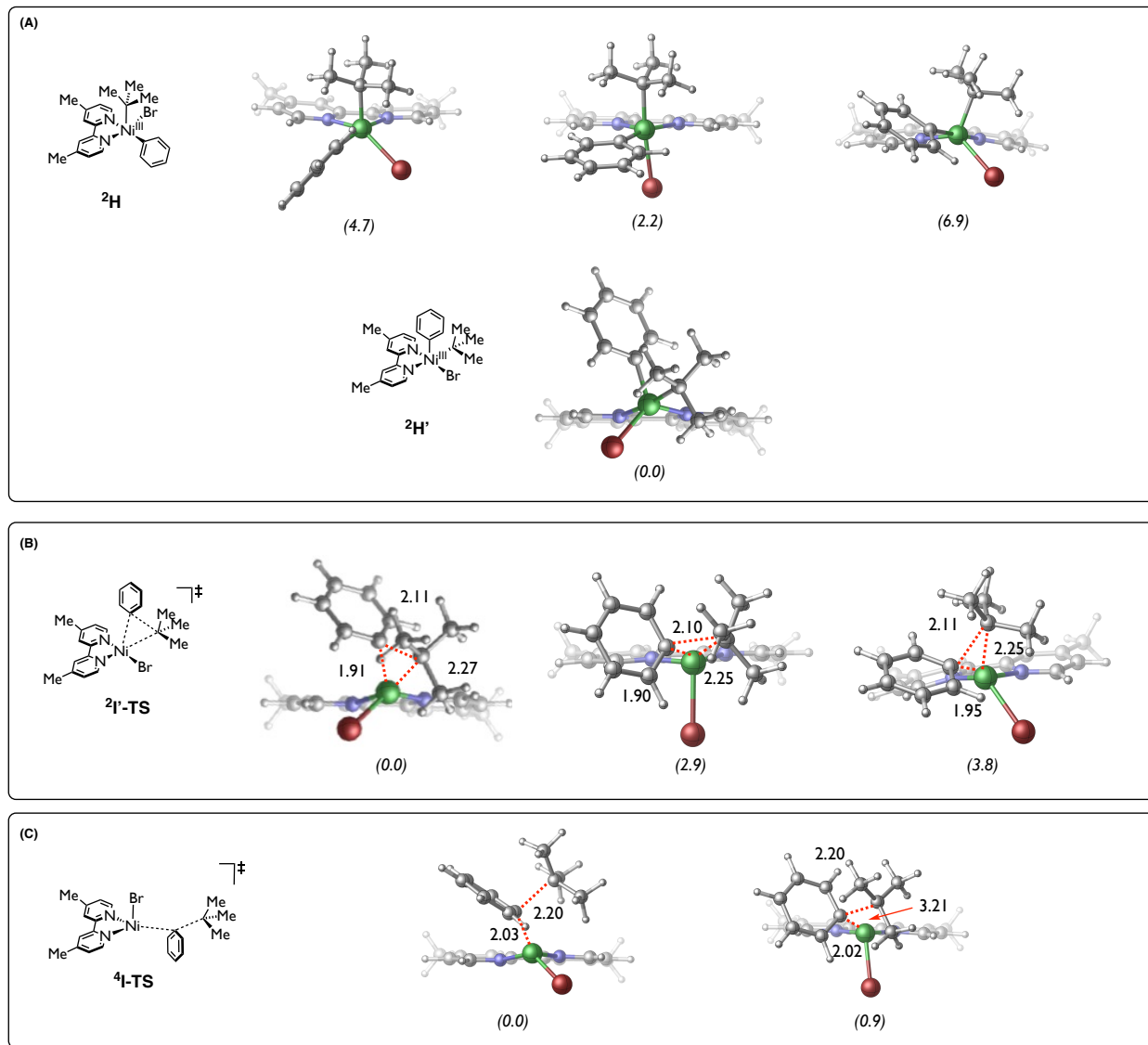
**Figure S2.** Conformational search of key intermediates and transition states of Ni-catalyzed cross-coupling between *tert*-butyl radical and phenyl bromide using anionic TMHD as ligand. Relative free energies (kcal/mol) were calculated with respect to corresponding lowest-energy level structure at the UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) level of theory.



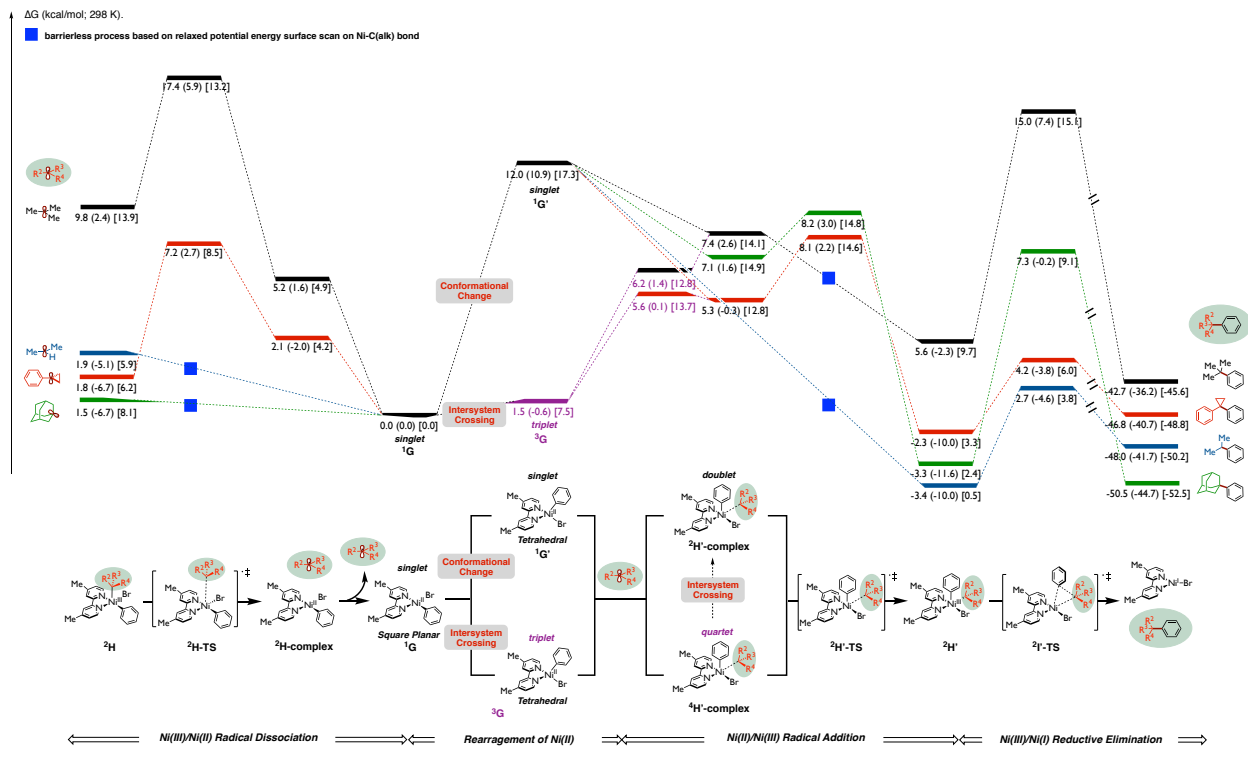
**Figure S3.** Calculated energetics of the Ni-catalyzed cross-coupling between *tert*-butyl radical and 4-bromobenzonitrile using anionic acac as ligand. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.



**Figure S4.** Calculated energetics of the Ni-catalyzed cross-coupling between *tert*-butyl radical and phenyl bromide using neutral bipyridine as ligand. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis), and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

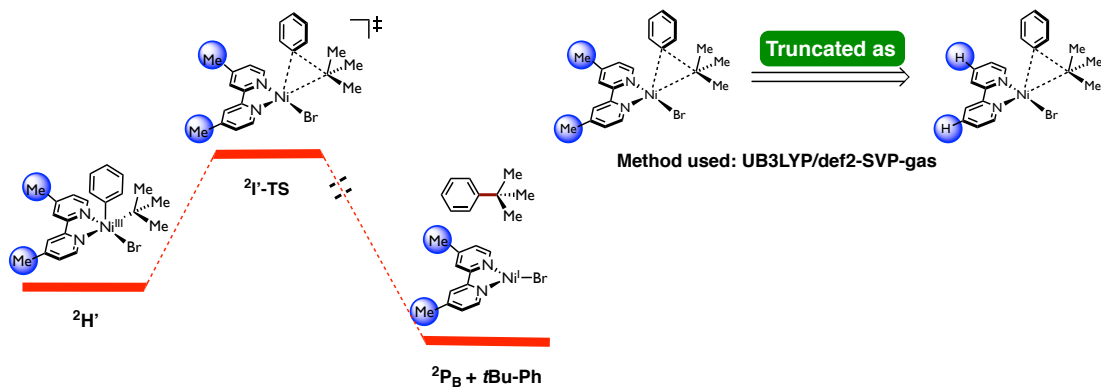


**Figure S5.** Conformational search of key intermediates and transition states of Ni-catalyzed cross-coupling between *tert*-butyl radical and phenyl bromide using neutral bipyridine as ligand. Relative free energies (kcal/mol) were calculated with respect to corresponding lowest-energy level structure at the UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) level of theory.

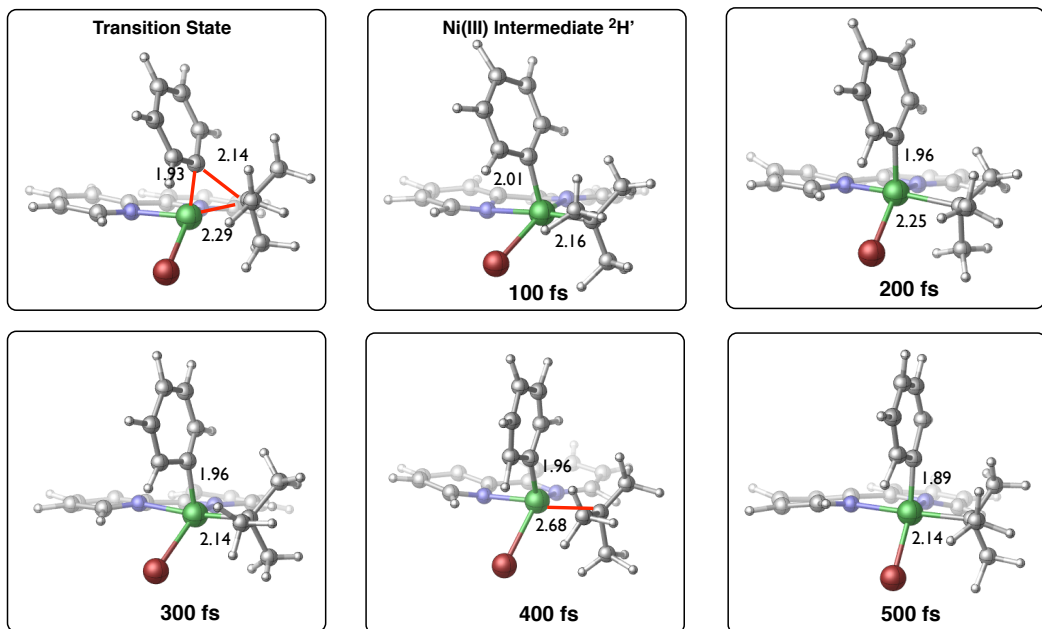


**Figure S6.** Calculated energetics of the key pathway using neutral bipyridine as ligand and *tert*-butyl radical (black), *iso*-propyl radical (blue), 1-phenylcyclopropyl radical (red) and adamantly radical (green) as radical substrates. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis), and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

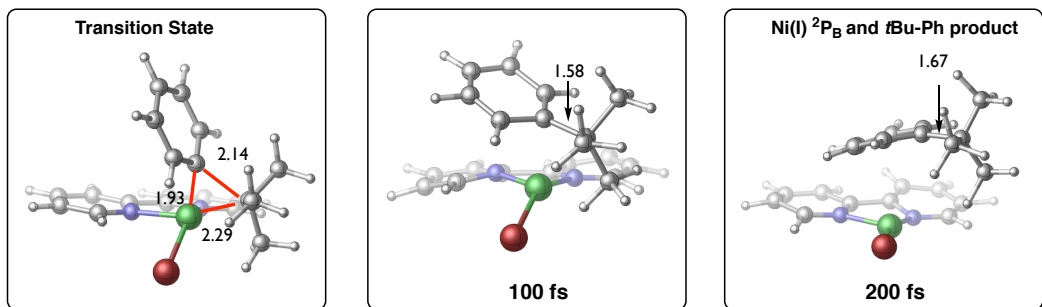
Starting from *singlet spin square planar* Ni(II) species **1G**, *tert*-butyl radical can undergo radical addition from axial position via **2H-TS** to generate the endergonic Ni(III) intermediate **2H**, which is a reversible process so *tert*-butyl radical is less favorable to be bound to Ni center. Alternatively, **1G** could undergo conformational change or intersystem crossing to generate *singlet spin tetrahedral* Ni(II) species **1G'** and *triplet spin tetrahedral* **3G**, respectively. Upon interaction with *tert*-butyl radical, Ni species can form doublet or quartet (purple energy values) complex, then after barrierless radical addition (blue squares, based on relaxed potential energy surface scan on Ni-C bond) Ni(III) species **2H'** could be formed. More importantly, the barrier of reductive elimination step of *tert*-butyl radical (15.0 kcal/mol with respect to **1G**) is much higher than those with other alkyl radicals. This might explain the observed failure with *tert*-butyl radical substrate and the success of adamantly and 1-phenylcyclopropyl radicals in experiment.<sup>13</sup>



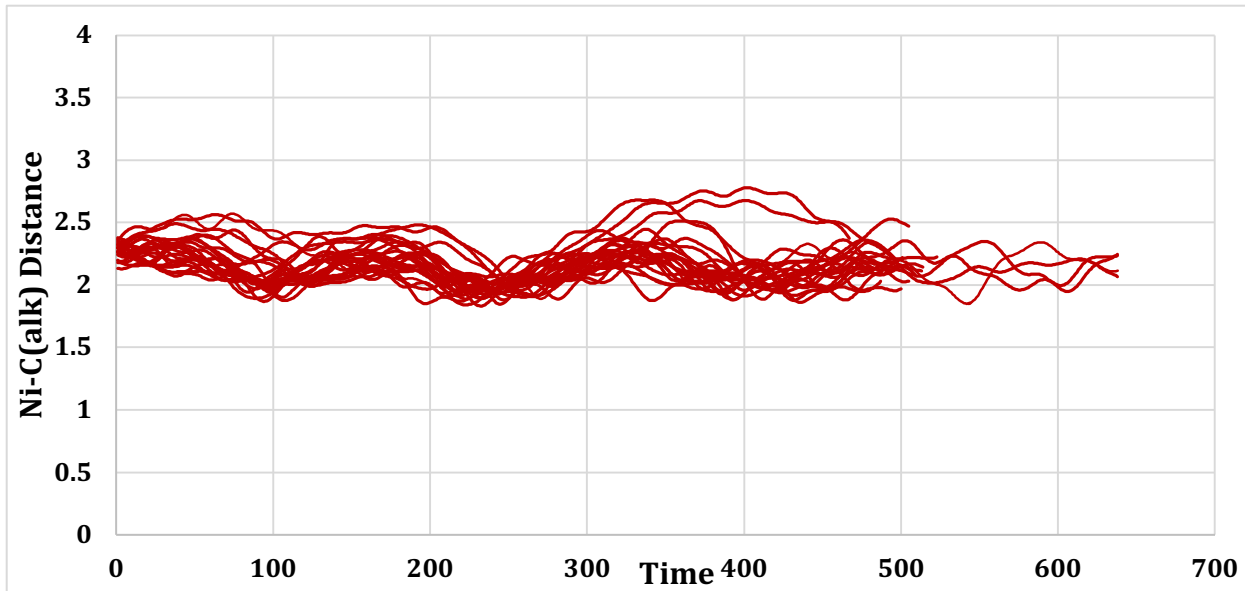
(A) example of reverse direction from  $^2\text{I}'\text{-TS}$



(B) example of forward direction from  $^2\text{I}'\text{-TS}$

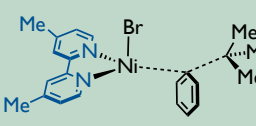
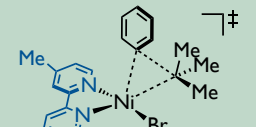
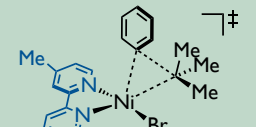
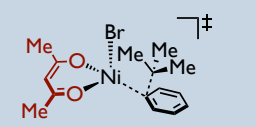
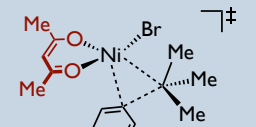
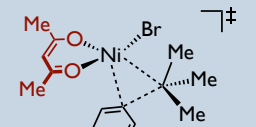


**Figure S7.** Snapshots of representative examples of quasi-classical dynamic calculation of truncated system from reductive elimination transition state  $^2\text{I}'\text{-TS}$  in both forward and backward directions of Ni-catalyzed cross-coupling between *tert*-butyl radical and phenyl bromide in Ni-bipyridine system.

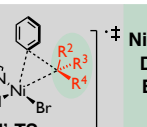
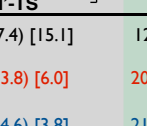
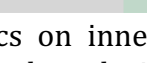


**Figure S8.** Time (fs) plot of trajectories versus Ni-C(alkyl radical) bond length ( $\text{\AA}$ ) of truncated system initiated at the reductive elimination transition state  $^2\text{I}'\text{-TS}$  in the reverse direction.

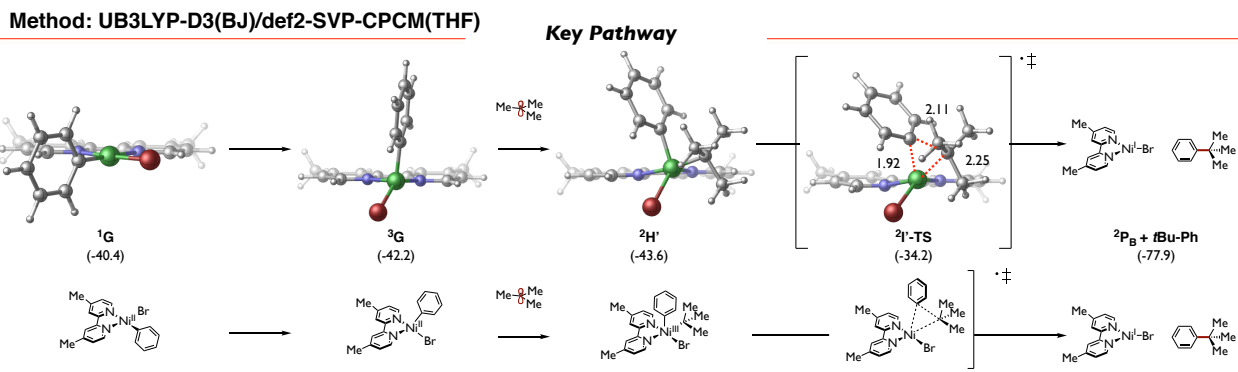
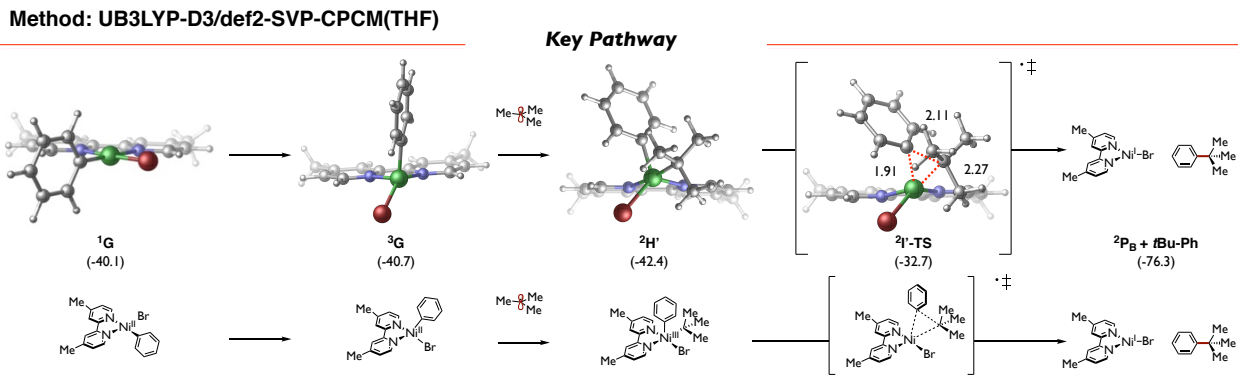
The red lines represent the results of more than 20 trajectories (ran for 500 to 700 fs) obtained, all of which show that the Ni-C(alk) bond length lies in the range of 1.8 to 2.8  $\text{\AA}$ . These results suggest that after initiated in the reverse direction at  $^2\text{I}'\text{-TS}$ , Ni(III) intermediate  $^2\text{H}'$  was formed and maintained until the end of trajectory, without the observation of *tert*-butyl radical dissociating from Ni center and generating corresponding Ni(II) intermediate (as shown in Figure S7). Overall, the results of quasi-classical dynamic calculation is consistent with the proposed pathway that the dissociation of *tert*-butyl radical from Ni(III) intermediate  $^2\text{H}'$  is uphill in energy. Also, the radical addition process of *tert*-butyl radical to Ni center occurs on *singlet spin tetrahedral* Ni species  $^1\text{G}'$  or *triplet spin tetrahedral* Ni species  $^3\text{G}$  rather than *singlet spin square planar* Ni species  $^1\text{G}$ , because the former Ni(II) species is either higher in energy level ( $^1\text{G}'$ ) than Ni(III) intermediate  $^2\text{H}'$  while the latter Ni(II) is lower, and the dynamic simulations operate on single-spin state potential energy surface thus cannot reveal the performances of system when spin state is changed. The exploration in which intermediate does the alkyl radical add to is ongoing in our group.

			$\Delta E_{\text{activation}}$ (kcal/mol)	$\Delta E_{\text{distortion}}$ (kcal/mol)	$\Delta E_{\text{interaction}}$ (kcal/mol)
<i>Outer-Sphere</i>		<b>2I-TS</b>	15.0 (11.6) [18.3]	24.6 (24.8) [29.9]	-9.6 (-13.2) [-11.6]
		<b>4I-TS</b>	13.6 (7.1) [19.8]	27.6 (25.4) [33.1]	-14.0 (-18.3) [-13.3]
<i>Inner-Sphere</i>		<b>2I'-TS</b>	-3.5 (-11.2) [-3.4]	37.1 (35.7) [42.6]	-40.6 (-46.9) [-46.0]
		<b>1C-TS</b>	-14.5 (-11.1) [-18.2]	16.0 (16.4) [18.3]	-30.5 (-27.6) [-36.5]
<i>Outer-Sphere</i>		<b>3C-TS</b>	-17.4 (-19.2) [-17.1]	17.4 (18.0) [19.8]	-34.8 (-37.2) [-36.9]
		<b>1C'-TS</b>	-12.3 (-17.7) [-16.6]	18.6 (19.2) [19.6]	-30.9 (-36.9) [-36.2]

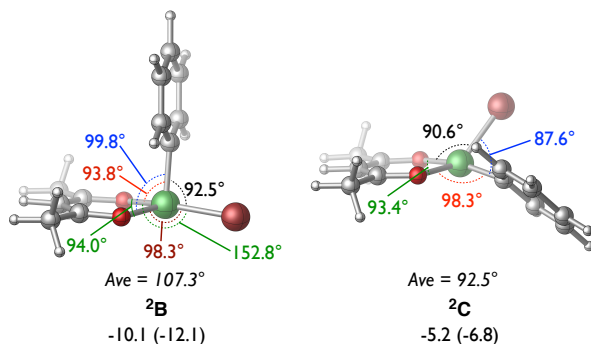
**Table S1.** Activation Strain-Distortion/Interaction analysis of C(sp<sup>2</sup>)-C(sp<sup>3</sup>) bond formation step. Relative electronic energy values were computed with respect to separate corresponding phenyl-bromo-Ni-ligand species and *tert*-butyl radical at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

	$\Delta G$ (kcal/mol; 298K)			$\Delta E_{\text{activation}}$ (kcal/mol)	$\Delta E_{\text{distortion}}$ (kcal/mol)	$\Delta E_{\text{interaction}}$ (kcal/mol)	
	<b>1G</b>	<b>2H'</b>					
	5.6 (-2.3) [9.7]	15.0 (7.4) [15.1]		12.8 (20.7) [8.7]	-3.5 (-11.2) [-3.4]	37.1 (35.7) [42.6]	-40.6 (-46.9) [-46.0]
	0.0 (0.0) [0.0]	-2.3 (-10.0) [3.3]		20.1 (27.8) [14.5]	-13.2 (-21.2) [-11.4]	33.6 (32.7) [53.4]	-46.8 (-53.9) [-64.8]
	-3.4 (-10.0) [0.5]	2.7 (-4.6) [3.8]		21.5 (28.1) [17.6]	-14.6 (-22.0) [-13.6]	29.8 (28.1) [35.4]	-44.4 (-50.1) [-49.0]
	-3.3 (-11.6) [2.4]	7.3 (-0.2) [9.1]		19.2 (27.5) [13.5]	-8.7 (-16.2) [-6.8]	28.2 (27.3) [48.0]	-36.9 (-43.5) [-54.8]

**Table S2.** Grey: calculated energetics on inner-sphere reductive elimination step with different alkyl radicals; Green: related analysis on bond dissociation energy of Ni(III) intermediate; Yellow: activation strain-distortion/interaction analysis on reductive elimination transition state (relative electronic energy values was calculated with respect to separate corresponding Ni(II) species and *tert*-butyl radical). All relative energy values were computed with respect to Ni(II) species at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

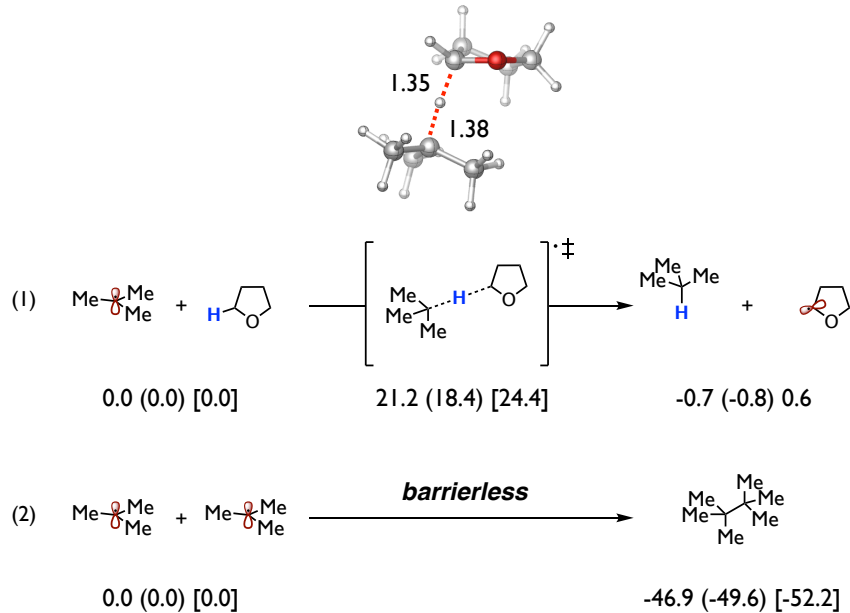


**Figure S9.** Comparison of key pathway in Ni-bipyridine system with original zero damping and Becke-Johnson damping of Grimme's dispersion correction. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) level of theory.



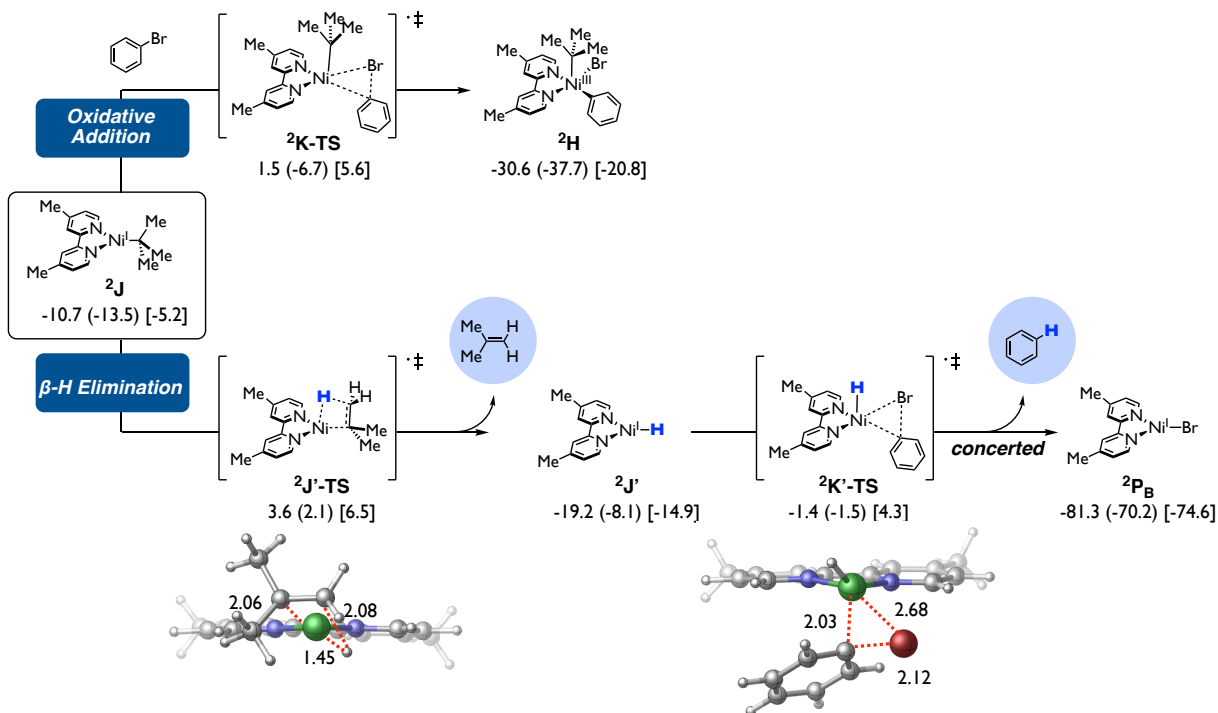
**Figure S10.** Structural parameters of  $^2\text{B}$  and  $^2\text{C}$ .

We have described "distorted tetrahedral" and "distorted square planar" as defined by the average angle deviations from a true tetrahedral geometry ( $109^\circ$ ) and square planar ( $90^\circ$ ).<sup>12</sup> As shown above, the average angle of the "distorted tetrahedral" structure  $^2\text{B}$  is  $107.3^\circ$  which is close to the angle of standard tetrahedral structure ( $109^\circ$ ). Also, the average angle of the "distorted square planar" structure  $^2\text{C}$  is  $92.5^\circ$  which is close to the angle of standard square planar structure ( $90^\circ$ ).



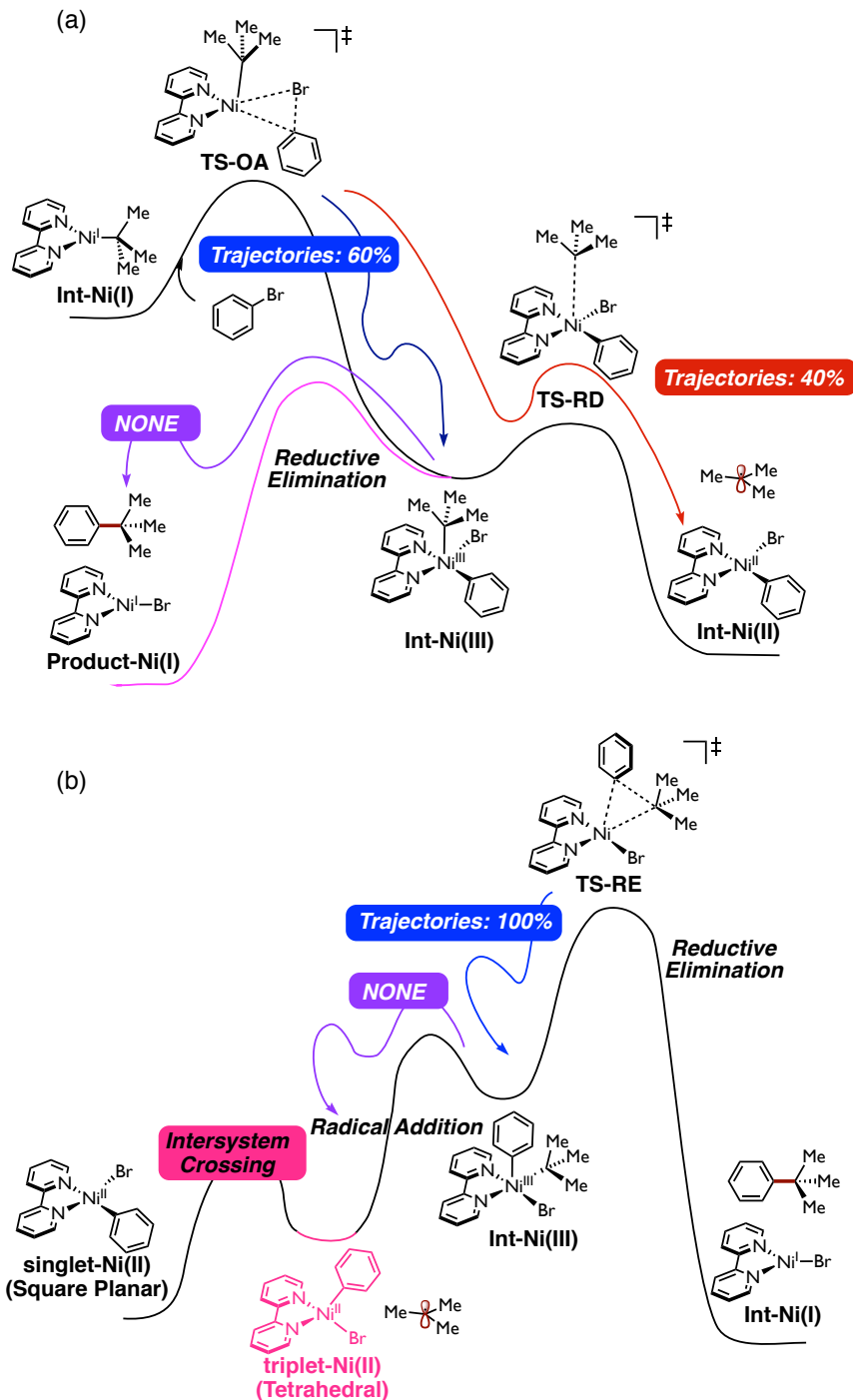
**Figure S11.** Calculated energetics of (1) HAT reaction between *tert*-butyl radical and THF, and (2) dimerization of two *tert*-butyl radicals. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

We computed the HAT pathways between the presumably formed *tBu* radical and THF solvent (eq.1) and found that the HAT of *tBu*• from THF is 0.7 kcal/mol downhill in energy with barrier of 21.1 kcal/mol. In comparison, the barrier of HAT of *tBu*• from THF is only slightly higher than that of radical addition of *tBu*• to Ni(II)(bpy) intermediate **1G** (17.3 kcal/mol at **2H-TS**), thus HAT pathway is feasible and stays competitive to the radical addition pathway. This could explain the observed experimental results<sup>13</sup> that both Ar-H (3-15% yield) and THF-Ar were observed in the stoichiometric studies using (bpy)Ni(Ar)(Br) complex and *tBu*-BF<sub>3</sub>K reagent. Moreover, we considered possible dimerization between two *tert*-butyl radicals (eq.2) and found that the dimerization process was barrierless based on relaxed potential energy surface scan on the newly formed C-C bond, resulting in the formation of 2,2-4,4-tetramethylbutane which is 46.9 kcal/mol downhill. This suggests that under higher concentration of *tBu* radical, as expected, these radicals will likely lead to dimerization instead of undergoing radical addition to Ni(II) intermediate to generate Ni(III) intermediate, which is uphill in energy. Given the efficiency of the system, these calculations support low concentration of *tBu* radical which lead to the desired cross-coupling product and only minor formation of side products such as HAT from solvent (due to higher barrier) or dimerization (due to low radical concentration).

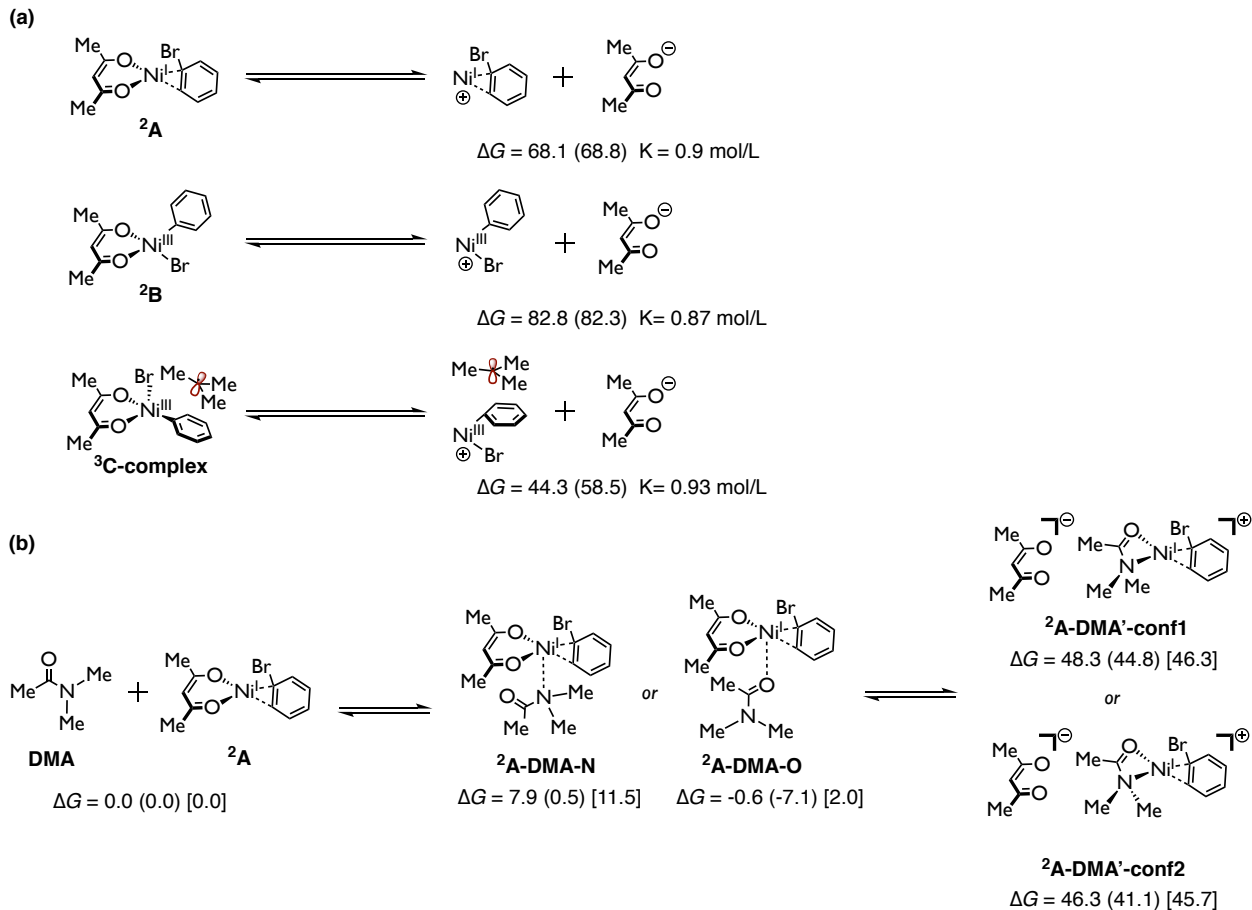


**Figure S12.** Calculated energetics of oxidative addition and  $\beta$ -H elimination pathways from Ni(I)-tBu intermediate **2J**. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

We also performed calculation on possible  $\beta$ -H elimination pathways (*vide infra*) from (bpy)Ni(tBu) intermediate **2J**, and found that oxidative addition to PhBr (via **2K'-TS**, with barrier of 12.2 kcal/mol with respect to **2J**) is comparable with the  $\beta$ -H elimination pathway (via **2J'-TS**, with barrier of 14.3 kcal/mol). Moreover, since the difference between barrier of oxidative addition pathway (12.2 kcal/mol) and that of  $\beta$ -H elimination pathway (14.3 kcal/mol) is only 2.1 kcal/mol, the  $\beta$ -H elimination pathway from (bpy)Ni(tBu) intermediate **2J** is a possible side reaction pathway in this reaction system leading to the generation of metal hydride species (bpy)Ni(H) and isobutylene as side products. Unfortunately, isobutylene has a low boiling point (-7.9 °C) and it is highly volatile thus challenging to assess experimentally. Nonetheless, the newly formed (bpy)Ni(H) species **2J'** can undergo oxidative addition with phenyl bromide (via **2K'-TS**), followed by concerted Ph-H bond formation (observed via intrinsic reaction coordinate at **2K'-TS**) to generate the experimentally observed side product benzene (Ph-H). Taken together, these computational and experimental results provide strong support for the formation of observed side products in experiment (i.e., THF-Ar adduct and Ar-H)<sup>13</sup> due to H atom transfer of tBu radical from THF solvent followed by C-C bond formation and  $\beta$ -H elimination followed by Ar-H bond formation, respectively.

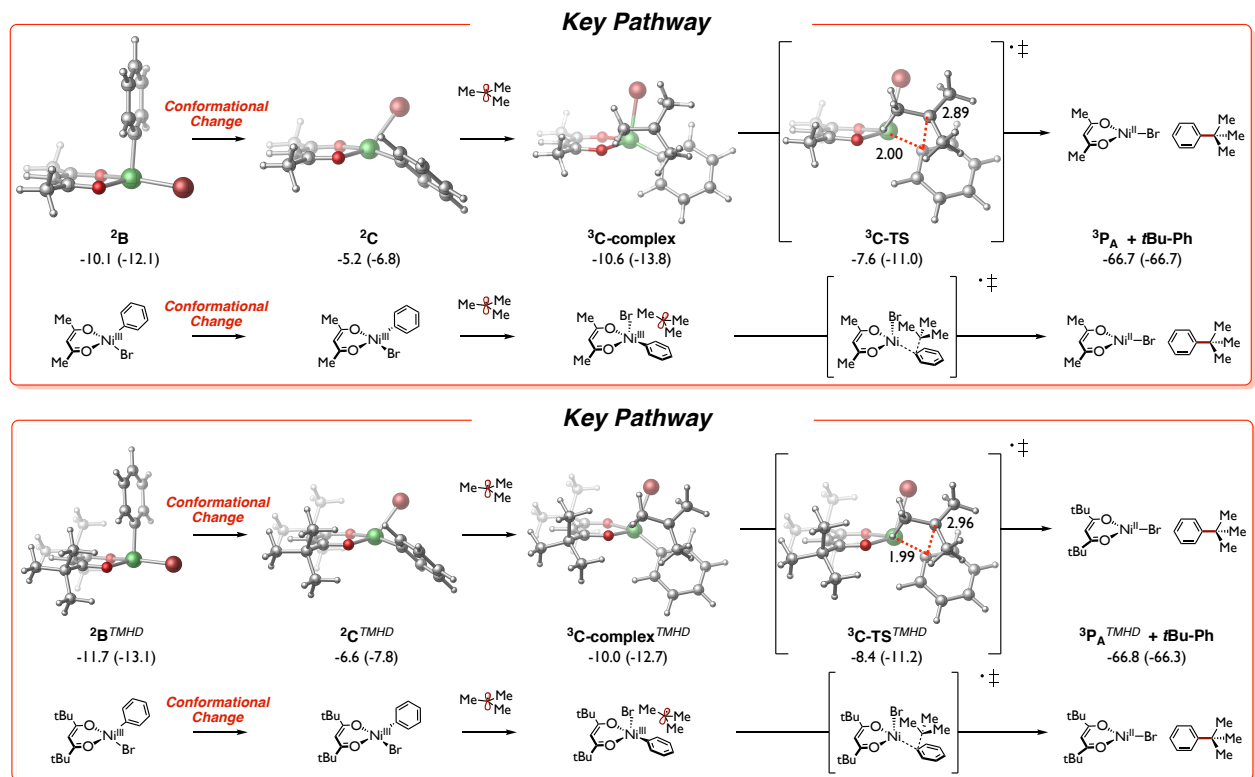


**Figure S13.** Preliminary results of quasi-classical dynamic simulations starting from (a)  ${}^2\text{K-TS}$  in forward direction and (b)  ${}^2\text{I}'\text{-TS}$  in reverse direction.



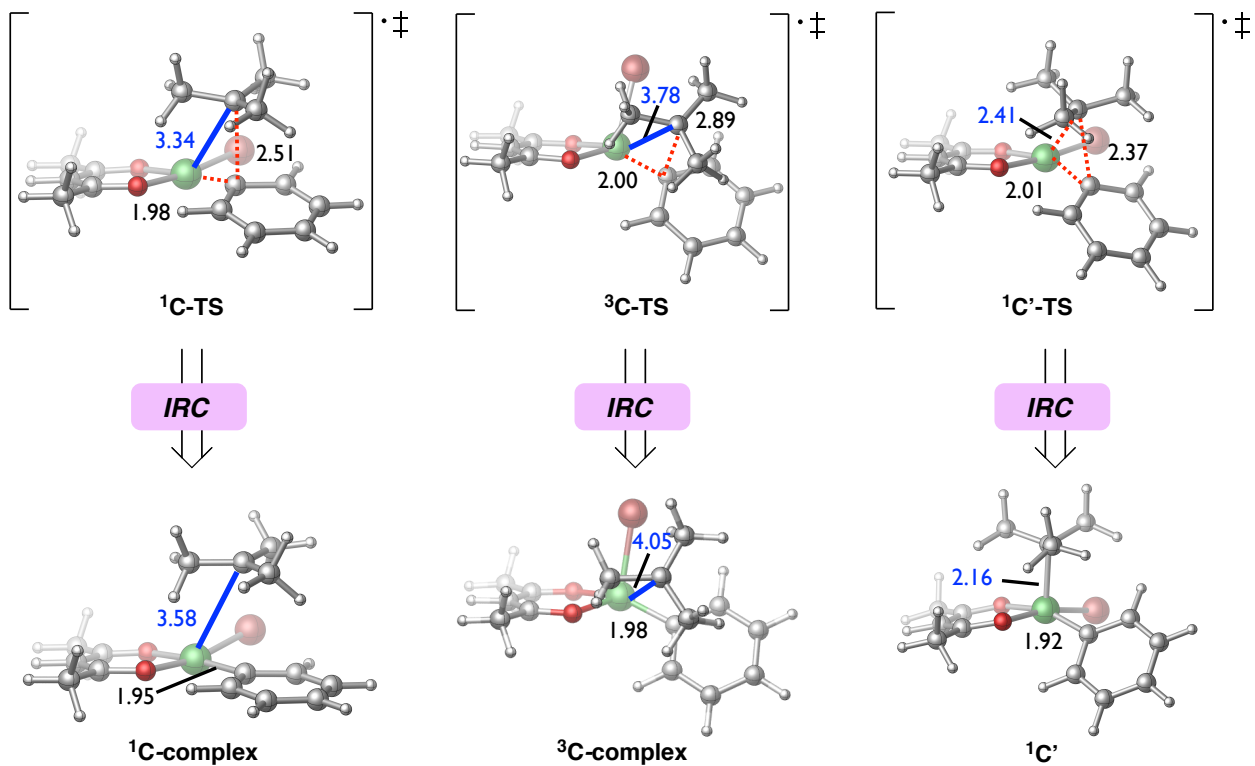
**Figure S14.** Calculated energetics of (a) ligand dissociation from key Ni(acac) species and (b) displacement and coordination of DMA solvent on key Ni(acac) species. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

To assess the likelihood of ligand dissociation, we have performed addition calculations to estimate the energy associated with ligand dissociation (shown above, Figure S14a), which showed there is an extremely energy uphill if the ligand dissociates from the Ni atom and the equilibrium constant for ligand dissociation is small. As a result, ligand dissociation pathways are unlikely. We also performed computations to gain insight at the propensity of DMA solvent to coordinate to the nickel active species and displace the acac (TMHD) ligand. As shown above (Figure S14b), the coordination of DMA to the nickel active species (prior to substrate activation) is energetically disfavored via N-mode coordination, but only weakly (dependent on the method) favorable via O-mode coordination. However, displacement of THMD ligand (here modeled as acac) is found highly unfavorable (> 40 kcal/mol). Based on these results, TMHD ligand is likely to stay coordinated to the Ni center during the whole catalytic process, and the coordination of DMA solvent to Ni center is possible via O-mode but it could not displace TMHD ligand of key Ni catalytic species.



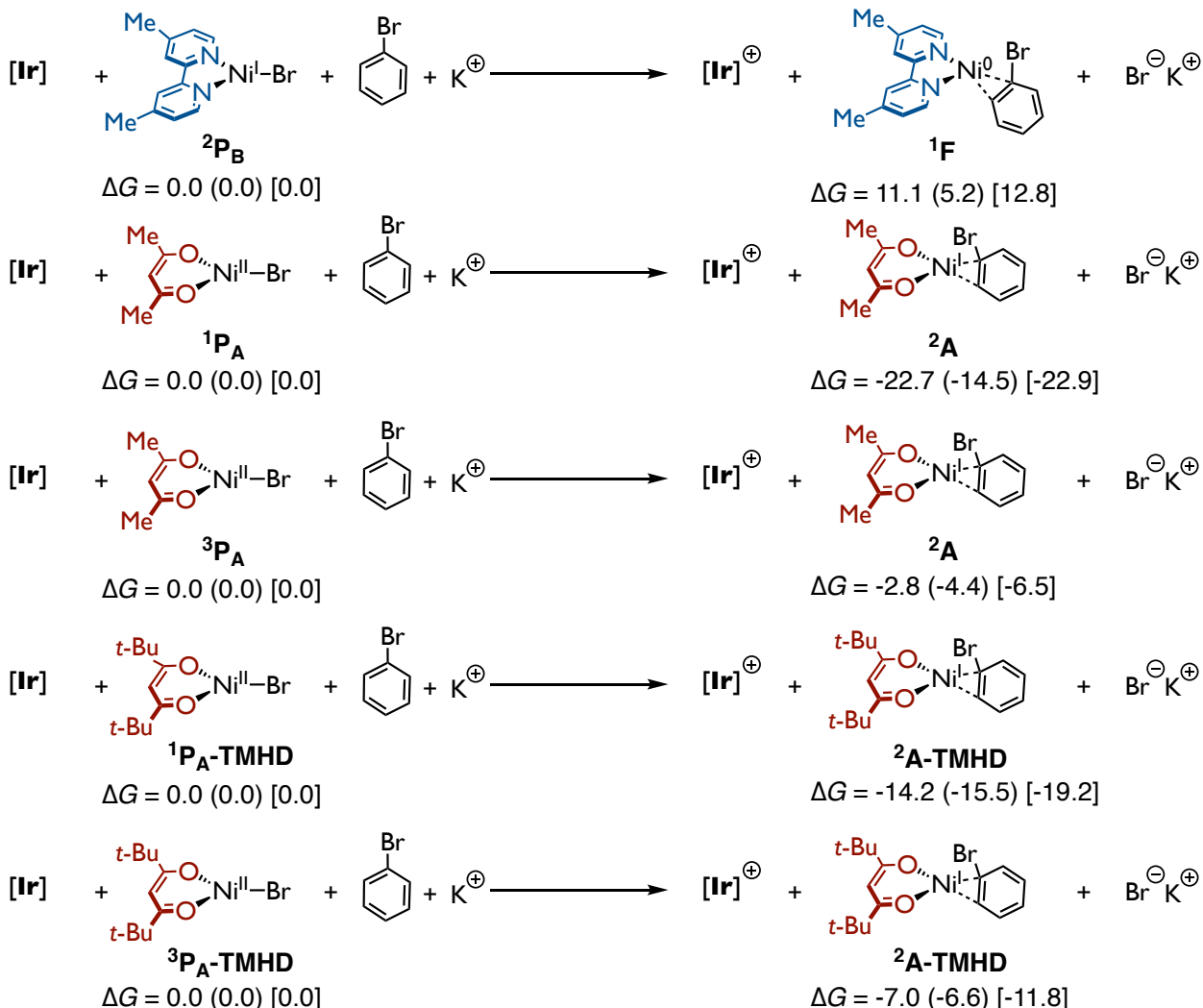
**Figure S15.** Comparison of energetics of the key pathways using truncated and full anionic TMHD as ligands. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) and UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) levels of theory.

We also computed the lowest energy pathway using the full TMHD ligand and found that the results of the full system are consistent with the pathway with truncated ligand (see below). Therefore, using acac ligand as computational model is a reasonable strategy.



**Figure S16.** Comparison of three transition states of C-C formation in Ni(acac) system and their corresponding intermediates obtained from intrinsic reaction coordinate calculations (Unit: Å).

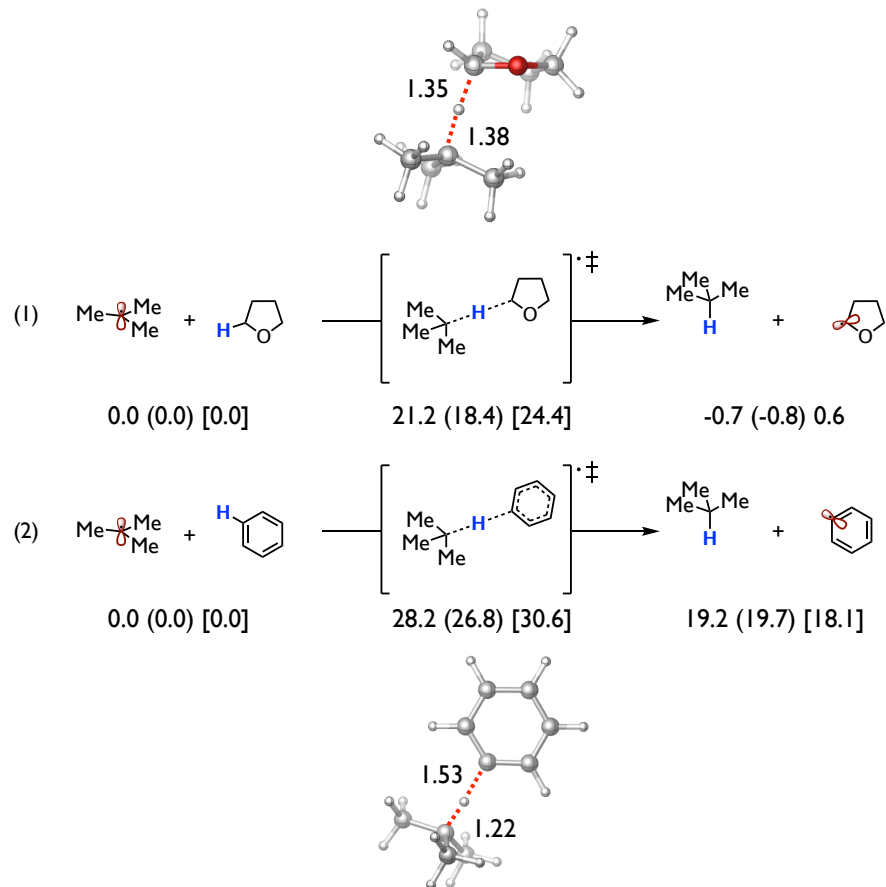
The distance between the *tert*-butyl and the Ni atom (in blue) in the *outer-sphere* transition states  $^1\text{C-TS}$  and  $^3\text{C-TS}$  are relative long (3.34 and 3.78 Å, respectively). Thus, based on these distances, there is no Ni-*tert*-butyl interaction in the *outer sphere* transition states. However, for the *inner-sphere* TS  $^1\text{C}'\text{-TS}$ , the Ni-*tBu* distance is much shorter, 2.41 Å, than that in the outer-sphere transition states. Moreover, the intrinsic reaction coordinate calculations of  $^1\text{C-TS}$ ,  $^3\text{C-TS}$  and  $^1\text{C}'\text{-TS}$  showed that the *outer-sphere* transition states  $^1\text{C-TS}$  and  $^3\text{C-TS}$  lead to Ni•••*tBu* radical complexes ( $^1\text{C-complex}$  and  $^3\text{C-complex}$ ) rather than Ni-alkyl intermediates ( $^1\text{C}'$ ) as determined by examining the corresponding distances between Ni-*tBu* and the hybridization of the optimized intermediates (from the IRCs), respectively.



**Figure S17.** Calculation of single electron transfer steps between bromo Ni-ligand species and Ir photocatalyst to regenerate corresponding starting Ni species. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

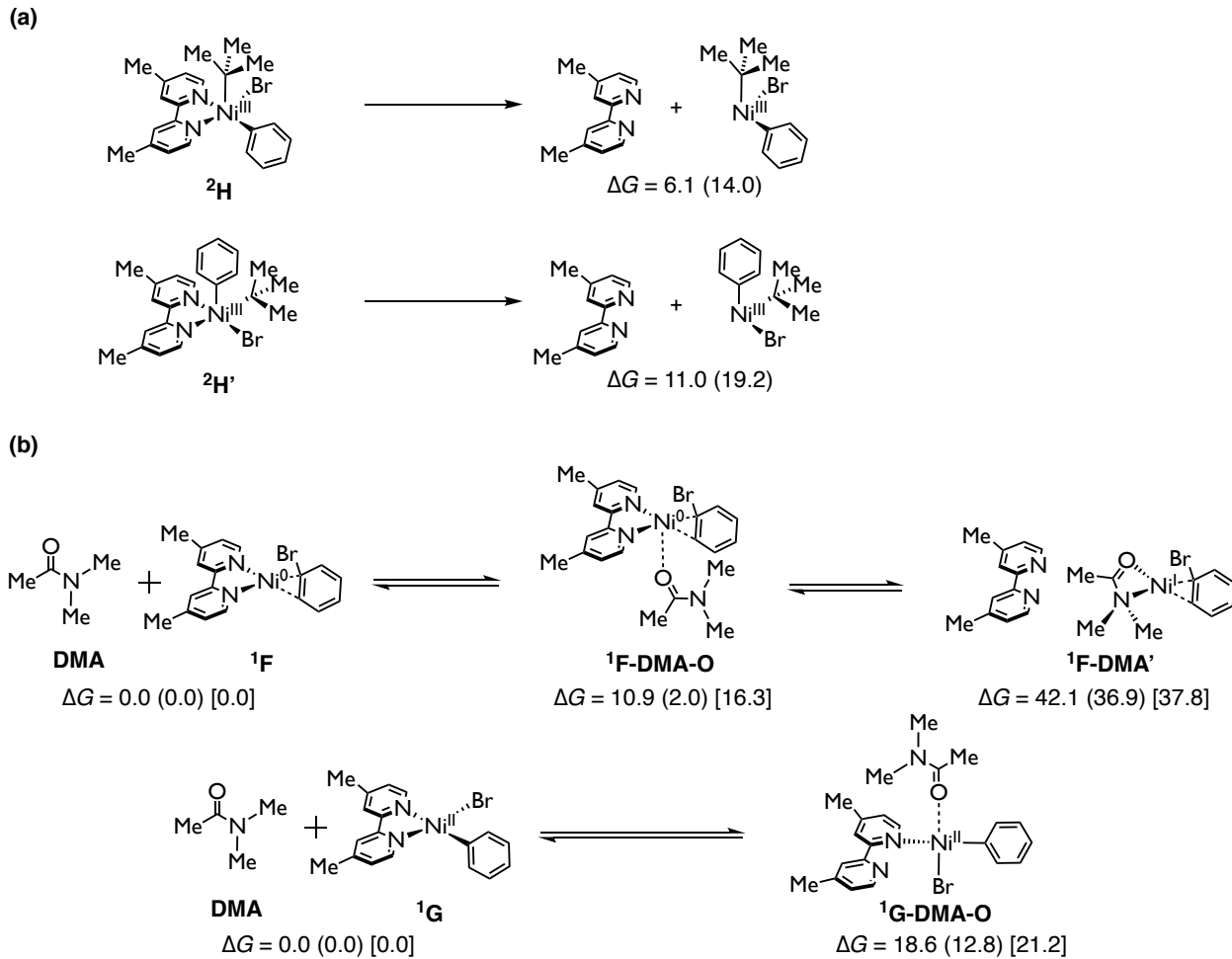
Based on the calculated thermodynamic data of SET of bromo-Ni-ligand species with reduced Ir(II) photocatalyst, the single electron transfer steps should be not problematic. Specifically, for the Ni(bpy) system, while this process is slightly uphill in energy, based on free energy span model<sup>14</sup> the reaction can still progress with turnover of catalytic cycle since the following steps from Ni(0) 1F have great thermodynamic drive with low barrier. As for Ni(acac) system, the SET steps between (acac)Ni-Br and photocatalyst are downhill in energy for both singlet and triplet spin states of (acac)Ni-Br, thus should be energetically feasible. Calculated results of full Ni-TMHD system are similar with those of the truncated Ni-acac system, both of which are downhill in energy. Therefore, in the absence of any direct experimental evidence (reduction potentials) for any of these mono-ligand bromo Ni

species to the best of our knowledge, computations predict that this step should not be problematic for both (acac)Ni(II)-Br and (bpy)Ni(I)-Br to retrieve (acac)Ni(I) complex **2A** and (bpy)Ni(0) complex **1F**, respectively.



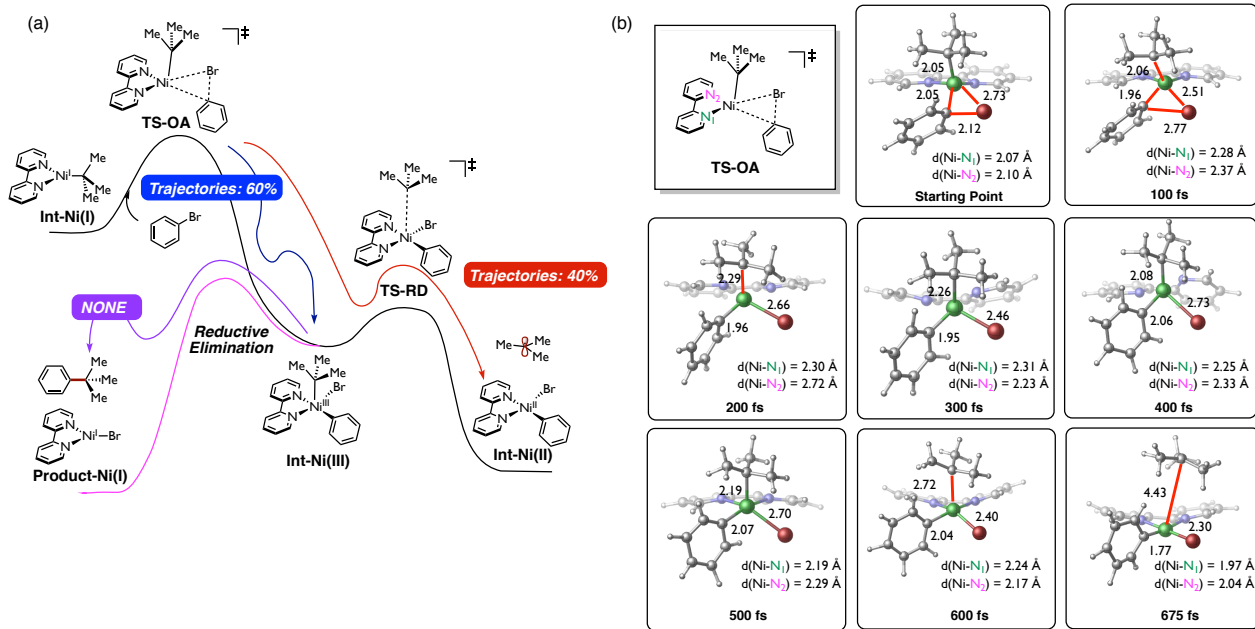
**Figure S18.** Calculated energetics of (1) HAT reaction between *tert*-butyl radical and THF, and (2) benzene. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

Calculations of HAT reactions between *tert*-butyl radical and different solvent molecules (i.e., THF and benzene) here were intended to show that the mechanism of this type of reaction might differ depending on the nature of solvent, electrophile, nucleophile even though the same *tert*-butyl radical was presumed to exist in the Ni-bipyridine system reported by Fu.<sup>15</sup>



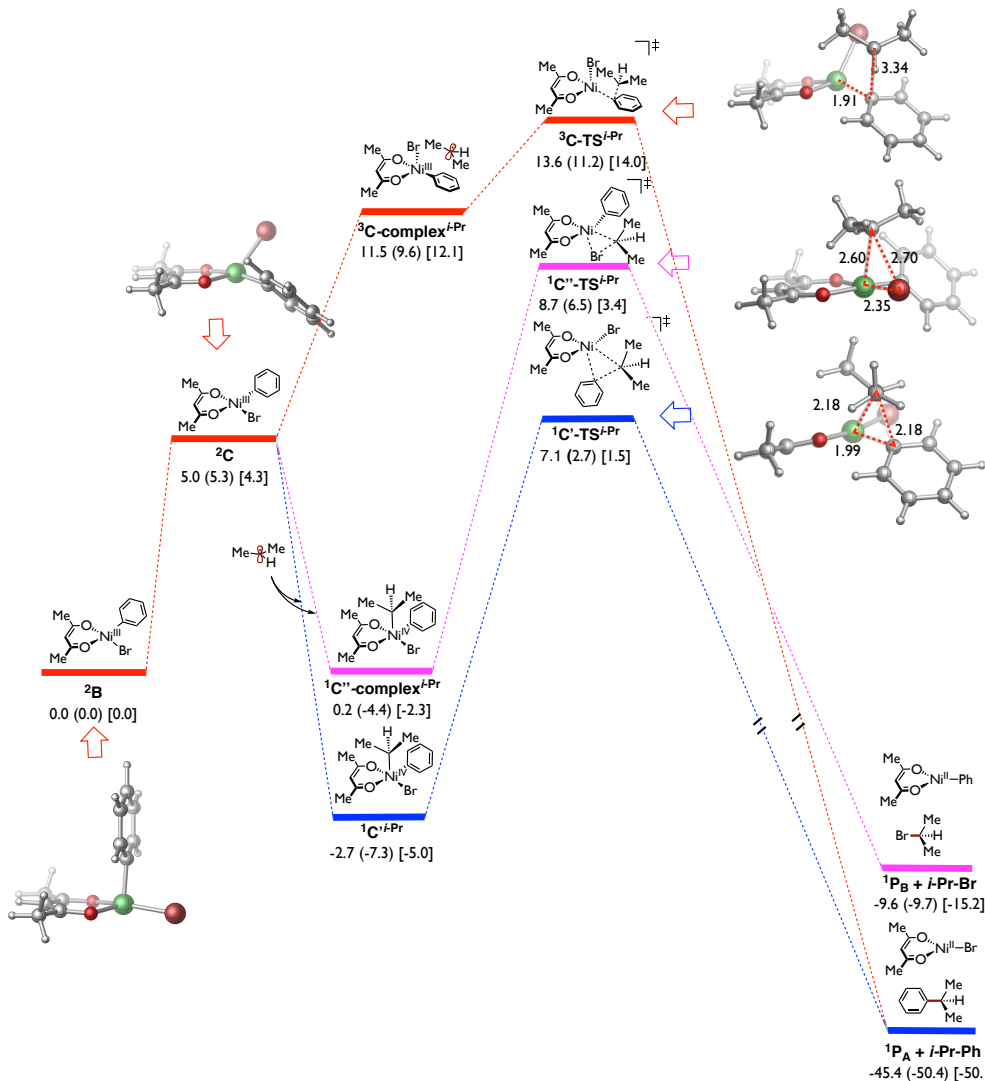
**Figure S19.** Energetics of (a) complete ligand dissociation from key Ni(III) intermediates  $^2\text{H}$  and  $^2\text{H}'$ , and (b) coordination and displacement with DMA solvent. Relative Gibbs free energy values were computed at the UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

These results show that complete ligand dissociation from Ni(III) intermediates is unlikely to occur since it's uphill in energy level from  $^2\text{H}$  and  $^2\text{H}'$  by 6.1 and 11.0 kcal/mol, respectively. Moreover, we explored the tendency of displacement or coordination with DMA solvent for the bypass-nickel system, and found that DMA coordination to Ni(0) or Ni(II) is found energetically unfavorable ( $^1\text{F-DMA-O}$  and  $^1\text{G-DMA-O}$ ) and, like the TMHD ligand, highly unfavorable to cause the dissociation of bipyridine ligand ( $^1\text{F-DMA}'$ ).



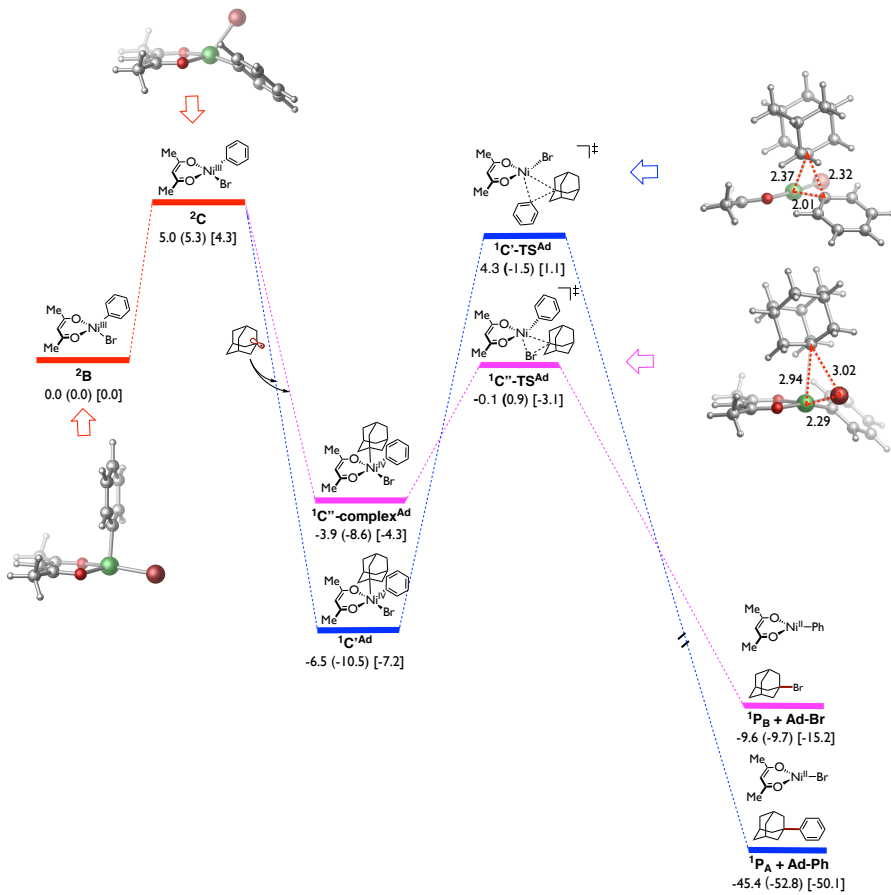
**Figure S20.** (a) Preliminary results of quasi-classical dynamic simulations starting from <sup>2</sup>K-TS in forward direction and (b) snapshots of one example trajectory undergoing radical dissociation.

To explore the likelihood of transient ligand dissociation during the reaction, we also performed quasi-classical dynamic calculations which allow us to “see” if ligand dissociation occurs, and if so, at what time step (Figure S20). Dynamic calculations starting from TS-Oxidative Addition from Ni(I) showed that complete ligand dissociation is unlikely to happen within the lifetime of Ni(III) intermediate. In addition, although transient decoordination of one of the pyridine ligands was observed during some simulated trajectories (see Figure S20(b) for an example trajectory), *the Ni(III) intermediate doesn't undergo any structural rearrangement*. Rather, Ni(III) intermediate either stayed the same conformation until the end of simulation (60% of trajectories) or underwent radical dissociation of tBu radical from axial position of Ni center (40% of trajectories).



**Figure S21.** Calculated energetics of the Ni-catalyzed cross-coupling between *iso*-propyl radical and phenyl bromide using anionic acac as ligand. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

We performed computations to investigate the effect of secondary radical in the reaction pathway and found that the favored pathway is via inner-sphere C-C bond formation with overall barriers for C-C bond formation of ~9 kcal/mol. Thus, the cross-coupling of secondary alkyl radical is also applicable in Ni-TMHD system, which is shown in the experiment results (see part D, Page S31). Presumably, the less sterically hindered secondary alkyl radical (in contrast to the tertiary alkyl radical) does not pay a penalty to form the LNi-alkyl-Ar-Br species and can quickly undergo inner sphere C-C bond formation. Thus, from these results, we hypothesize that for acac/TMHD-Ni systems, sterics more than electronics influence the nature of the C-C bond formation and whether inner- or outer-sphere C(sp<sup>2</sup>)-C(sp<sup>3</sup>) is operative.



**Figure S22.** Calculated energetics of the Ni-catalyzed cross-coupling between adamantyl radical and phenyl bromide using anionic acac as ligand. Free energies (kcal/mol) were computed at the UB3LYP-D3/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF), UB3LYP-D3/def2-SVP-CPCM(THF) (in parenthesis) and UM06/def2-TZVPP-CPCM(DMA)//UB3LYP-D3/def2-SVP-CPCM(THF) (in bracket) levels of theory.

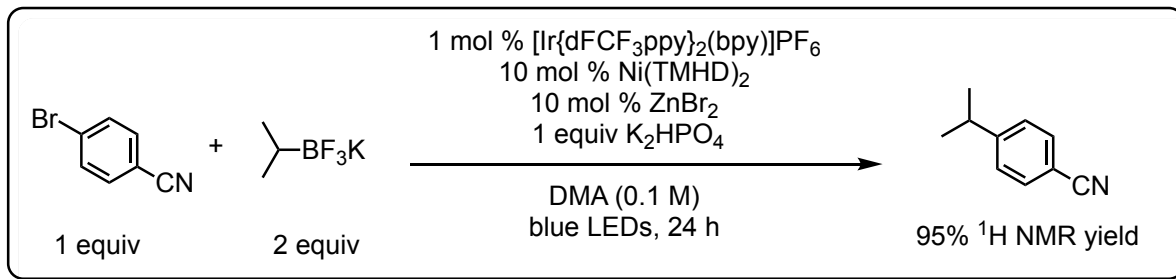
We performed computations to investigate the effect of cyclic tertiary radical in the reaction pathway, showing that, if the cyclic radical were to engage with the nickel in the cross-coupling cycle, formation of 1-bromoadamantane via inner-sphere C-Br bond formation is favored over C-C bond formation. However, since this step is reversible with only 5.7 kcal/mol in thermodynamic drive with low barrier of 3.8 kcal/mol, the acac-Ni species is likely to also undergo other pathways in this system. Presumably, the sterically hindered cyclic adamantyl radical (compared with the acyclic tertiary radical and acyclic secondary radical) needs to pay a penalty to undergo inner-sphere C-C bond formation to form the desired product 1-phenyladamantane. As a result, we assume that sterics plays an important role in the C-C bond formation step and have a significant effect on the final product. Compared with the experimental results (see part D, Page S33) where no cross-coupling product was observed and aryl bromide was recovered, our calculations suggest that *if the aryl halide substrate were activated*, the reactions of adamantyl radical should be feasible to form many possible products, including 1-bromoadamantane and phenyladamantane. While regarding the failure of activating aryl bromide substrate in this system, more exploration of its properties in the Ni-TMHD system is ongoing in our group.

## D. Experimental Details

**General:** All chemical transformations were carried out under an inert atmosphere of argon unless otherwise noted. Standard Schlenk line techniques with a 4- or 5-port dual-bank manifold were used for the manipulation of solvents and reagents. LED irradiation was accomplished as described in our previous reports.<sup>16</sup> Reactions were monitored by GC/MS and <sup>1</sup>H NMR.

**Chemicals:** DMA was purchased as 99.9%, extra dry. K<sub>2</sub>HPO<sub>4</sub> and Na<sub>2</sub>CO<sub>3</sub> were stored and dispensed in a dry glovebox. Deuterated NMR solvents were purchased and stored over 4 Å molecular sieves. IrCl<sub>3</sub>·xH<sub>2</sub>O, and Ni(TMHD)<sub>2</sub> were purchased from commercial sources. The iridium photocatalyst Ir[dFCF<sub>3</sub>ppy]<sub>2</sub>(bpy)PF<sub>6</sub> was prepared in-house by the procedure outlined in our previous publication.<sup>17</sup> Organotrifluoroborates were used as purchased without further purification.

### General Procedure I: Cross-Coupling of Secondary Trifluoroborates<sup>18</sup>

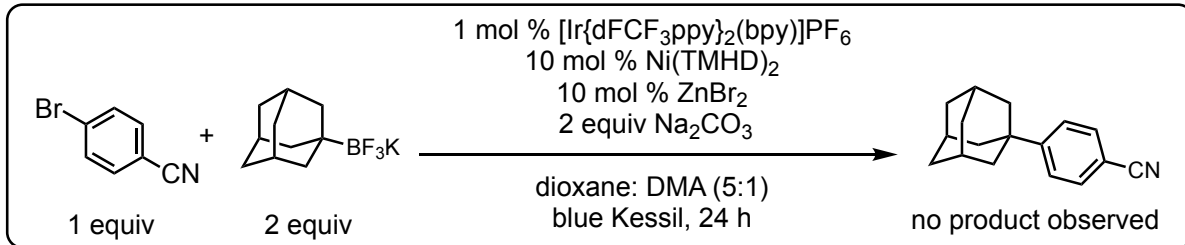


To an 8 mL reaction vial equipped with a Teflon-coated magnetic stir bar were added Ir[dFCF<sub>3</sub>ppy]<sub>2</sub>(bpy)PF<sub>6</sub> (3.0 mg, 0.003 mmol, 1 mol %), isopropyltrifluoroborate (90 mg, 0.6 mmol, 2.0 equiv), and 4-bromobenzonitrile (55 mg, 0.3 mmol, 1.0 equiv). The vial was then transferred into a glove box where Ni(TMHD)<sub>2</sub> (12.8 mg, 0.03 mmol, 10 mol %), ZnBr<sub>2</sub> (6.8 mg, 0.03 mmol, 10 mol %), and anhydrous K<sub>2</sub>HPO<sub>4</sub> (52 mg, 0.3 mmol, 1.0 equiv) were added. The vial was then capped and removed from the glovebox. Anhydrous DMA (3 mL) was added to the vial via syringe under inert atmosphere. The vial was subsequently sparged with argon for 10 min. The vial was sealed with parafilm and stirred for 24 h approximately 1 cm away from a ring of blue LED lights strips (see below). A fan was used to maintain the reaction at temperature. After completion, the crude reaction mixture was diluted with ether, and <sup>1</sup>H NMR yield was determined using 1,3,5-trimethoxybenzene as internal standard (95% yield).

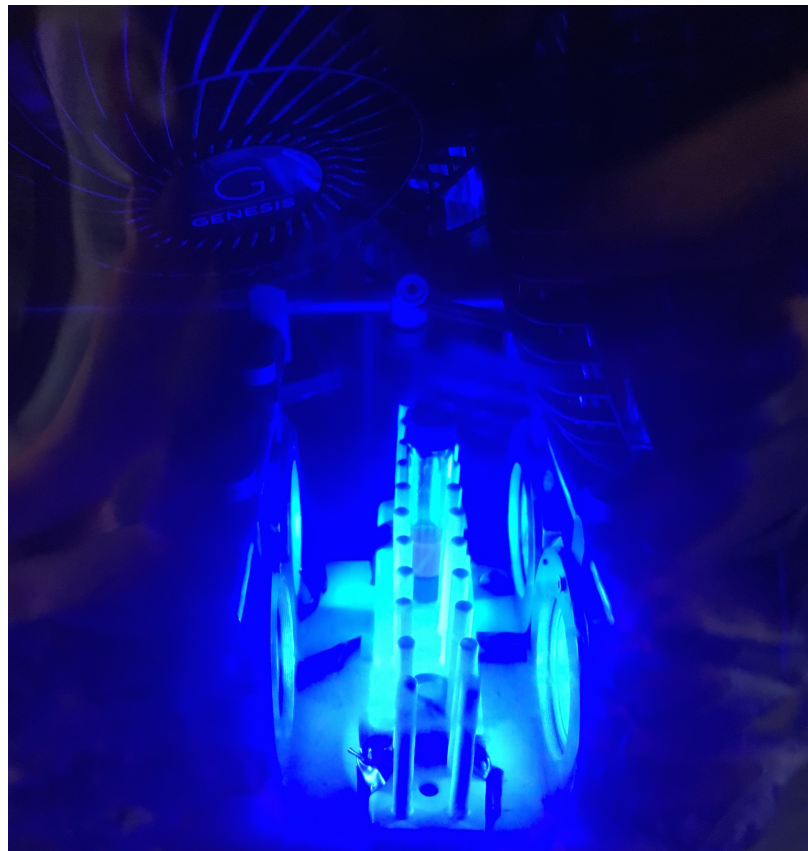


**Figure S23.** Reaction set-up using blue LEDs.

**General Procedure II: Cross-Coupling of Adamantyltrifluoroborate<sup>16a</sup>**



To an 8 mL reaction vial equipped with a Teflon-coated magnetic stir bar were added  $[\text{Ir}(\text{dFCF}_3\text{ppy})_2(\text{bpy})]\text{PF}_6$  (3.0 mg, 0.003 mmol, 1 mol %), adamantyltrifluoroborate (145 mg, 0.6 mmol, 2.0 equiv), and 4-bromobenzonitrile (55 mg, 0.3 mmol, 1.0 equiv). The vial was then transferred into a glove box where  $\text{Ni}(\text{TMHD})_2$  (12.8 mg, 0.03 mmol, 10 mol %),  $\text{ZnBr}_2$  (6.8 mg, 0.03 mmol, 10 mol %), and anhydrous  $\text{Na}_2\text{CO}_3$  (64 mg, 0.6 mmol, 2.0 equiv) were added. A mixture of anhydrous DMA and dioxane (5:1, 3 mL) was then added to the vial via syringe under inert atmosphere. The vial was then capped, removed from the glovebox, and sealed with parafilm. The reaction was stirred for 24 h using blue Kessil lamps (see below). A fan was used to maintain the reaction around 40 °C. After completion, the crude reaction mixture was diluted with ether, and  $^1\text{H}$  NMR yield was determined using 1,3,5-trimethoxybenzene as internal standard (no product observed, recovery of aryl bromide was detected).

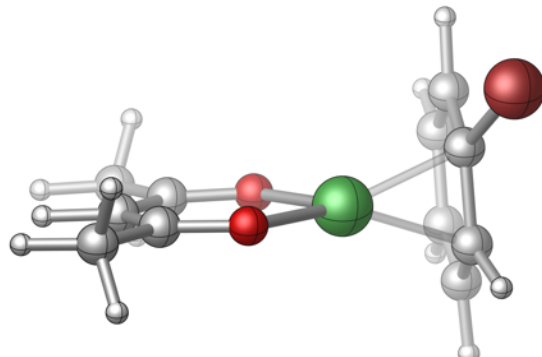


**Figure S24.** Reaction set-up using blue Kessil lamps.

## E. Coordinates and Energies of Optimized Structures in Computation

Figure S1

<sup>2</sup>A



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202379 (Hartree/Particle)

Thermal correction to Energy= 0.219143

Thermal correction to Enthalpy= 0.220087

Thermal correction to Gibbs Free Energy= 0.154245

Sum of electronic and zero-point Energies= -4658.354829

Sum of electronic and thermal Energies= -4658.338065

Sum of electronic and thermal Enthalpies= -4658.337121

Sum of electronic and thermal Free Energies= -4658.402963

C -2.53178800 -0.43393100 1.44289900

C -2.49128200 -0.70770400 0.04310200

C -2.82017300 0.30666700 -0.89855000

C -3.12620200 1.57753200 -0.44707400

C -3.16201600 1.86786400 0.93761200

C -2.88072000 0.87951200 1.86361900

Br -2.54192800 -2.55709300 -0.56147600

H -2.81567400 0.07267600 -1.96415900

H -2.52967100 -1.25152100 2.17009600

H -2.94103100 1.08783300 2.93438200

H -3.42275400 2.87405900 1.27307400

H -3.35065300 2.36310200 -1.17251700

Ni -0.56734700 -0.47208200 0.70356300

O 1.21828000 -1.16628900 0.90826800

O 0.02037900 1.33193300 0.22988800

C 1.23370500 1.69364400 0.14842900

C 2.29158200 -0.49781500 0.74328500

C 3.57343000 -1.27103100 0.96263700

H 3.59819100 -2.13082300 0.27315600

H 4.47667500 -0.66388700 0.81587100

H 3.57522100 -1.68356600 1.98485100

C 1.45377000 3.14181900 -0.23613100

H 0.94702500 3.78917700 0.49802400

H 2.51434800 3.42083000 -0.29455600

H 0.97410500 3.33085400 -1.21040400  
C 2.35582500 0.86357500 0.38303500  
H 3.34439500 1.31082700 0.27594500

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01015289$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.7324910

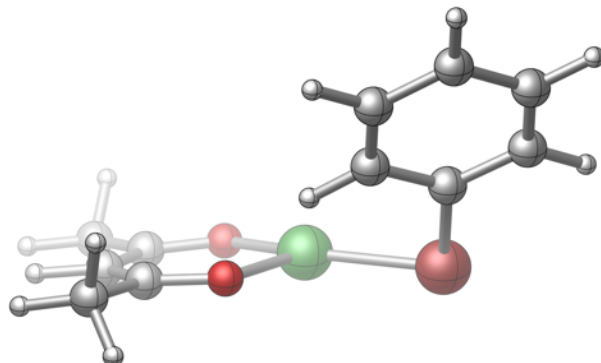
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.08046134

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4656.139208237236

<sup>2</sup>A'



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202973 (Hartree/Particle)

Thermal correction to Energy= 0.219947

Thermal correction to Enthalpy= 0.220891

Thermal correction to Gibbs Free Energy= 0.153592

Sum of electronic and zero-point Energies= -4658.339107

Sum of electronic and thermal Energies= -4658.322134

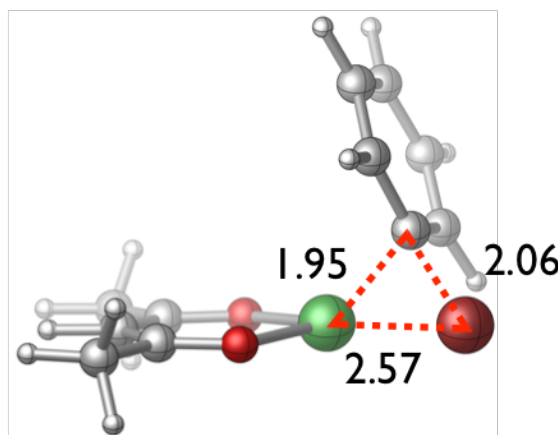
Sum of electronic and thermal Enthalpies= -4658.321189

Sum of electronic and thermal Free Energies= -4658.388488

C	-2.39558900	-1.29660400	1.46528600
O	-3.01344300	-0.46132500	2.20271700
C	-2.62407300	-1.50135300	0.08499500
H	-2.00279700	-2.24497400	-0.41559300
C	-3.57722900	-0.82185700	-0.70626700
O	-4.39832100	0.05707200	-0.28385800
C	-3.65895500	-1.12503700	-2.18735400
H	-3.36882900	-0.22458200	-2.75493600
H	-3.01600500	-1.96028300	-2.49641100
H	-4.70403500	-1.34785900	-2.45572700
C	-1.33179000	-2.11378000	2.16762300
H	-1.79735200	-2.68589100	2.98713700
H	-0.80630300	-2.80818600	1.49836800
H	-0.59918600	-1.43325400	2.63169000
Ni	-4.40417900	0.65782000	1.56899600
C	-5.84502100	3.55560900	0.40545300
C	-5.11523300	3.21248800	-0.73216100
C	-6.31983600	4.84551300	0.63839800
C	-4.85440300	4.21613700	-1.67210700
H	-4.76971000	2.18660000	-0.87699500
C	-6.04643900	5.83341500	-0.31547300
H	-6.88723200	5.08057900	1.54072600
C	-5.31538900	5.52154900	-1.46676400
H	-4.28514800	3.96951900	-2.57183700
H	-6.40818700	6.85128500	-0.15063500
H	-5.10506800	6.29799200	-2.20597100

Br -6.22832200 2.16943000 1.71547300  
 $\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01025185$   
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.7317488  
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.07809925  
DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4656.147502274090

<sup>2</sup>A-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.201937 (Hartree/Particle)

Thermal correction to Energy= 0.218204

Thermal correction to Enthalpy= 0.219148

Thermal correction to Gibbs Free Energy= 0.154635

Sum of electronic and zero-point Energies= -4658.347249

Sum of electronic and thermal Energies= -4658.330981

Sum of electronic and thermal Enthalpies= -4658.330037

Sum of electronic and thermal Free Energies= -4658.394550

C -2.97553100 0.34941000 1.52032600

C -2.28096300 -0.59136300 0.72958800

C -2.22260500 -0.47104200 -0.67640700

C -2.76148000 0.66671600 -1.26751200

C -3.39786600 1.65092700 -0.48816400

C -3.50377600 1.48785900 0.89375700

Br -1.98440100 -2.44446100 1.57505900

H -1.71192700 -1.23486700 -1.26342300

H -3.09258000 0.19415400 2.59402000

H -4.01492600 2.23971400 1.50025600

H -3.82107400 2.53564400 -0.96881000

H -2.68434600 0.79329200 -2.35040300

Ni -0.52357700 -0.33275800 1.52376600

O 1.01536800 -1.09755800 0.59651700

O 0.21836200 1.44706400 1.68152500

C 1.32888400 1.80411800 1.16900400

C 2.01888200 -0.41530700 0.21813600

C 3.08153200 -1.19043800 -0.53120500

H 2.61808300 -1.68999300 -1.39733500

H 3.91530000 -0.56422100 -0.87564500

H 3.47357700 -1.98682000 0.12262600

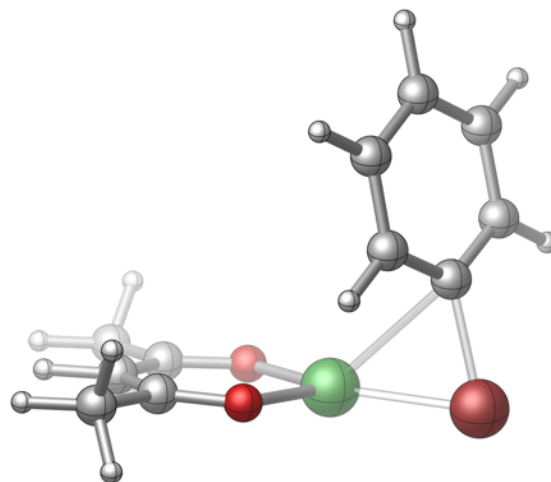
C 1.69846700 3.25582900 1.38174300

H 1.77581900 3.45268200 2.46373500

H 2.64190000 3.53881800 0.89617300

H 0.88502700 3.89360000 0.99959700  
C 2.20761400 0.96722700 0.44827700  
H 3.12192600 1.41793300 0.06166900  
 $\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01048663$   
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.7254964  
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.07173765  
DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4656.13047753174

<sup>2</sup>A'-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202302 (Hartree/Particle)

Thermal correction to Energy= 0.218638

Thermal correction to Enthalpy= 0.219582

Thermal correction to Gibbs Free Energy= 0.154463

Sum of electronic and zero-point Energies= -4658.336430

Sum of electronic and thermal Energies= -4658.320094

Sum of electronic and thermal Enthalpies= -4658.319150

Sum of electronic and thermal Free Energies= -4658.384269

C	-2.66543400	-0.54021800	1.31203400
O	-3.54977000	0.16165100	1.90193600
C	-2.75745500	-1.04436600	-0.00579100
H	-1.90838400	-1.62150400	-0.37388500
C	-3.85190100	-0.88199200	-0.88726100
O	-4.92957500	-0.25548300	-0.62718400
C	-3.77392900	-1.48775300	-2.27347800
H	-3.80891100	-0.67890700	-3.02257300
H	-2.86623300	-2.08441200	-2.43690400
H	-4.66111800	-2.11855300	-2.44475900
C	-1.42598100	-0.83438100	2.13127200
H	-1.71780700	-1.37441000	3.04723200
H	-0.67674900	-1.42635100	1.58862700
H	-0.97157900	0.11643800	2.45493400
Ni	-5.19980300	0.62850900	1.07948300
C	-6.10163200	2.98309900	0.22911100
C	-5.54528100	2.78780600	-1.03909300
C	-5.83997500	4.12340300	0.99130100
C	-4.67424300	3.75969200	-1.53660500
H	-5.75086700	1.87337400	-1.59511700
C	-4.96455900	5.08430900	0.47191200
H	-6.29762700	4.25476200	1.97282800

C -4.38118000 4.90613300 -0.78654100  
H -4.21726900 3.61269900 -2.51828900  
H -4.74398900 5.97824200 1.06040800  
H -3.69838300 5.66021300 -1.18447700  
Br -7.38844100 1.67802400 0.90565300

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01078544$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.7247785

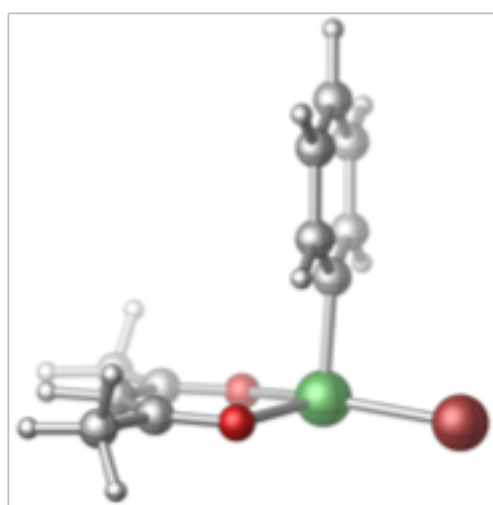
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.07196787

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4656.14180760118

<sup>2</sup>B



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202983 (Hartree/Particle)

Thermal correction to Energy= 0.219779

Thermal correction to Enthalpy= 0.220724

Thermal correction to Gibbs Free Energy= 0.154827

Sum of electronic and zero-point Energies= -4658.374184

Sum of electronic and thermal Energies= -4658.357387

Sum of electronic and thermal Enthalpies= -4658.356443

Sum of electronic and thermal Free Energies= -4658.422339

C -3.02059400 -0.20232900 1.26979400

O -4.07185600 0.32063600 1.77961800

C -2.95186100 -0.78793100 -0.00600300

H -1.99375400 -1.19622900 -0.32547500

C -4.04737800 -0.91164300 -0.88740700

O -5.22036900 -0.47009500 -0.65790600

C -3.86952100 -1.60621900 -2.21422100

H -4.05556200 -0.87921700 -3.02177500

H -2.86939100 -2.04016200 -2.34084500

H -4.63093300 -2.39592900 -2.31225800

C -1.80066000 -0.15896100 2.15343300

H -2.04085200 -0.61999500 3.12457700

H -0.93424800 -0.66604800 1.71000300

H -1.54579800 0.89479300 2.35266900

Ni -5.71248500 0.61864700 0.87298400

C -5.19472900 2.32978100 0.23203000

C -5.19034500 2.57300000 -1.13601600

C -4.75413100 3.25115500 1.17378500

C -4.69931500 3.80998800 -1.57972800

H -5.55481400 1.82299900 -1.83842000

C -4.26938600 4.48369700 0.71022200

H -4.76930400 3.02260600 2.24103100

C -4.24174700 4.76093100 -0.66091600  
H -4.68212400 4.02467100 -2.65162900  
H -3.91042100 5.22314300 1.43107900  
H -3.86301800 5.72289400 -1.01479200  
Br -7.96348000 1.16584000 0.81094000

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01656274$

UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4659.7503056

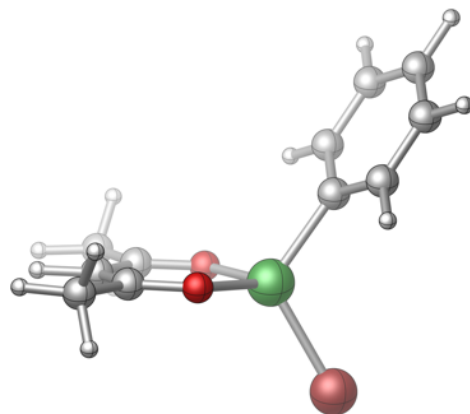
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4659.08182533

DLPNO-CCSD(T)/def2-TZVPP-gas// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4656.15666595836

**<sup>4</sup>B**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202208 (Hartree/Particle)

Thermal correction to Energy= 0.219348

Thermal correction to Enthalpy= 0.220292

Thermal correction to Gibbs Free Energy= 0.151895

Sum of electronic and zero-point Energies= -4658.348616

Sum of electronic and thermal Energies= -4658.331476

Sum of electronic and thermal Enthalpies= -4658.330532

Sum of electronic and thermal Free Energies= -4658.398929

C -2.53491700 -0.04438600 0.44720700

O -3.45994600 0.83791000 0.44218300

C -2.67146600 -1.36031800 -0.03946100

H -1.80695100 -2.01811300 0.03895100

C -3.83428900 -1.88275600 -0.63716800

O -4.92384800 -1.23471600 -0.81410200

C -3.86040600 -3.30632900 -1.12795900

H -2.89759000 -3.81763200 -1.00407400

H -4.64043000 -3.85609300 -0.57657600

H -4.15140600 -3.31515500 -2.19015100

C -1.21873800 0.41733700 1.01577600

H -1.38537500 0.82720900 2.02416700

H -0.46996900 -0.38341500 1.05983800

H -0.83424800 1.24120400 0.39257500

Ni -5.22881600 0.54852000 -0.23443600

C -5.46095600 1.86181000 -1.72835600

C -6.55428600 1.78500400 -2.59514300

C -4.48140700 2.84937200 -1.87097100

C -6.66247600 2.71521700 -3.63763400

H -7.32014100 1.01517300 -2.46594800

C -4.59858000 3.77508300 -2.91779300

H -3.63388100 2.89464700 -1.18267100

C -5.68550100 3.70639500 -3.79706900

H -7.51247100 2.66496400 -4.32370400

H -3.83656200 4.54916800 -3.04445300  
H -5.77334100 4.43004600 -4.61156800  
Br -6.84790200 0.65981100 1.44015000

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01265173$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.7252207

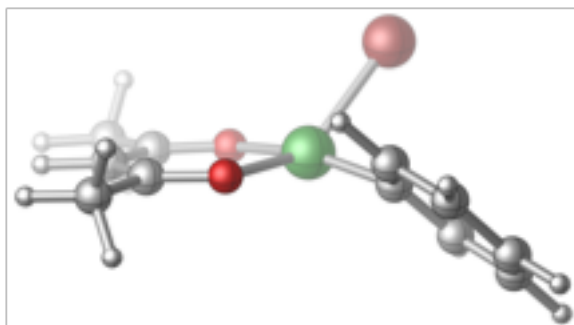
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4659.05442964

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4656.12009381385

<sup>2</sup>C



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.203033 (Hartree/Particle)

Thermal correction to Energy= 0.219788

Thermal correction to Enthalpy= 0.220732

Thermal correction to Gibbs Free Energy= 0.154904

Sum of electronic and zero-point Energies= -4658.365732

Sum of electronic and thermal Energies= -4658.348977

Sum of electronic and thermal Enthalpies= -4658.348033

Sum of electronic and thermal Free Energies= -4658.413861

C -3.04514500 -1.46216400 -1.03628300

O -3.57397700 -0.29461400 -1.12691700

C -3.74079000 -2.67377400 -0.94109100

H -3.15807600 -3.59216900 -0.88087600

C -5.15014200 -2.77270900 -0.87397300

O -5.94095200 -1.78133000 -0.92750100

C -5.80154400 -4.12411100 -0.71697400

H -6.48528600 -4.29041600 -1.56490200

H -5.07672100 -4.94657800 -0.66475800

H -6.41865100 -4.12034900 0.19580100

C -1.53706500 -1.45059300 -1.03019100

H -1.10420300 -2.45547900 -0.94566500

H -1.17843300 -0.97177500 -1.95544400

H -1.18747700 -0.82794600 -0.19108900

Ni -5.43346800 0.08164600 -1.20696600

C -5.08026700 1.95510000 -1.05476400

C -3.99209900 2.54261800 -1.69347000

C -5.86831600 2.65173300 -0.14046100

C -3.66331500 3.86730200 -1.37849400

H -3.39494400 1.96999400 -2.40527500

C -5.52134000 3.97451900 0.17688300

H -6.74517300 2.18891100 0.31987700

C -4.42433500 4.58064000 -0.44321000

H -2.80610100 4.34068800 -1.86514000

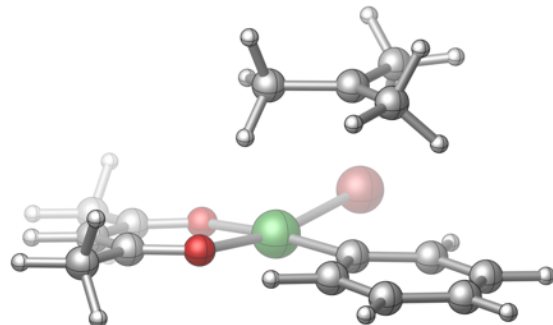
H -6.12317000 4.52971300 0.90150700

H -4.16500000 5.61492900 -0.20352600

Br -6.96918100 0.60457500 -2.86262600

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01550038$   
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.7424670  
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4659.07503520  
DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -4656.14246856537

### <sup>1</sup>C-Complex



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.321793 (Hartree/Particle)

Thermal correction to Energy= 0.345372

Thermal correction to Enthalpy= 0.346316

Thermal correction to Gibbs Free Energy= 0.266666

Sum of electronic and zero-point Energies= -4815.961361

Sum of electronic and thermal Energies= -4815.937782

Sum of electronic and thermal Enthalpies= -4815.936838

Sum of electronic and thermal Free Energies= -4816.016488

C	-2.39389900	-0.51158900	0.85223300
O	-3.16522000	0.50574200	0.78907600
C	-2.71944800	-1.79987800	0.41085300
H	-1.98340700	-2.59607000	0.51874000
C	-3.97684100	-2.09793400	-0.15803300
O	-4.89931000	-1.24925400	-0.33500800
C	-4.29154200	-3.50971500	-0.59734200
H	-5.16367100	-3.87591700	-0.03156700
H	-4.58235000	-3.49963100	-1.66004200
H	-3.45120300	-4.20207400	-0.45432900
C	-1.03817200	-0.23166100	1.45705400
H	-1.17136600	0.17955600	2.47068300
H	-0.40097400	-1.12445700	1.50836900
H	-0.52952400	0.54356600	0.86098000
Ni	-4.91195500	0.64605000	0.07844800
C	-4.99953900	2.50811500	0.65906200
C	-6.12341400	3.35603800	0.49553300
C	-3.91788700	3.08391300	1.37120400
C	-6.15450200	4.67390600	0.96723000
H	-7.00179300	2.98136900	-0.02977600
C	-3.93725600	4.39577400	1.85616400
H	-3.02687800	2.48050200	1.54351300
C	-5.05541000	5.21046400	1.64826300
H	-7.04731600	5.28507700	0.80167100
H	-3.06751200	4.78736500	2.39322600

H	-5.07368500	6.24054700	2.01495700
Br	-6.93045800	0.69849500	-1.24748000
C	-3.78836200	3.27854900	-2.07598600
C	-2.83200500	2.16456600	-1.96954600
H	-3.23484300	1.21227300	-2.33469500
H	-1.94892600	2.45282100	-2.57626200
H	-2.47887700	2.05448900	-0.93337400
C	-3.44417400	4.59300800	-1.51780700
H	-4.29727800	5.00047100	-0.94601400
H	-2.54094700	4.58697700	-0.89899700
H	-3.31618400	5.27966000	-2.37812800
C	-5.01093400	3.10074300	-2.82892700
H	-5.63665200	2.37048200	-2.23075300
H	-5.58515500	4.01886200	-2.99881900
H	-4.82514200	2.53406000	-3.75685000

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02574488$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.6391478

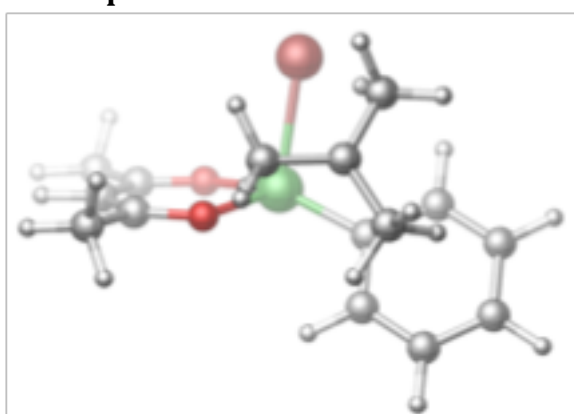
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82739558

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.64164587687

### <sup>3</sup>C-Complex



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.319982 (Hartree/Particle)

Thermal correction to Energy= 0.344022

Thermal correction to Enthalpy= 0.344966

Thermal correction to Gibbs Free Energy= 0.262848

Sum of electronic and zero-point Energies= -4815.972389

Sum of electronic and thermal Energies= -4815.948349

Sum of electronic and thermal Enthalpies= -4815.947405

Sum of electronic and thermal Free Energies= -4816.029523

C	-4.15367600	-1.41276300	2.26173000
O	-4.35116300	-0.19214200	2.52854000
C	-3.73006500	-1.91348600	1.00408200
H	-3.60922400	-2.99215100	0.90413100
C	-3.43197400	-1.12180400	-0.11561500
O	-3.50121800	0.15626900	-0.15480400
C	-2.99001800	-1.78044500	-1.40201000
H	-3.75492200	-1.60320000	-2.17660300
H	-2.06059400	-1.30531600	-1.75349600
H	-2.83458800	-2.86246800	-1.30039500
C	-4.40031600	-2.38362100	3.39540800
H	-3.77611000	-2.09581800	4.25669500
H	-5.45031300	-2.29249200	3.71819700
H	-4.19265200	-3.42806000	3.12751500
Ni	-4.26902900	1.34948400	1.25137300
C	-3.56788700	3.12039600	0.70705800
C	-4.28449200	4.33358900	0.78157700
C	-2.21259500	3.21856100	0.32091500
C	-3.69088500	5.57106100	0.49728400
H	-5.34321100	4.30433700	1.06194300
C	-1.59909800	4.44647300	0.04030500
H	-1.61930500	2.30107200	0.22867700
C	-2.34013100	5.63197100	0.12702100
H	-4.27959300	6.49178900	0.56253000

H	-0.54563900	4.48189000	-0.25554800
H	-1.87154100	6.59461500	-0.09671300
Br	-6.71638600	1.45750200	0.76756600
C	-4.65053800	2.79903500	-2.50779300
C	-3.54389200	3.74091800	-2.71956500
H	-2.55928900	3.26066300	-2.75496800
H	-3.75079400	4.19638600	-3.71156300
H	-3.56377800	4.56803800	-1.99440600
C	-5.95245700	3.28546600	-2.08653800
H	-6.12663400	2.82740200	-1.06818200
H	-6.02785300	4.37627600	-2.01441200
H	-6.76080600	2.84453600	-2.69183100
C	-4.43574000	1.37782900	-2.69432000
H	-4.00735400	1.03277100	-1.70193000
H	-5.35326300	0.79746600	-2.85305000
H	-3.65027900	1.15190900	-3.42980100

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02679875$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.6442305

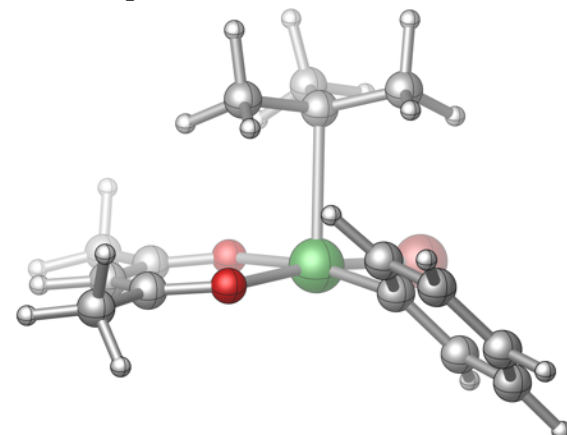
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82575635

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.65211594193

**<sup>1</sup>C''-Complex**



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.324614 (Hartree/Particle)

Thermal correction to Energy= 0.347556

Thermal correction to Enthalpy= 0.348500

Thermal correction to Gibbs Free Energy= 0.272116

Sum of electronic and zero-point Energies= -4815.971390

Sum of electronic and thermal Energies= -4815.948448

Sum of electronic and thermal Enthalpies= -4815.947503

Sum of electronic and thermal Free Energies= -4816.023888

C -2.56409800 -0.05189600 1.57097900

O -3.24509600 1.01608200 1.52880700

C -2.66238000 -1.10373900 0.63246700

H -2.02670000 -1.97852900 0.76495400

C -3.53212100 -1.07621300 -0.46587900

O -4.34521900 -0.13186300 -0.75350300

C -3.58261900 -2.24368800 -1.42137600

H -2.83716500 -3.01577200 -1.19078700

H -4.59053800 -2.68868300 -1.38594200

H -3.43024800 -1.87715000 -2.44866900

C -1.61501700 -0.17841800 2.73942100

H -2.20639500 -0.29521500 3.66331200

H -0.93160100 -1.03275600 2.64636500

H -1.03796200 0.75312900 2.84200900

Ni -4.64120100 1.52916100 0.15583700

C -5.73390400 2.01241500 -1.36217900

C -6.62198700 1.07590000 -1.91342500

C -5.52757900 3.21485500 -2.05731600

C -7.32192500 1.34630500 -3.09518700

H -6.75783100 0.10436400 -1.43632300

C -6.21143900 3.48070700 -3.25325500

H -4.83534000 3.96122800 -1.66685400

H -8.01675100 0.60121800 -3.49385500

H -6.03070100 4.42495700 -3.77559300  
 C -6.40956000 0.77147400 1.74155100  
 C -5.86660600 1.18283900 3.07146000  
 H -4.81572200 0.90108400 3.19701600  
 H -5.98730400 2.25399300 3.26155500  
 H -6.45979300 0.61732700 3.81893900  
 C -7.62285400 1.49747100 1.27064300  
 H -7.48049900 2.58460200 1.29571500  
 H -7.96088300 1.18106700 0.27916400  
 H -8.41877000 1.24568600 2.00229500  
 C -6.28906000 -0.67665700 1.40965800  
 H -6.53065400 -0.89919400 0.36531800  
 H -5.30444800 -1.08750700 1.66770200  
 H -7.03971500 -1.19230200 2.04333800  
 C -7.12004700 2.55461800 -3.77175500  
 H -7.66094000 2.76739300 -4.69778500  
 Br -4.74380600 3.68847500 1.10334400

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01640250$

UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4817.64084424

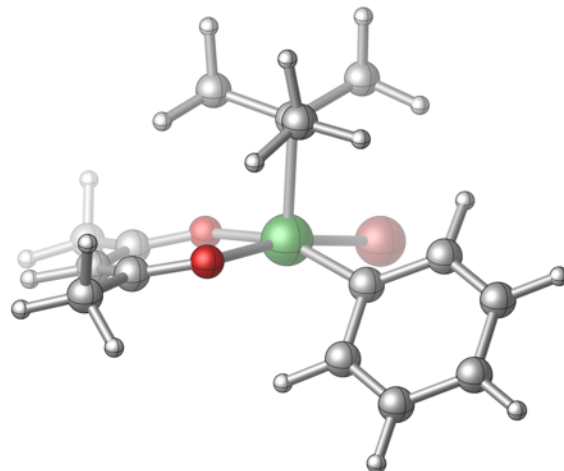
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4816.82646240

DLPNO-CCSD(T)/def2-TZVPP-gas// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4813.63811608447

<sup>1</sup>C'



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.325423 (Hartree/Particle)

Thermal correction to Energy= 0.347952

Thermal correction to Enthalpy= 0.348896

Thermal correction to Gibbs Free Energy= 0.273698

Sum of electronic and zero-point Energies= -4815.975127

Sum of electronic and thermal Energies= -4815.952597

Sum of electronic and thermal Enthalpies= -4815.951653

Sum of electronic and thermal Free Energies= -4816.026851

C -4.30533800 -0.34406200 0.82260900

O -5.46671300 0.16780700 0.64013700

C -3.50074900 -0.92262900 -0.16399200

H -2.52904700 -1.31736900 0.12959100

C -3.88626900 -1.00908500 -1.52163100

O -4.98712200 -0.58324800 -1.97837200

C -2.94568800 -1.61997800 -2.53197800

H -3.48526700 -2.38441400 -3.11230700

H -2.63518100 -0.83676300 -3.24324500

H -2.05215600 -2.06364000 -2.07366200

C -3.82097100 -0.25596200 2.24854800

H -2.85698700 -0.75844500 2.39998400

H -3.72549700 0.80600700 2.52813200

H -4.57859300 -0.69714900 2.91499900

Ni -6.50227900 0.19844700 -0.94819300

C -8.09615900 0.31863700 0.12195900

C -9.22403100 1.12080100 -0.08108500

C -8.10532000 -0.61378400 1.16896200

C -10.32913100 1.02564200 0.77250400

H -9.26388400 1.81980500 -0.91581100

C -9.22086600 -0.72184300 2.01314400

H -7.24535900 -1.26051000 1.34753700

C	-10.33225000	0.10288000	1.82498600
H	-11.19653700	1.66954700	0.60193500
H	-9.20915400	-1.45434300	2.82537700
H	-11.19751200	0.02494000	2.48836200
Br	-7.85869200	-0.23305700	-2.81430900
C	-6.12142100	2.32226000	-1.09046200
C	-4.65707600	2.26823100	-1.45419300
H	-4.03287600	1.80718900	-0.68059400
H	-4.48784900	1.76506300	-2.41280300
H	-4.32700000	3.31993900	-1.56103000
C	-6.39950500	2.92315200	0.26541100
H	-7.47184500	3.01331100	0.47024700
H	-5.92118600	2.35653400	1.07135600
H	-5.96755000	3.94255300	0.25597000
C	-6.97276700	2.88665500	-2.19907600
H	-6.73225300	2.45386700	-3.17629900
H	-8.04707600	2.78417000	-2.01539400
H	-6.75013600	3.97167700	-2.23111000

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01592820$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.64556732

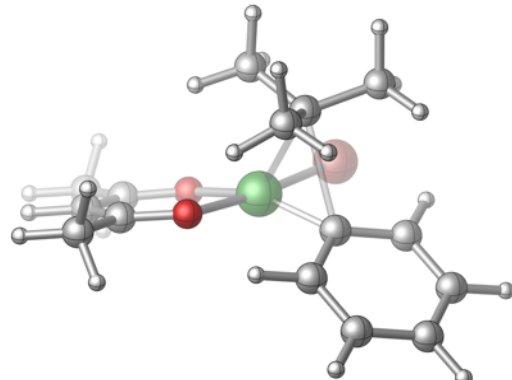
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82999570

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.66328229734

<sup>1</sup>C'-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.325398 (Hartree/Particle)

Thermal correction to Energy= 0.347305

Thermal correction to Enthalpy= 0.348249

Thermal correction to Gibbs Free Energy= 0.274271

Sum of electronic and zero-point Energies= -4815.962361

Sum of electronic and thermal Energies= -4815.940454

Sum of electronic and thermal Enthalpies= -4815.939510

Sum of electronic and thermal Free Energies= -4816.013487

C	-4.07305100	-0.47179400	0.72299900
O	-5.21739300	0.05547100	0.50807100
C	-3.31981100	-1.16659600	-0.23020200
H	-2.35284000	-1.57653600	0.05776300
C	-3.79081300	-1.37457500	-1.54133500
O	-4.89846000	-0.94266300	-1.98728700
C	-2.96580700	-2.16396700	-2.52823400
H	-3.54296700	-3.04484700	-2.85257600
H	-2.79267100	-1.54729000	-3.42439100
H	-2.00315300	-2.49127900	-2.11454000
C	-3.55362000	-0.28675400	2.12694000
H	-2.57966800	-0.76748100	2.28527800
H	-3.46679600	0.79119100	2.33909100
H	-4.28610700	-0.69772300	2.83994900
Ni	-6.24064000	0.13838100	-1.10969200
C	-7.89699300	0.65210700	-0.09676100
C	-9.14944900	0.79710000	-0.71833000
C	-7.87409500	0.28009600	1.26002400
C	-10.33457300	0.54289500	-0.02069300
H	-9.20968600	1.09428200	-1.76283400
C	-9.05935700	0.02313200	1.95769500
H	-6.92269700	0.18118000	1.77724700
C	-10.29717800	0.15377400	1.32188800
H	-11.29244300	0.65356700	-0.53631500
H	-9.00771500	-0.27860500	3.00751500

H	-11.22395300	-0.04056900	1.86803200
Br	-7.25628200	0.20116400	-3.31752100
C	-6.49527800	2.48171500	-0.62929300
C	-5.17178100	2.47401100	-1.38310800
H	-4.37219700	1.92770600	-0.86779100
H	-5.26263500	2.13048600	-2.42187300
H	-4.85246400	3.53388100	-1.42398900
C	-6.33285300	2.87239800	0.81735300
H	-7.29265800	2.95925100	1.33988000
H	-5.67778800	2.17634200	1.35265400
H	-5.84606500	3.86603400	0.82104900
C	-7.56261900	3.24373600	-1.37123800
H	-7.76831100	2.80584400	-2.35563500
H	-8.49221000	3.33671100	-0.79839800
H	-7.16021300	4.26105800	-1.53518800

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01578531$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.63196236

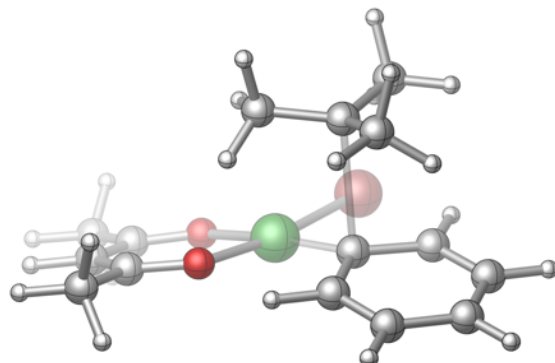
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82172077

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.65006456835

<sup>1</sup>C-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.323859 (Hartree/Particle)

Thermal correction to Energy= 0.346214

Thermal correction to Enthalpy= 0.347158

Thermal correction to Gibbs Free Energy= 0.271504

Sum of electronic and zero-point Energies= -4815.958572

Sum of electronic and thermal Energies= -4815.936217

Sum of electronic and thermal Enthalpies= -4815.935273

Sum of electronic and thermal Free Energies= -4816.010927

C	-2.31677600	-0.52043500	0.71526300
O	-3.09316900	0.48973200	0.61191500
C	-2.66193400	-1.83578400	0.38478000
H	-1.92370800	-2.62590900	0.51776900
C	-3.94469700	-2.16533600	-0.10222400
O	-4.87180500	-1.32585600	-0.30197000
C	-4.28888000	-3.60301700	-0.41523600
H	-5.11266400	-3.92526100	0.24255800
H	-4.66226100	-3.66774500	-1.44950100
H	-3.43791500	-4.28519100	-0.28750900
C	-0.93617200	-0.19863500	1.23552200
H	-1.02328100	0.29140100	2.21861600
H	-0.29783400	-1.08722100	1.32771500
H	-0.45437600	0.52495600	0.55767100
Ni	-4.86499600	0.58582400	-0.04958400
C	-4.90608600	2.52000300	0.35886900
C	-6.04267300	3.35988500	0.18257800
C	-3.91875200	3.02813100	1.25252500
C	-6.18028700	4.59194300	0.82211500
H	-6.83748700	3.03000000	-0.48546600
C	-4.04636800	4.25246500	1.90523600
H	-3.02458700	2.43032400	1.42612500
C	-5.17815400	5.04973800	1.68712100
H	-7.07112600	5.20082000	0.64448300
H	-3.25862500	4.59507100	2.58225300
H	-5.27734800	6.01794800	2.18550100

Br -7.00945500 0.52068600 -1.13207000  
 C -3.84272600 3.24560100 -1.79453200  
 C -2.85038500 2.14864400 -1.92283300  
 H -3.29711300 1.20688400 -2.26338500  
 H -2.12888200 2.49671900 -2.69059600  
 H -2.29171800 1.98245100 -0.99492400  
 C -3.34809200 4.57349000 -1.35644300  
 H -4.12468400 5.16280300 -0.84725200  
 H -2.44764300 4.51311700 -0.73492000  
 H -3.10468400 5.12032300 -2.28966100  
 C -5.02868200 3.20251600 -2.68368200  
 H -5.59850300 2.27022600 -2.55158300  
 H -5.68496100 4.07287000 -2.56557700  
 H -4.62680000 3.19333100 -3.71704800

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02375768$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4817.63659718

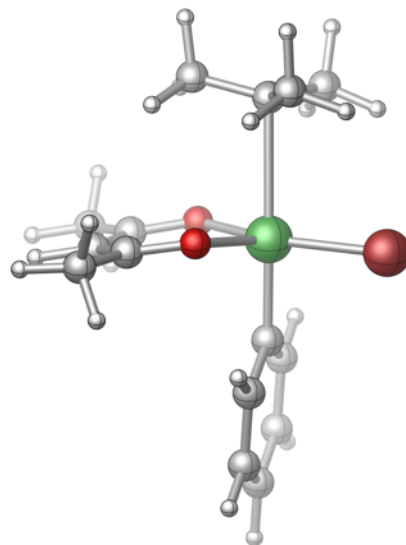
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4816.82544584

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4816.801688160

<sup>3</sup>C'



Zero-point correction= 0.322394 (Hartree/Particle)

Thermal correction to Energy= 0.346206

Thermal correction to Enthalpy= 0.347150

Thermal correction to Gibbs Free Energy= 0.265944

Sum of electronic and zero-point Energies= -4815.969566

Sum of electronic and thermal Energies= -4815.945754

Sum of electronic and thermal Enthalpies= -4815.944810

Sum of electronic and thermal Free Energies= -4816.026016

C -2.71674100 0.15818900 0.46731300

O -3.68961800 0.76472700 1.01871000

C -2.84124300 -0.89722800 -0.46363200

H -1.92410400 -1.32440800 -0.86729900

C -4.05853200 -1.47163300 -0.87047100

O -5.21790200 -1.08617400 -0.49479200

C -4.06055500 -2.65359500 -1.80629400

H -3.05063800 -2.96868500 -2.09848800

H -4.58026500 -3.49553100 -1.32115400

H -4.64035000 -2.39382100 -2.70661200

C -1.34435300 0.63578500 0.87462200

H -1.27105600 0.62861200 1.97340500

H -0.53425200 0.03103100 0.44706900

H -1.22480200 1.68294100 0.55117000

Ni -5.57704200 0.53911500 0.46401600

C -5.46032000 1.54197700 -1.17637500

C -6.30204000 1.16259200 -2.21934600

C -4.51593300 2.55468800 -1.32304700

C -6.18500300 1.81631800 -3.45483700

H -7.03210300 0.36077100 -2.08662300

C -4.40562300 3.19862400 -2.56419000

H -3.87784800 2.84052900 -0.48566300

C	-5.23777300	2.83176700	-3.62770800
H	-6.83712900	1.52338800	-4.28243800
H	-3.66636900	3.99430900	-2.69264000
H	-5.14927800	3.33860400	-4.59196600
Br	-6.88502600	2.24706700	1.39887900
C	-5.85136200	-1.03027400	2.56522100
C	-5.44340900	-0.08607000	3.65131700
H	-4.42897300	0.30321700	3.47266700
H	-6.13525600	0.75959600	3.75236300
H	-5.41939200	-0.63468600	4.61627100
C	-7.30547000	-1.27937200	2.31401200
H	-7.90724800	-0.36629700	2.41787800
H	-7.47114900	-1.71116400	1.31364800
H	-7.68088500	-2.02395000	3.04851200
C	-4.90689700	-2.15583500	2.28389100
H	-5.21957800	-2.75774100	1.42086700
H	-3.87937800	-1.79936300	2.11999200
H	-4.87840500	-2.82348200	3.16979100

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01168905$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.63659718

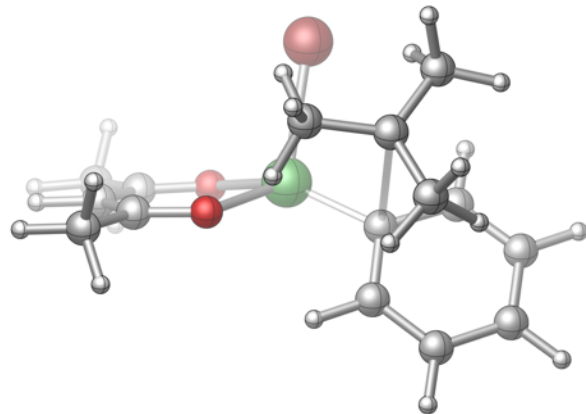
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82544584

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.66623270917

<sup>3</sup>C-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.320766 (Hartree/Particle)

Thermal correction to Energy= 0.344184

Thermal correction to Enthalpy= 0.345128

Thermal correction to Gibbs Free Energy= 0.264977

Sum of electronic and zero-point Energies= -4815.969306

Sum of electronic and thermal Energies= -4815.945888

Sum of electronic and thermal Enthalpies= -4815.944944

Sum of electronic and thermal Free Energies= -4816.025095

C -4.17569500 -1.41235000 2.26028700

O -4.45944600 -0.18562600 2.40579100

C -3.60900800 -1.98301900 1.09566100

H -3.41889300 -3.05637000 1.09739000

C -3.224444000 -1.25305200 -0.04437100

O -3.375333000 0.00339800 -0.20266800

C -2.56208900 -1.96938400 -1.20010100

H -3.17651000 -1.83288600 -2.10524300

H -1.58602000 -1.49942100 -1.40242800

H -2.41819300 -3.04292200 -1.01976200

C -4.47673300 -2.30100600 3.44709800

H -3.95559700 -1.90616400 4.33410100

H -5.55654600 -2.25081300 3.66312500

H -4.18688000 -3.34853900 3.29038100

Ni -4.38550600 1.22748000 0.99835600

C -3.69264400 2.99837000 0.39281200

C -4.33254400 4.23459000 0.63232500

C -2.31962000 3.05650300 0.05320800

C -3.65690000 5.45632800 0.53510600

H -5.39649800 4.23584400 0.89250400

C -1.62390700 4.26538500 -0.03657000

H -1.78874700 2.11986800 -0.15035700

C -2.29683700 5.47446400 0.19508000

H -4.18578800 6.39627300 0.71990000

H -0.56153800 4.27464600 -0.29885800  
H -1.76414900 6.42571200 0.11003600  
Br -6.83441500 1.24573900 0.56164400  
C -4.70877100 3.03216600 -2.30928500  
C -3.59189600 3.87447000 -2.76245400  
H -2.66149900 3.31955800 -2.92536100  
H -3.92879800 4.32085900 -3.72098200  
H -3.42884100 4.72688000 -2.08357800  
C -5.99990900 3.64971600 -1.99038000  
H -6.44553000 3.14142100 -1.11290700  
H -5.95342000 4.73686700 -1.86120500  
H -6.67907700 3.39382200 -2.82949800  
C -4.64388000 1.57410500 -2.48245000  
H -3.72960500 1.15204300 -2.03797500  
H -5.52063200 1.05891000 -2.07283800  
H -4.57697000 1.40007400 -3.57653100

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02583388$

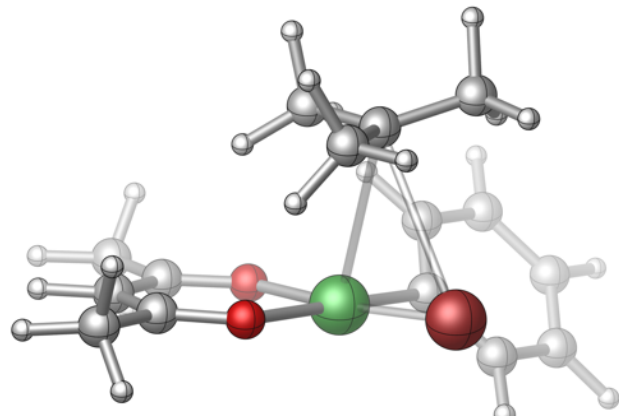
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.64143428

UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82379643

**<sup>1</sup>C''-TS**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.323194 (Hartree/Particle)

Thermal correction to Energy= 0.345704

Thermal correction to Enthalpy= 0.346648

Thermal correction to Gibbs Free Energy= 0.270392

Sum of electronic and zero-point Energies= -4815.963060

Sum of electronic and thermal Energies= -4815.940550

Sum of electronic and thermal Enthalpies= -4815.939606

Sum of electronic and thermal Free Energies= -4816.015862

C	-2.61804000	-0.11587600	1.50279400
O	-3.28107000	0.96028700	1.36471500
C	-2.62710600	-1.17982700	0.57954600
H	-2.02814200	-2.06374500	0.79722900
C	-3.36956400	-1.15248000	-0.61512500
O	-4.10550200	-0.18872600	-1.00732800
C	-3.33683000	-2.33974000	-1.54909400
H	-2.68961400	-3.15038800	-1.18927200
H	-4.36200100	-2.72257200	-1.67947400
H	-2.99105500	-2.00886900	-2.54157800
C	-1.81107900	-0.22777000	2.77527900
H	-2.50250600	-0.29135000	3.63256000
H	-1.14990300	-1.10450300	2.78907900
H	-1.21601900	0.68866400	2.91009200
Ni	-4.45396200	1.44872600	-0.12523600
C	-5.56794100	1.92570100	-1.58861800
C	-6.50501600	1.00619300	-2.09445700
C	-5.46396400	3.16350700	-2.25127500
C	-7.32553600	1.31348300	-3.18889900
H	-6.57766400	0.01011700	-1.64810500
C	-6.26175600	3.47232000	-3.36084400
H	-4.75242000	3.90936400	-1.88744500
H	-8.05176700	0.57764300	-3.54881400
H	-6.14997500	4.44165400	-3.85710400

C	-6.55526900	1.09284500	2.13554300
C	-5.84629700	1.32869700	3.40960500
H	-4.76541100	1.18169700	3.26734700
H	-6.05186700	2.31252100	3.84399600
H	-6.18952900	0.53314900	4.10327300
C	-7.78354200	1.83570800	1.78339100
H	-7.82823400	2.83379700	2.23076800
H	-7.92518800	1.88416200	0.69458200
H	-8.61811200	1.22308900	2.18706900
C	-6.34665000	-0.21996800	1.47810200
H	-6.65932400	-0.22871800	0.42893000
H	-5.33118700	-0.61240300	1.59721400
H	-7.03180800	-0.90144800	2.02842400
C	-7.20594000	2.55126300	-3.83021700
H	-7.83821100	2.79415900	-4.68894100
Br	-5.03147700	3.39640500	1.07445000

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02358562$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.63883416

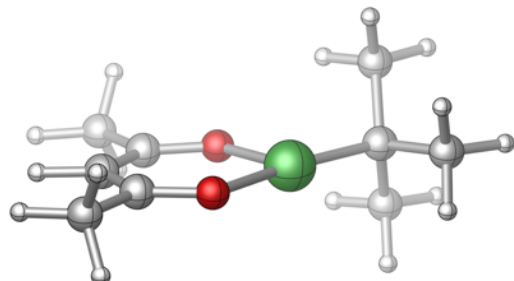
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.82668057

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.64502028160

**<sup>1</sup>D**



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.231479 (Hartree/Particle)

Thermal correction to Energy= 0.247258

Thermal correction to Enthalpy= 0.248202

Thermal correction to Gibbs Free Energy= 0.187876

Sum of electronic and zero-point Energies= -2010.679481

Sum of electronic and thermal Energies= -2010.663702

Sum of electronic and thermal Enthalpies= -2010.662758

Sum of electronic and thermal Free Energies= -2010.723084

C -3.04816600 -1.58856000 -0.57249400

O -3.60656400 -1.11182600 0.47582600

C -3.70755300 -2.17290100 -1.66924000

H -3.09228100 -2.53376600 -2.49332400

C -5.10941900 -2.32678100 -1.77738000

O -5.94271900 -1.95284300 -0.89416000

C -5.69777000 -2.97457500 -3.00927000

H -6.40228000 -2.27226900 -3.48390600

H -4.93822100 -3.27644600 -3.74245500

H -6.28308000 -3.85839000 -2.70744500

C -1.54066400 -1.48285400 -0.57486500

H -1.08171800 -1.90585300 -1.47800800

H -1.25437500 -0.42241300 -0.48396900

H -1.14133700 -1.99867100 0.31328800

Ni -5.45834700 -1.07899900 0.75321600

C -5.36148700 -0.20184800 2.44527100

C -6.85048200 -0.01437400 2.74608700

H -7.34072500 0.64322600 2.00600100

H -7.39456400 -0.97745200 2.75785000

H -7.01234600 0.44873200 3.74164700

C -4.68369100 -1.13415500 3.43934400

H -5.21201700 -2.10018300 3.51694800

H -3.64852000 -1.34277500 3.12473100

H -4.64536400 -0.69906200 4.46062000

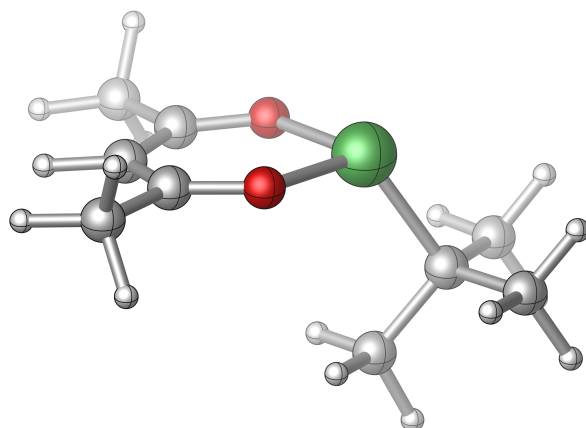
C -4.62877300 1.12534200 2.32230600

H -3.59170500 0.96134600 1.99039200

H -5.11526600 1.79012400 1.58796800

H -4.59442700 1.67192500 3.28840700  
 $\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01003141$   
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -2011.69926554  
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -2011.20902185  
DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)  
HF = -2009.74405799902

<sup>3</sup>D



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.230992 (Hartree/Particle)

Thermal correction to Energy= 0.246945

Thermal correction to Enthalpy= 0.247889

Thermal correction to Gibbs Free Energy= 0.185693

Sum of electronic and zero-point Energies= -2010.689470

Sum of electronic and thermal Energies= -2010.673517

Sum of electronic and thermal Enthalpies= -2010.672573

Sum of electronic and thermal Free Energies= -2010.734769

C 2.88294900 -0.62000300 -0.03246900

O 1.91881300 -0.10380500 0.62159600

C 2.75216000 -1.55531800 -1.08246100

H 3.67019200 -1.91116500 -1.55012700

C 1.53232200 -2.03671200 -1.60597800

O 0.37074000 -1.73345800 -1.17690700

C 1.55555100 -2.98975900 -2.77931900

H 2.57087600 -3.24883400 -3.10707900

H 1.01343900 -3.90930400 -2.50563400

H 1.00744700 -2.53559800 -3.62102700

C 4.26177700 -0.15780400 0.38065000

H 4.40134800 -0.36319800 1.45430900

H 5.06838900 -0.63674200 -0.18980100

H 4.32697900 0.93529300 0.25450700

Ni 0.04955000 -0.57161400 0.34993800

C -0.70065700 -1.48494200 1.93145700

C -0.49386500 -0.56712600 3.13475300

H 0.57087200 -0.30870000 3.26301700

H -1.05574900 0.37752100 3.03218300

H -0.83381100 -1.04675900 4.07922700

C -2.18021400 -1.78639700 1.70272400

H -2.77245400 -0.86448100 1.57134700

H -2.33162400 -2.41077700 0.80525400

H -2.62238800 -2.33825000 2.56176600

C 0.10745800 -2.77042800 2.07677300  
H -0.00376700 -3.41607400 1.18970100  
H 1.18320200 -2.55750200 2.19646200  
H -0.21282800 -3.36266100 2.96191300

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01003141$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2011.69926554

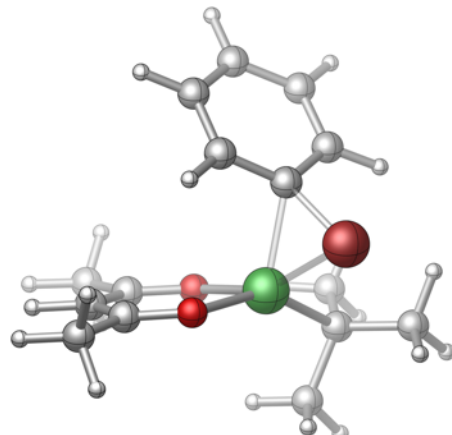
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2011.20902185

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2009.74405799902

<sup>1</sup>E-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.322368 (Hartree/Particle)

Thermal correction to Energy= 0.345098

Thermal correction to Enthalpy= 0.346042

Thermal correction to Gibbs Free Energy= 0.269644

Sum of electronic and zero-point Energies= -4815.950397

Sum of electronic and thermal Energies= -4815.927667

Sum of electronic and thermal Enthalpies= -4815.926723

Sum of electronic and thermal Free Energies= -4816.003121

C -3.00307000 -1.22231000 -0.44759300

O -3.69710000 -0.57016400 0.39855700

C -3.50532800 -2.02518000 -1.48790600

H -2.78502000 -2.54792500 -2.11693700

C -4.87949500 -2.24543200 -1.72767000

O -5.82153700 -1.67147400 -1.10282900

C -5.30048000 -3.21663400 -2.80853300

H -5.92334300 -2.68493600 -3.54627500

H -4.45170200 -3.68744000 -3.32225900

H -5.93451400 -3.99931100 -2.36133900

C -1.50734100 -1.09884400 -0.25894100

H -0.93093600 -1.64707800 -1.01580000

H -1.23129700 -0.03259200 -0.29008900

H -1.23818200 -1.47080900 0.74304300

Ni -5.61982400 -0.35771600 0.40848800

C -5.90709600 1.45818600 -0.58928400

C -5.36013200 1.26412600 -1.87017900

C -5.64389400 2.63195200 0.13258700

C -4.43037100 2.18592200 -2.35379000

H -5.63258900 0.38379100 -2.44920300

C -4.70978600 3.54169800 -0.36896500

H -6.14351500 2.81977400 1.08192500

C -4.09048700 3.32024600 -1.60481400

H -3.97233200 2.01315200 -3.33120000

H	-4.47578700	4.43687400	0.21312900
H	-3.36454500	4.03843500	-1.99252600
Br	-7.80850500	0.56785400	-0.21395000
C	-5.56419400	0.00627800	2.42335200
C	-6.89048000	0.51468900	2.95920100
H	-7.12606700	1.52936500	2.60747500
H	-7.72918200	-0.14284500	2.69049600
H	-6.83591200	0.55586700	4.06544200
C	-5.32913200	-1.45368400	2.79451400
H	-6.12726000	-2.11848900	2.42136800
H	-4.36071200	-1.82387700	2.43036800
H	-5.33366500	-1.55297300	3.89912500
C	-4.40834300	0.90721100	2.80702900
H	-3.47282800	0.58559900	2.33408300
H	-4.59640300	1.95486700	2.53150400
H	-4.27091300	0.87836700	3.90643200

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.00708735$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.61341725

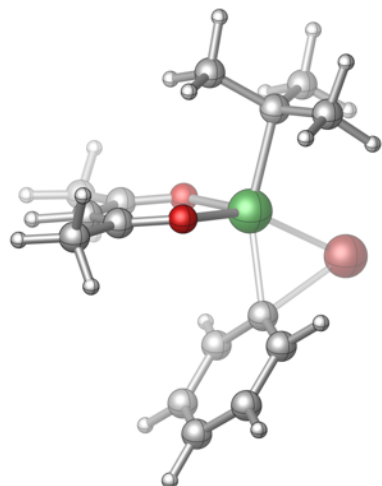
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.80636293

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.63643423474

<sup>3</sup>E-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.321470 (Hartree/Particle)

Thermal correction to Energy= 0.344473

Thermal correction to Enthalpy= 0.345417

Thermal correction to Gibbs Free Energy= 0.266848

Sum of electronic and zero-point Energies= -4815.950515

Sum of electronic and thermal Energies= -4815.927512

Sum of electronic and thermal Enthalpies= -4815.926568

Sum of electronic and thermal Free Energies= -4816.005138

C	-2.61196700	0.18283800	0.53402100
O	-3.57685300	0.71560400	1.16547700
C	-2.73328900	-0.77010100	-0.50178800
H	-1.81115800	-1.12268900	-0.96317200
C	-3.94146300	-1.31825200	-0.98262700
O	-5.10643500	-1.03825600	-0.55492800
C	-3.90068300	-2.33891400	-2.09702900
H	-2.88247200	-2.56208300	-2.44168100
H	-4.38065000	-3.26873200	-1.75113200
H	-4.49871500	-1.96485800	-2.94377800
C	-1.23439700	0.65037500	0.94795600
H	-1.12598000	0.53442400	2.03789200
H	-0.42350700	0.11249100	0.43932800
H	-1.14444000	1.72759400	0.73009000
Ni	-5.46981700	0.36182000	0.73736900
C	-5.74173500	1.86079900	-0.71154500
C	-6.72503600	1.46517500	-1.62506000
C	-4.55920600	2.48765400	-1.11722800
C	-6.45737200	1.59675900	-2.99090900
H	-7.66105800	1.02595800	-1.27433300
C	-4.31227400	2.61389600	-2.48716200
H	-3.83302600	2.81934500	-0.37499100

C	-5.25373500	2.16734500	-3.42426800
H	-7.19999000	1.25695800	-3.71758300
H	-3.37438700	3.06516200	-2.82153700
H	-5.05435000	2.27606500	-4.49283900
Br	-6.55256900	2.46067300	1.30294000
C	-5.88803100	-0.89029700	2.36315000
C	-5.63925700	-0.06327600	3.60582300
H	-4.60750700	0.32183800	3.62181300
H	-6.32698500	0.79121100	3.68432400
H	-5.77669800	-0.68835000	4.51294200
C	-7.34150600	-1.22579900	2.09787500
H	-7.98078500	-0.32864200	2.11583400
H	-7.46946400	-1.72707100	1.12416200
H	-7.72628600	-1.92120900	2.87410300
C	-4.95430600	-2.07471900	2.25267900
H	-5.13629500	-2.65515100	1.33788500
H	-3.90021200	-1.75808000	2.25673300
H	-5.09940500	-2.75296700	3.11923200

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.00793865$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4817.61456950

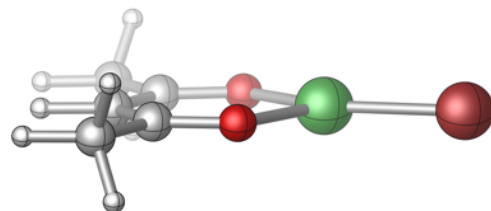
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4816.80487186

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -4813.63413962550

**<sup>1</sup>P<sub>A</sub>**



UB3LYP-D3/def2-SVP-CPCM(THF)

<S<sup>2</sup>> = 0.0000

Zero-point correction= 0.113187 (Hartree/Particle)

Thermal correction to Energy= 0.124416

Thermal correction to Enthalpy= 0.125360

Thermal correction to Gibbs Free Energy= 0.073570

Sum of electronic and zero-point Energies= -4426.989320

Sum of electronic and thermal Energies= -4426.978091

Sum of electronic and thermal Enthalpies= -4426.977147

Sum of electronic and thermal Free Energies= -4427.028936

C -2.69078700 -0.69335300 0.04535900

O -3.10573800 0.05502500 -0.90451100

C -3.52303100 -1.49102600 0.85184900

H -3.05780900 -2.09108800 1.63271400

C -4.92182300 -1.55970500 0.70982100

O -5.60859700 -0.91850000 -0.15687200

C -5.73285500 -2.44487400 1.62186500

H -6.27668200 -3.18374600 1.01159500

H -5.11746100 -2.96656100 2.36552200

H -6.49134400 -1.83242700 2.13483000

C -1.20113200 -0.67764400 0.26909300

H -0.69442000 -0.97056400 -0.66433700

H -0.88303500 0.35244000 0.49720500

H -0.88699300 -1.34500100 1.08144700

Ni -4.89234400 0.25158000 -1.44258200

Br -6.68390100 1.00741000 -2.74760800

ΔG<sub>solvation</sub><sup>UM06</sup>= -0.01991587

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4427.55542114

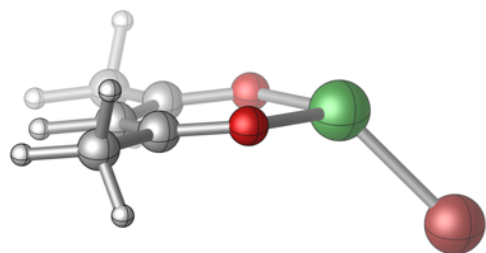
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4428.01914375

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4424.94554428170

$^3\text{P}_\text{B}$



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.112925 (Hartree/Particle)

Thermal correction to Energy= 0.124251

Thermal correction to Enthalpy= 0.125196

Thermal correction to Gibbs Free Energy= 0.071801

Sum of electronic and zero-point Energies= -4427.003874

Sum of electronic and thermal Energies= -4426.992547

Sum of electronic and thermal Enthalpies= -4426.991603

Sum of electronic and thermal Free Energies= -4427.044998

C -4.20757300 -0.40855400 2.63898700

O -5.19577900 0.33291600 2.31336500

C -3.26815500 -0.95076300 1.73918100

H -2.47361500 -1.57001100 2.15421500

C -3.24996600 -0.71386500 0.34959900

O -4.10948800 -0.01850100 -0.29141900

C -2.14189100 -1.29426900 -0.49341100

H -2.58428600 -1.91486600 -1.28911500

H -1.60283000 -0.47216300 -0.99154900

H -1.43116100 -1.89547600 0.08788200

C -4.07828000 -0.68320200 4.11637000

H -3.22289600 -1.32584400 4.36080800

H -3.98111300 0.27525500 4.65145700

H -5.00727100 -1.15576500 4.47399700

Ni -5.66814500 0.76594500 0.50679800

Br -7.78070400 0.13913100 -0.29285100

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.02500545$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4428.04619084

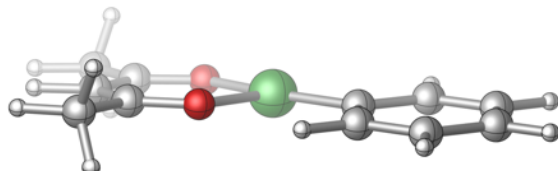
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4427.57712008

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4424.90484789000

**<sup>1</sup>P<sub>B</sub>**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.202049 (Hartree/Particle)

Thermal correction to Energy= 0.216762

Thermal correction to Enthalpy= 0.217706

Thermal correction to Gibbs Free Energy= 0.158091

Sum of electronic and zero-point Energies= -2084.454886

Sum of electronic and thermal Energies= -2084.440173

Sum of electronic and thermal Enthalpies= -2084.439229

Sum of electronic and thermal Free Energies= -2084.498844

C -3.48194900 -1.13581000 -0.48983600

O -4.33939100 -0.48231100 -1.17667500

C -3.70619700 -1.71587200 0.77299500

H -2.87856800 -2.24759800 1.24151500

C -4.93100400 -1.65226000 1.47069500

O -5.97348900 -1.06389700 1.03670900

C -5.06421500 -2.30948900 2.82250600

H -5.87443900 -3.05549000 2.78090800

H -4.13886900 -2.79642700 3.15644800

H -5.36615700 -1.55135400 3.56309700

C -2.12617400 -1.26197300 -1.14288800

H -2.24489600 -1.74270900 -2.12716400

H -1.72101800 -0.25367500 -1.32600300

H -1.41193800 -1.83698000 -0.53957300

Ni -6.12012700 -0.15017800 -0.63501200

C -6.64257400 0.84499400 -2.17224800

C -7.98622500 1.26568400 -2.23167400

C -5.80052100 1.20696400 -3.23827300

C -8.47194300 2.02031700 -3.30933100

H -8.67812600 1.00321400 -1.42072400

C -6.27457400 1.96044600 -4.31967100

H -4.75447400 0.88763800 -3.21292900

C -7.61378000 2.37016400 -4.35809600

H -9.52006300 2.33493900 -3.33048000

H -5.59874200 2.23079900 -5.13727500

H -7.98651200 2.95857200 -5.20135600

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.01662909$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

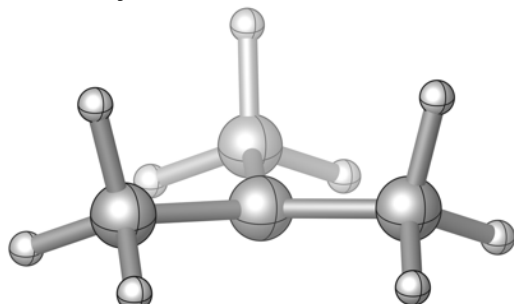
HF= -2085.51525078

UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2084.97451907

DLPNO-CCSD(T)/def2-TZVPP-gas// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2083.39540036421

**tert-Butyl radical**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.115524 (Hartree/Particle)

Thermal correction to Energy= 0.121853

Thermal correction to Enthalpy= 0.122797

Thermal correction to Gibbs Free Energy= 0.086138

Sum of electronic and zero-point Energies= -157.575218

Sum of electronic and thermal Energies= -157.568888

Sum of electronic and thermal Enthalpies= -157.567944

Sum of electronic and thermal Free Energies= -157.604604

C 0.05439000 1.47296400 1.28749000

H 1.15371300 1.47939600 1.19271400

H -0.30833600 2.51095700 1.19567200

H -0.16750400 1.15507600 2.33125000

C -2.04795000 0.72978000 0.00109100

H -2.34155300 1.79222600 -0.04829300

H -2.34037500 0.24384600 -0.94544000

H -2.67577200 0.26835000 0.79647700

C 0.05291600 -0.75676200 0.00135100

H -0.30729100 -1.19427100 -0.94544200

H 1.15254900 -0.68014400 -0.04738600

H -0.17378700 -1.50250000 0.79647000

C -0.58858200 0.56453700 0.28731600

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.00051217$

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -157.86998506

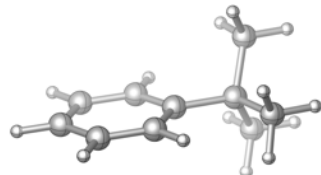
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -157.72008051

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -157.49166663063

**tBu-Ph**



Zero-point correction= 0.212567 (Hartree/Particle)

Thermal correction to Energy= 0.222196

Thermal correction to Enthalpy= 0.223140

Thermal correction to Gibbs Free Energy= 0.178263

Sum of electronic and zero-point Energies= -389.034497

Sum of electronic and thermal Energies= -389.024869

Sum of electronic and thermal Enthalpies= -389.023924

Sum of electronic and thermal Free Energies= -389.068801

C -2.58176900 1.21472200 0.38690900

C -1.22824900 1.17869200 -0.00409700

C -0.63158200 2.39658000 -0.37075900

C -1.35282800 3.59764200 -0.34863500

C -2.69310700 3.61187300 0.04204300

C -3.30571500 2.40859500 0.41120700

H -3.08621700 0.29102200 0.68019000

H 0.41272100 2.42820900 -0.68195400

H -0.85703000 4.52750500 -0.64054300

H -3.25607900 4.54860800 0.05950800

H -4.35445400 2.39895700 0.72043100

C -0.47089600 -0.16262400 -0.01661400

C -0.47838100 -0.76833300 1.40543100

H 0.05896700 -1.73089600 1.41423100

H 0.01717500 -0.09334900 2.12156900

H -1.50102700 -0.95357000 1.76740000

C 0.99279300 -0.00337300 -0.46382900

H 1.49247000 -0.98475200 -0.45386700

H 1.06693700 0.39712100 -1.48696000

H 1.55563000 0.66298200 0.20845000

C -1.17186700 -1.13447600 -0.99315200

H -2.21471300 -1.32907300 -0.69996500

H -1.17865300 -0.72448000 -2.01584700

H -0.64427600 -2.10218800 -1.01578600

$\Delta G_{\text{solvation}}^{\text{UM06}}$ = -0.00210233

UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -389.32908106

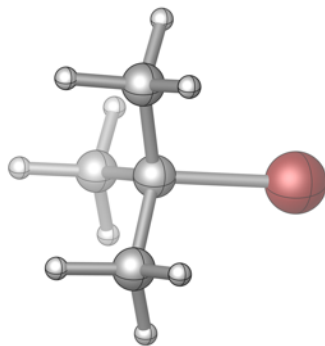
UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -389.32942074

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -388.77305156643

**tBu-Br**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.121383 (Hartree/Particle)

Thermal correction to Energy= 0.127998

Thermal correction to Enthalpy= 0.128942

Thermal correction to Gibbs Free Energy= 0.090831

Sum of electronic and zero-point Energies= -2731.509375

Sum of electronic and thermal Energies= -2731.502760

Sum of electronic and thermal Enthalpies= -2731.501816

Sum of electronic and thermal Free Energies= -2731.539927

C -0.80126600 0.79228800 -0.05867000

C -2.32358400 0.83072300 0.00721800

H -2.69950800 1.86412100 0.00978500

H -2.72789300 0.31834600 -0.88296400

H -2.70045000 0.31407300 0.90179900

C -0.25779100 -0.63025600 0.00764600

H -0.60744900 -1.18347300 -0.88130800

H 0.84182700 -0.64079900 0.00849400

H -0.61779200 -1.15675100 0.90345100

C -0.25801100 1.56088200 -1.25757100

H -0.62018200 2.59917700 -1.26690400

H 0.84159800 1.56916000 -1.26547000

H -0.60583300 1.06605800 -2.18093300

Br -0.11938200 1.75722800 1.61166300

$\Delta G_{\text{solvation}}^{\text{UM06}} = -0.00339442 \text{ shang}$

UB3LYP-D3/def2-TZVPP-CPCM(THF) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2732.12476199

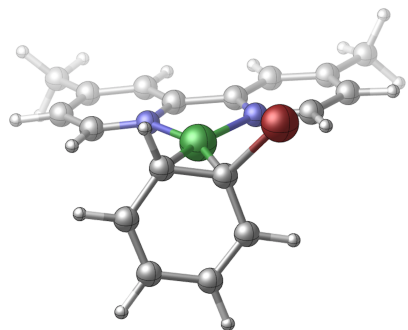
UM06/def2-TZVPP-CPCM(THF) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2731.85678123

DLPNO-CCSD(T)/def2-TZVPP-gas // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2730.27021223902

**Figure S4**  
 $^1\text{F}$



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.303924 (Hartree/Particle)

Thermal correction to Energy= 0.325242

Thermal correction to Enthalpy= 0.326187

Thermal correction to Gibbs Free Energy= 0.250047

Sum of electronic and zero-point Energies= -4886.916607

Sum of electronic and thermal Energies= -4886.895289

Sum of electronic and thermal Enthalpies= -4886.894345

Sum of electronic and thermal Free Energies= -4886.970485

C -2.35713200 0.66818000 1.26433000

C -2.03043400 -0.65370700 0.71070300

C -2.62076800 -1.02021400 -0.55416500

C -3.44410600 -0.14832400 -1.22864300

C -3.81708500 1.11177900 -0.66154400

C -3.31437600 1.48551300 0.56283800

Br -1.96632800 -2.28793700 2.02852200

H -2.38920200 -2.00165300 -0.97713300

H -2.27581600 0.83379700 2.34714300

H -3.66010700 2.41098000 1.03589400

H -4.52466100 1.75435400 -1.19237200

H -3.83293100 -0.43097800 -2.21173200

C 2.21375300 0.23800000 0.02640300

C 1.28434400 -1.89145700 0.08627200

C 2.50108900 -2.47921900 -0.24527300

C 3.63211900 -1.67205600 -0.45328500

C 3.46510300 -0.28824000 -0.30866900

C 1.93080300 1.68321100 0.20974400

C 0.27895400 3.23495100 0.72077800

C 1.17039700 4.29550300 0.59312500

C 2.51051000 4.03996100 0.25709500

C 2.87800700 2.70142300 0.06452300

H 0.38679600 -2.48706000 0.26939100

H 2.56598100 -3.56530600 -0.34084200  
H 4.31685000 0.37601000 -0.45956400  
H -0.77124200 3.39503000 0.97473900  
H 0.82053000 5.31726600 0.75559100  
H 3.90785100 2.45862100 -0.19952300  
N 1.13741300 -0.56421400 0.21612900  
N 0.64473000 1.95767100 0.53838200  
Ni -0.50805500 0.38037200 0.64864600  
C 3.49880600 5.16258500 0.09922700  
H 4.52387400 4.78948200 -0.03455000  
H 3.24147300 5.78012500 -0.77752400  
H 3.47932900 5.82800300 0.97684100  
C 4.96407200 -2.27782300 -0.80119200  
H 5.70898400 -1.51094000 -1.05612000  
H 5.35276300 -2.86437800 0.04800500  
H 4.86880100 -2.97073500 -1.65217600

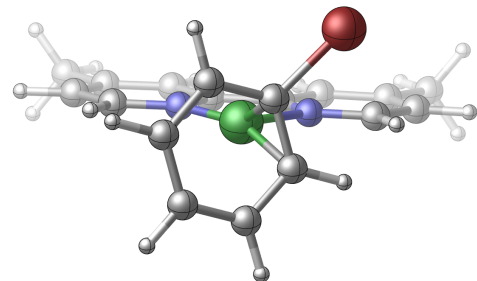
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.748303

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.625241

<sup>3</sup>F



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.303037 (Hartree/Particle)

Thermal correction to Energy= 0.324280

Thermal correction to Enthalpy= 0.325224

Thermal correction to Gibbs Free Energy= 0.249695

Sum of electronic and zero-point Energies= -4886.907048

Sum of electronic and thermal Energies= -4886.885805

Sum of electronic and thermal Enthalpies= -4886.884861

Sum of electronic and thermal Free Energies= -4886.960390

C -2.86004700 0.53399500 1.63679200

C -2.15805200 -0.57910400 1.08878500

C -2.26351900 -0.85908500 -0.30596100

C -3.10501400 -0.04467300 -1.10984200

C -3.79390400 1.02973300 -0.56253900

C -3.64814000 1.32896700 0.81157400

Br -1.52628800 -1.94341300 2.30708200

H -1.85251500 -1.78191100 -0.71809400

H -2.77869900 0.74475100 2.70382100

H -4.16773000 2.18873400 1.24078100

H -4.43714500 1.64913600 -1.19054800

H -3.21198200 -0.28990300 -2.16894500

C 2.15628700 0.25623200 -0.12325500

C 1.20467500 -1.89164300 -0.22112200

C 2.43029100 -2.49963900 -0.42579900

C 3.60169000 -1.68391100 -0.48870600

C 3.44226400 -0.31924600 -0.33446200

C 1.90132600 1.65745700 0.05116200

C 0.24662700 3.29745600 0.41826600

C 1.16875300 4.32630100 0.40858600

C 2.55027900 4.01179700 0.20563700

C 2.88970500 2.68307500 0.03207200

H 0.29296100 -2.49250900 -0.15567100

H 2.49366400 -3.58438700 -0.53297200

H 4.31863800 0.33053900 -0.37358800

H	-0.81895300	3.49727300	0.57204000
H	0.84330100	5.35821500	0.55466200
H	3.93614900	2.41393200	-0.12380200
N	1.03958100	-0.56588900	-0.07685900
N	0.57121700	2.00171800	0.24642200
Ni	-0.64075000	0.46304200	0.21118700
C	3.57627900	5.11229600	0.18684700
H	4.59281000	4.72480500	0.02373200
H	3.35484800	5.84380100	-0.60967300
H	3.57029400	5.67354700	1.13742000
C	4.94888900	-2.31570000	-0.71341400
H	5.75680500	-1.56926700	-0.72674000
H	5.17458600	-3.05339200	0.07598500
H	4.97190800	-2.86455500	-1.67084600

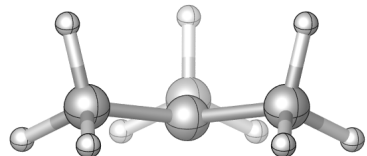
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7324304

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.61024

***tert*-butyl radical**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.115524 (Hartree/Particle)

Thermal correction to Energy= 0.121853

Thermal correction to Enthalpy= 0.122797

Thermal correction to Gibbs Free Energy= 0.086138

Sum of electronic and zero-point Energies= -157.575218

Sum of electronic and thermal Energies= -157.568888

Sum of electronic and thermal Enthalpies= -157.567944

Sum of electronic and thermal Free Energies= -157.604604

C -1.40592600 -5.80885900 -0.81665200

C -0.76293300 -4.90023200 -1.81663000

H 0.33639100 -4.89385900 -1.72186500

H -0.98485100 -5.21787700 -2.86045900

H -1.12562200 -3.86224700 -1.72457600

C -2.86529500 -5.64365300 -0.53040600

H -3.49311600 -6.10490200 -1.32589700

H -3.15773800 -6.12978300 0.41601800

H -3.15888000 -4.58121200 -0.48079900

C -0.76445100 -7.13022500 -0.53094900

H 0.33518400 -7.05363400 -0.48219000

H -1.12467000 -7.56791800 0.41575500

H -0.99116000 -7.87580100 -1.32621900

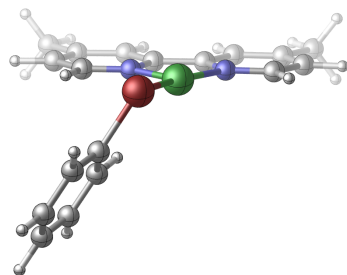
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-157.7200251

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-157.8699418

**<sup>1</sup>F'**



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 1.0136$  (before annihilation, 0.1453 after)

Zero-point correction= 0.303179 (Hartree/Particle)

Thermal correction to Energy= 0.324700

Thermal correction to Enthalpy= 0.325644

Thermal correction to Gibbs Free Energy= 0.249331

Sum of electronic and zero-point Energies= -4886.889735

Sum of electronic and thermal Energies= -4886.868214

Sum of electronic and thermal Enthalpies= -4886.867270

Sum of electronic and thermal Free Energies= -4886.943582

Ni	0.45097900	0.75422500	-0.51615600
C	-2.61923300	1.79943100	0.69437800
C	-3.80753400	2.52186400	0.78846000
C	-2.19830900	0.91052900	1.68156500
C	-4.60388600	2.33902600	1.92544600
H	-4.11043300	3.21117600	-0.00201500
C	-3.00841600	0.74202800	2.80967800
H	-1.26345000	0.35964700	1.56075100
C	-4.20732800	1.45287100	2.93326100
H	-5.53949000	2.89587200	2.01842400
H	-2.69559000	0.04769500	3.59335200
H	-4.83495500	1.31567000	3.81678000
Br	-1.50765300	2.03093300	-0.89197600
N	2.28718000	0.21033200	-0.31399000
C	4.84792900	-0.93207900	0.03142900
C	3.39401500	0.97541600	-0.30108000
C	2.41655000	-1.16425800	-0.14525500
C	3.71534600	-1.72516400	0.02368700
C	4.67452600	0.47726100	-0.13626800
N	0.03521700	-1.16120900	-0.33288500
C	-0.13784100	-3.96464000	-0.05814400
C	-1.15006500	-1.78968900	-0.37876300
C	1.19027000	-1.91125100	-0.16131900

C 1.08842900 -3.32614300 -0.02063400  
C -1.30380400 -3.15959400 -0.24831900  
H 3.22394300 2.04850100 -0.43475300  
H 3.81785500 -2.80464900 0.15126000  
H 5.53174800 1.15357700 -0.13713000  
H -2.01951200 -1.13964800 -0.52399200  
H 1.99565600 -3.91758100 0.11796900  
H -2.29792000 -3.60929800 -0.28986200  
C -0.27003300 -5.45646000 0.09238600  
H 0.70641800 -5.94504900 0.22688500  
H -0.90393100 -5.71038000 0.95979200  
H -0.75831100 -5.90086800 -0.79227800  
C 6.22827000 -1.50473400 0.20983000  
H 6.72258100 -1.07366300 1.09787100  
H 6.20833200 -2.59852100 0.32607300  
H 6.87116100 -1.26235900 -0.65425100

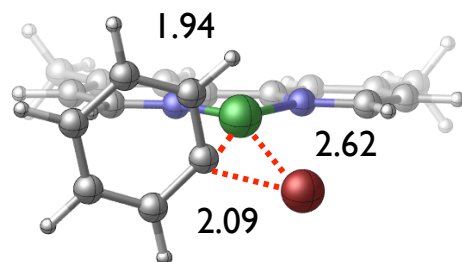
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7258715

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.6064849

<sup>1</sup>F-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.301918 (Hartree/Particle)

Thermal correction to Energy= 0.322858

Thermal correction to Enthalpy= 0.323802

Thermal correction to Gibbs Free Energy= 0.249700

Sum of electronic and zero-point Energies= -4886.898920

Sum of electronic and thermal Energies= -4886.877980

Sum of electronic and thermal Enthalpies= -4886.877036

Sum of electronic and thermal Free Energies= -4886.951138

C	-2.70242500	-0.07554900	1.35175300
C	-2.48593000	-0.39023000	-0.01498100
C	-3.05621300	0.39728800	-1.04546700
C	-3.73353300	1.55588600	-0.69934200
C	-3.89777500	1.92626000	0.65600600
C	-3.38781700	1.11697300	1.66513800
Br	-2.12202100	-2.41383100	-0.41069800
H	-2.92249600	0.10757100	-2.08902100
H	-2.45280700	-0.78360900	2.14542300
H	-3.54905800	1.37642100	2.71467400
H	-4.44490500	2.83755500	0.90673400
H	-4.13919000	2.19419300	-1.48866800
C	2.19635700	-0.18511600	0.13256200
C	1.61021300	-2.43581700	0.48578500
C	2.91959700	-2.85802600	0.37086400
C	3.94211800	-1.89307300	0.11678500
C	3.55615000	-0.56848700	-0.00168500
C	1.69785800	1.16386500	0.02495300
C	-0.22368200	2.51283400	0.03841500
C	0.51265500	3.67144500	-0.13362700
C	1.93122600	3.58101900	-0.23607400
C	2.50121600	2.32069200	-0.14974900
H	0.79904300	-3.14721000	0.66887500
H	3.16380100	-3.91739100	0.47412500
H	4.30957700	0.19363200	-0.20966200
H	-1.31199400	2.54355600	0.12135900
H	0.00621600	4.63712200	-0.19221500

H 3.58624200 2.21663100 -0.20901400  
N 1.23237700 -1.14498000 0.38190200  
N 0.32544800 1.28692800 0.10394600  
Ni -0.59521100 -0.44560400 0.41120500  
C 2.75756800 4.82441400 -0.42441900  
H 3.83290500 4.59969700 -0.47785000  
H 2.46931800 5.35042400 -1.35077400  
H 2.59448200 5.53480400 0.40408600  
C 5.37543700 -2.32903300 -0.02113200  
H 6.04717400 -1.47871200 -0.20990800  
H 5.72017300 -2.84371400 0.89234900  
H 5.49120200 -3.04913900 -0.84959000

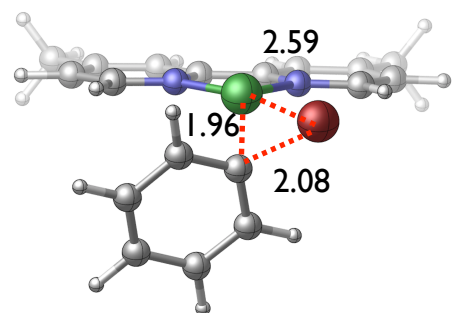
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7224721

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.5888467

<sup>3</sup>F-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.302378 (Hartree/Particle)

Thermal correction to Energy= 0.323155

Thermal correction to Enthalpy= 0.324099

Thermal correction to Gibbs Free Energy= 0.249623

Sum of electronic and zero-point Energies= -4886.899261

Sum of electronic and thermal Energies= -4886.878484

Sum of electronic and thermal Enthalpies= -4886.877540

Sum of electronic and thermal Free Energies= -4886.952015

C	-2.98196100	0.53763500	1.38884700
C	-2.33565200	-0.52608900	0.72079200
C	-2.36186400	-0.63350000	-0.68707300
C	-2.91143900	0.41192200	-1.42107900
C	-3.48624400	1.52278900	-0.77458900
C	-3.52523800	1.57702300	0.61953700
Br	-2.08238700	-2.26280700	1.84526800
H	-1.90111900	-1.49363900	-1.17414800
H	-3.04337300	0.55394600	2.47831900
H	-3.99362100	2.42496800	1.12564600
H	-3.91880800	2.33250400	-1.36618300
H	-2.89098200	0.36639800	-2.51298700
C	2.05129500	-0.07640600	0.27065900
C	1.46030100	-2.33969800	0.54765700
C	2.64594000	-2.78631300	-0.00664700
C	3.60420100	-1.82406500	-0.45249700
C	3.28789500	-0.48615500	-0.30556800
C	1.64408500	1.28764500	0.46008000
C	-0.06051900	2.71784700	1.23765500
C	0.65030600	3.85620700	0.90591100
C	1.94513700	3.71219100	0.31631600
C	2.41633100	2.43014600	0.10215400
H	0.70141500	-3.04605900	0.89799400
H	2.84014600	-3.85685000	-0.09874500
H	3.99931200	0.27131700	-0.63996600
H	-1.05796400	2.78637800	1.68256700

H 0.22330000 4.84404900 1.09008300  
H 3.39904400 2.29004700 -0.35189200  
N 1.14828700 -1.03991000 0.69664600  
N 0.39751300 1.46825400 1.04211400  
Ni -0.52639600 -0.24001900 1.41846100  
C 2.74293800 4.93376200 -0.05191900  
H 3.71690200 4.67299800 -0.49169400  
H 2.19380200 5.55869000 -0.77748000  
H 2.92340600 5.56765700 0.83347900  
C 4.90367700 -2.28019500 -1.05894700  
H 5.54204100 -1.43300400 -1.35015000  
H 5.46934000 -2.91019800 -0.35082700  
H 4.72425000 -2.89977300 -1.95472800

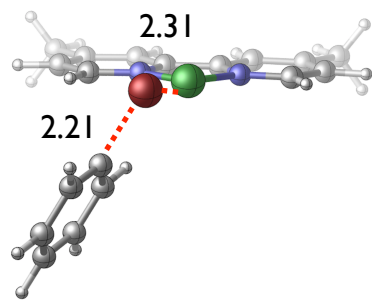
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7206594

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.6028485

<sup>1</sup>F'-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.302070 (Hartree/Particle)

Thermal correction to Energy= 0.323321

Thermal correction to Enthalpy= 0.324265

Thermal correction to Gibbs Free Energy= 0.248135

Sum of electronic and zero-point Energies= -4886.884886

Sum of electronic and thermal Energies= -4886.863635

Sum of electronic and thermal Enthalpies= -4886.862690

Sum of electronic and thermal Free Energies= -4886.938820

Ni -3.25440100 1.69922200 -0.86461000

C -6.35073000 2.74589100 0.53308000

C -7.53839400 3.44711300 0.67417300

C -5.86220800 1.86132000 1.48043900

C -8.28028700 3.24455700 1.84917500

H -7.88972100 4.13539600 -0.10011600

C -6.61321900 1.66746500 2.64876800

H -4.91949200 1.33067400 1.30643200

C -7.81837600 2.35839500 2.83071200

H -9.22162300 3.78253300 1.99505000

H -6.25559300 0.97493500 3.41641200

H -8.40158400 2.20507500 3.74213700

Br -5.14157500 2.96186900 -1.30211900

N -1.43673700 1.14855000 -0.56103500

C 1.13037600 0.04417700 -0.13811000

C -0.33715600 1.93142800 -0.52312200

C -1.28515500 -0.21011700 -0.37455300

C 0.00054600 -0.76171700 -0.17187300

C 0.93910900 1.44332600 -0.31648700

N -3.65142800 -0.21949300 -0.60653600

C -3.85403800 -3.00901400 -0.27941300

C -4.84620800 -0.83417400 -0.65249500

C -2.52059100 -0.97139200 -0.40358000

C -2.61976500 -2.37340400 -0.23952300

C -5.00368800 -2.20043400 -0.49558900

H -0.51503800 3.00012000 -0.67276100

H 0.10638300 -1.83949400 -0.03613000  
H 1.78764300 2.13053100 -0.29576000  
H -5.70475700 -0.17691700 -0.81941800  
H -1.71623100 -2.96410700 -0.07912700  
H -6.00041000 -2.64452600 -0.53793000  
C -3.99416700 -4.49675500 -0.10296000  
H -3.02112100 -4.98599000 0.04915400  
H -4.63647800 -4.72965900 0.76334700  
H -4.47599200 -4.95299000 -0.98434100  
C 2.51010400 -0.51465700 0.07923900  
H 2.97467100 -0.07668900 0.97929200  
H 2.49559500 -1.60786000 0.19844700  
H 3.17260700 -0.26945500 -0.76839500

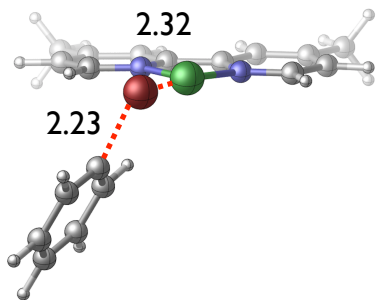
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.712787

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.5994182

<sup>3</sup>F'-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.301975 (Hartree/Particle)

Thermal correction to Energy= 0.323268

Thermal correction to Enthalpy= 0.324212

Thermal correction to Gibbs Free Energy= 0.246893

Sum of electronic and zero-point Energies= -4886.882832

Sum of electronic and thermal Energies= -4886.861540

Sum of electronic and thermal Enthalpies= -4886.860596

Sum of electronic and thermal Free Energies= -4886.937914

Ni -3.25263800 1.70106100 -0.95983300

C -6.35184300 2.70762400 0.47578100

C -7.55100900 3.38369900 0.63666300

C -5.83505700 1.82269900 1.40707900

C -8.27458100 3.15564700 1.81877700

H -7.92568500 4.07211900 -0.12667200

C -6.56771500 1.60345900 2.58285800

H -4.88479500 1.31169300 1.21578400

C -7.78363000 2.26932300 2.78604100

H -9.22429000 3.67396400 1.98147500

H -6.18755400 0.91068700 3.33952800

H -8.35277500 2.09594900 3.70278900

Br -5.16984200 2.93988300 -1.39971200

N -1.44440100 1.16785800 -0.57589000

C 1.12337200 0.07844800 -0.12970900

C -0.34815200 1.95332000 -0.55288900

C -1.28934600 -0.18483900 -0.37268100

C -0.00578600 -0.73140600 -0.15506900

C 0.93004300 1.47088600 -0.33441100

N -3.64223100 -0.21697000 -0.71040900

C -3.86427200 -2.97683700 -0.19545900

C -4.83578800 -0.83350300 -0.75824100

C -2.52536900 -0.95129400 -0.40433400

C -2.63166100 -2.33606400 -0.14348800

C -5.00027300 -2.18742000 -0.51776000

H -0.52837500 3.01905900 -0.71892100

H 0.10223800 -1.80749000 -0.00874000  
H 1.77655600 2.16077100 -0.32147800  
H -5.68585400 -0.18916700 -1.00093400  
H -1.73827800 -2.90972000 0.10926100  
H -5.99455300 -2.63537700 -0.57555400  
C -4.01310800 -4.44828300 0.08160900  
H -3.04713800 -4.92357100 0.30576000  
H -4.68840300 -4.61997400 0.93698100  
H -4.46063000 -4.96689600 -0.78309200  
C 2.50309700 -0.47512700 0.10090000  
H 2.48847200 -1.56499400 0.24673500  
H 3.16503000 -0.25058400 -0.75271500  
H 2.96709300 -0.01490200 0.98984600

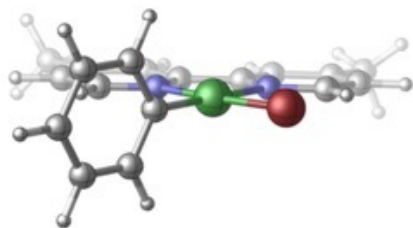
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.710583

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.5980175

**<sup>1</sup>G**



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.0000$

Zero-point correction= 0.306236 (Hartree/Particle)

Thermal correction to Energy= 0.327471

Thermal correction to Enthalpy= 0.328416

Thermal correction to Gibbs Free Energy= 0.251298

Sum of electronic and zero-point Energies= -4886.979456

Sum of electronic and thermal Energies= -4886.958220

Sum of electronic and thermal Enthalpies= -4886.957276

Sum of electronic and thermal Free Energies= -4887.034393

Ni -4.62164100 0.37276900 -0.35551800

C -6.48961200 0.30741100 -0.60607900

C -7.09110200 0.71952100 -1.80860200

C -7.32438700 -0.22393800 0.39359000

C -8.47025500 0.58409600 -2.01712600

H -6.47263100 1.15269500 -2.60117700

C -8.70484800 -0.36076100 0.19395800

H -6.89063800 -0.54663600 1.34577600

C -9.28439000 0.04067100 -1.01568600

H -8.91247300 0.90753700 -2.96479400

H -9.33112800 -0.78186000 0.98683500

H -10.36102100 -0.06552000 -1.17536300

Br -4.86562300 2.59801700 0.38673300

C -2.08305200 -0.93380400 -0.43222200

C -0.71910400 -1.20423500 -0.28347300

C 0.14074900 -0.22825800 0.23592100

C -0.43238600 1.00444300 0.58288700

C -1.79853600 1.20493800 0.40952800

C -3.06402100 -1.89823600 -0.97117700

C -2.72086900 -3.17719700 -1.42003300

C -3.69785200 -4.04180700 -1.92546000

C -5.01382400 -3.55700200 -1.95217400

C -5.29150600 -2.27595300 -1.49150600

H -0.31683100 -2.17600900 -0.56937000

H 0.18079400 1.81192400 0.98842400

H -2.28181300 2.14944200 0.66877300

H -1.68270700 -3.50593000 -1.37942800

H -5.83107600 -4.17356900 -2.33189100  
H -6.30644800 -1.88304700 -1.50502500  
N -2.61180200 0.26140400 -0.08511500  
N -4.34639000 -1.44945200 -1.00598000  
C 1.60662000 -0.49444200 0.43630000  
H 1.81183200 -0.69302500 1.50204700  
H 1.94610500 -1.36673200 -0.14000000  
H 2.21068100 0.37926600 0.14824900  
C -3.36195400 -5.41711900 -2.42902400  
H -3.53113800 -5.47644100 -3.51705900  
H -2.31471700 -5.68322400 -2.22918800  
H -4.01304700 -6.17284000 -1.96195100

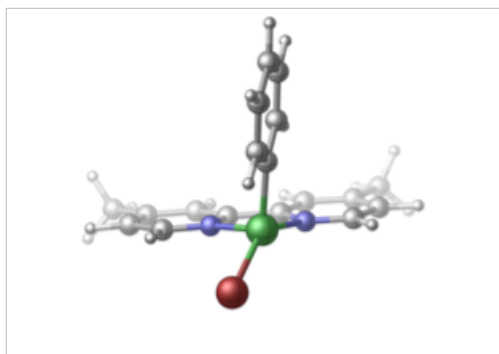
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.8048777

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.6907597

<sup>3</sup>G



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.305000 (Hartree/Particle)

Thermal correction to Energy= 0.326820

Thermal correction to Enthalpy= 0.327764

Thermal correction to Gibbs Free Energy= 0.248330

Sum of electronic and zero-point Energies= -4886.978664

Sum of electronic and thermal Energies= -4886.956844

Sum of electronic and thermal Enthalpies= -4886.955900

Sum of electronic and thermal Free Energies= -4887.035334

C	-1.73044200	0.93421700	-0.63476400
C	-1.87312800	0.01418300	0.42272300
C	-3.09150000	-0.69213400	0.48577300
C	-4.11710000	-0.48980600	-0.44888000
C	-3.94577100	0.43216300	-1.48845600
C	-2.74396600	1.14515500	-1.57994400
Br	-0.88450500	-2.21177200	3.23222000
H	-3.23786100	-1.42479400	1.28704600
H	-0.80136700	1.50807000	-0.73169700
H	-2.59751400	1.86676900	-2.39048300
H	-4.74106900	0.59286600	-2.22222600
H	-5.05158000	-1.05493100	-0.36868500
C	2.01065800	0.06705500	0.27402200
C	1.37737300	-2.16886900	0.19169000
C	2.45861300	-2.49492100	-0.62011900
C	3.36180300	-1.48973000	-1.00624600
C	3.11900900	-0.19089800	-0.54028200
C	1.66580800	1.40468700	0.82149200
C	0.17929400	2.58600300	2.15870500
C	0.84043600	3.78607700	1.92081700
C	1.97024400	3.79875800	1.08551300
C	2.37512800	2.57640700	0.53376900
H	0.65416600	-2.91549900	0.52937300
H	2.59867200	-3.52731600	-0.94686900
H	3.80224600	0.61390500	-0.81193400

H	-0.71171300	2.54155000	2.79011400
H	0.47461800	4.70615700	2.38100800
H	3.24287400	2.54531600	-0.12525700
N	1.16153900	-0.92126200	0.62231300
N	0.58416900	1.42588700	1.62949600
Ni	-0.43901900	-0.35359100	1.74233300
C	2.69511000	5.08093000	0.78436100
H	3.65064800	4.90126000	0.27232000
H	2.07555700	5.72408100	0.13705800
H	2.88953100	5.64723300	1.70840800
C	4.53562400	-1.80355900	-1.89196600
H	5.22460800	-0.95168800	-1.97632800
H	5.09354100	-2.67194100	-1.50816100
H	4.18856300	-2.06632300	-2.90518900

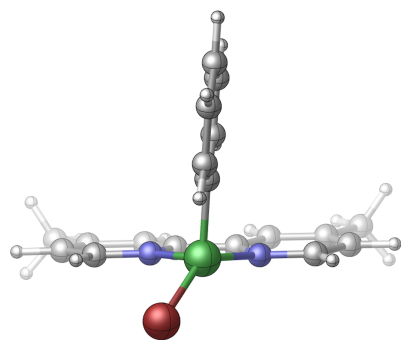
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7899161

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.6853609

**<sup>1</sup>G'**



UB3LYP-D3/def2-SVP-CPCM(THF)

$\langle S^2 \rangle = 0.9508$  (before annihilation, 0.0349 after)

Zero-point correction= 0.304833 (Hartree/Particle)

Thermal correction to Energy= 0.326760

Thermal correction to Enthalpy= 0.327704

Thermal correction to Gibbs Free Energy= 0.247505

Sum of electronic and zero-point Energies= -4886.959643

Sum of electronic and thermal Energies= -4886.937717

Sum of electronic and thermal Enthalpies= -4886.936773

Sum of electronic and thermal Free Energies= -4887.016971

Ni 1.09190800 0.85609700 -1.70045300

C 2.38443500 0.71457900 -0.24778600

C 4.32361600 0.52823400 1.82293400

C 2.00852900 0.65376700 1.10673000

C 3.76251500 0.67514900 -0.53491700

C 4.72335100 0.58681800 0.48277100

C 2.95825500 0.55992200 2.13304300

H 0.94689300 0.67949200 1.37739800

H 4.09105500 0.70806600 -1.57887300

H 5.78794100 0.56005200 0.22846600

H 2.63351000 0.51232000 3.17757600

N -0.25402700 2.12891000 -0.78486400

C -2.03722900 3.59640700 0.80378900

C -0.08589800 3.44506300 -0.61045800

C -1.28901800 1.50693000 -0.18117900

C -2.19373900 2.21728500 0.61705300

C -0.94786200 4.21283200 0.16501000

N -0.34638900 -0.47557600 -1.12802500

C -2.39020200 -2.15220500 -0.22052400

C -0.31543500 -1.78422300 -1.40004300

C -1.36863100 0.04049700 -0.41723000

C -2.40540200 -0.77721600 0.04796700

C -1.30952600 -2.65450600 -0.96577900

H 0.77765700 3.88987600 -1.11133800

H -3.01999300 1.69972700 1.10459300  
H -0.76936800 5.28462200 0.27348500  
H 0.53822100 -2.12410100 -1.99223700  
H -3.23277300 -0.34994300 0.61481500  
H -1.24475100 -3.71689000 -1.20969400  
Br 1.96750300 0.21137900 -3.84444000  
H 5.06948400 0.45680800 2.61980200  
C -3.48105700 -3.06467300 0.26779900  
H -3.89910200 -3.65146300 -0.56545100  
H -3.07828400 -3.78603900 0.99785100  
H -4.29730700 -2.50768600 0.74835000  
C -2.98421500 4.39357400 1.65712600  
H -3.78647000 3.76897300 2.07386900  
H -2.44347800 4.86833200 2.49191800  
H -3.44307200 5.20520200 1.06971000

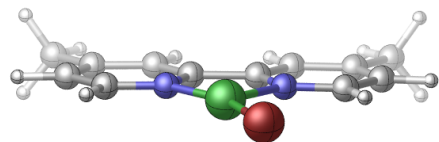
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4887.7735251

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4888.6678698

## <sup>1</sup>F'' (Ni-bromo species)



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.214708 (Hartree/Particle)

Thermal correction to Energy= 0.230876

Thermal correction to Enthalpy= 0.231820

Thermal correction to Gibbs Free Energy= 0.166322

Sum of electronic and zero-point Energies= -4655.579462

Sum of electronic and thermal Energies= -4655.563294

Sum of electronic and thermal Enthalpies= -4655.562349

Sum of electronic and thermal Free Energies= -4655.627848

Ni -3.99183200 1.30009300 -1.54653000

C -1.87359800 -0.39708200 -0.83451100

C -0.60463100 -0.81497600 -0.41773600

C 0.38038200 0.12429500 -0.09164500

C 0.02971800 1.48282700 -0.20377800

C -1.24637600 1.83242200 -0.62458000

C -2.97912600 -1.31121800 -1.19958100

C -2.90514100 -2.70937500 -1.17420400

C -4.01531200 -3.47922700 -1.54213400

C -5.17917400 -2.78948900 -1.92940000

C -5.18373000 -1.39947400 -1.93172800

H -0.37768600 -1.87943400 -0.34653600

H 0.75171100 2.26661600 0.03582700

H -1.54668200 2.87837300 -0.72245500

H -1.98301600 -3.20525100 -0.86858000

H -6.07708900 -3.33486700 -2.22820500

H -6.06484300 -0.82279400 -2.22618200

N -2.18757800 0.92199600 -0.93709400

N -4.11313300 -0.67488700 -1.57617200

C 1.75486100 -0.28579000 0.36089600

H 1.95781900 0.09097300 1.37696900

H 1.87445700 -1.37835300 0.36807700

H 2.52660300 0.14347000 -0.29878000

C -3.97959500 -4.98297800 -1.52479800

H -2.98906900 -5.36658100 -1.24261000

H -4.71990400 -5.37924000 -0.81035300

H -4.24149600 -5.38984900 -2.51492300

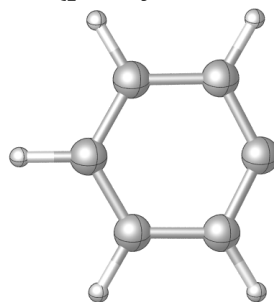
Br -5.83964900 2.56755300 -2.20535500

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4656.2756634

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF=-4656.9610627

**<sup>1</sup>F'' (phenyl radical)**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.087289 (Hartree/Particle)

Thermal correction to Energy= 0.091650

Thermal correction to Enthalpy= 0.092595

Thermal correction to Gibbs Free Energy= 0.059244

Sum of electronic and zero-point Energies= -231.315888

Sum of electronic and thermal Energies= -231.311527

Sum of electronic and thermal Enthalpies= -231.310583

Sum of electronic and thermal Free Energies= -231.343933

C -0.29103800 0.26151900 0.00004900

C 1.08781900 0.18926900 0.00053500

C 1.78056300 1.41499000 -0.00004200

C 1.07310400 2.62406400 -0.00104400

C -0.32767200 2.63227200 -0.00149100

C -1.04299800 1.41956500 -0.00093600

H 1.62938100 -0.76109600 0.00131300

H 2.87438700 1.41756800 0.00029600

H 1.61954800 3.57046000 -0.00149000

H -0.87218400 3.58094900 -0.00227600

H -2.13682400 1.41357900 -0.00127400

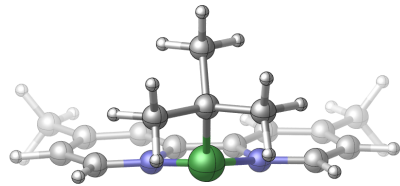
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-231.4580093

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-231.6566083

<sup>2</sup>J



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.331279 (Hartree/Particle)

Thermal correction to Energy= 0.351894

Thermal correction to Enthalpy= 0.352838

Thermal correction to Gibbs Free Energy= 0.280022

Sum of electronic and zero-point Energies= -2239.242341

Sum of electronic and thermal Energies= -2239.221726

Sum of electronic and thermal Enthalpies= -2239.220782

Sum of electronic and thermal Free Energies= -2239.293598

Ni	-4.10779300	1.25373400	-1.32019600
C	-5.28347400	2.10080600	0.03079000
C	-5.03401900	1.38406500	1.35564100
H	-5.25906000	0.30620400	1.28131900
H	-3.98154200	1.47642200	1.67430500
H	-5.66375000	1.79496700	2.17789100
C	-6.74299800	1.93257600	-0.39703800
H	-6.94980400	2.42389700	-1.36397600
H	-7.01521500	0.86793600	-0.50344000
H	-7.44565000	2.37383100	0.34714100
C	-4.92890800	3.58415100	0.15262900
H	-3.87438700	3.72864900	0.44643000
H	-5.08315600	4.12523800	-0.79738700
H	-5.55196400	4.09679500	0.92164600
C	-1.89191700	-0.41456100	-0.88502500
C	-0.61814100	-0.81574800	-0.41270300
C	0.35899600	0.12005500	-0.11030600
C	0.02631300	1.49591800	-0.28126800
C	-1.23753900	1.83555500	-0.71914600
C	-2.96893100	-1.31088500	-1.24418800
C	-2.91389500	-2.72433300	-1.16927600
C	-3.99501600	-3.50287200	-1.55282100
C	-5.16206000	-2.82501300	-2.01248100
C	-5.17490100	-1.44531200	-2.04011900
H	-0.40127800	-1.87823200	-0.28847900
H	0.75671600	2.27789200	-0.06274800
H	-1.53394200	2.88224600	-0.83746600
H	-2.00592900	-3.20789700	-0.80444100
H	-6.04307800	-3.38544700	-2.33253800

H -6.06335400 -0.89367900 -2.36210800  
N -2.18708600 0.92479600 -1.02438100  
N -4.11948700 -0.68522600 -1.67555800  
C 1.72370800 -0.27320800 0.38598300  
H 1.91667200 0.15250000 1.38561100  
H 1.83814000 -1.36510800 0.45040600  
H 2.51186200 0.11755900 -0.28004500  
C -3.96763700 -5.00572600 -1.48791700  
H -2.99793700 -5.38363700 -1.13244600  
H -4.75292300 -5.38446500 -0.81138900  
H -4.16727900 -5.44621100 -2.47965200

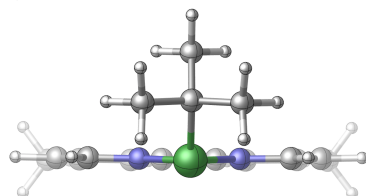
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2239.8687984

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2240.5868708

<sup>4</sup>J



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.331122 (Hartree/Particle)

Thermal correction to Energy= 0.351753

Thermal correction to Enthalpy= 0.352697

Thermal correction to Gibbs Free Energy= 0.279075

Sum of electronic and zero-point Energies= -2239.236155

Sum of electronic and thermal Energies= -2239.215524

Sum of electronic and thermal Enthalpies= -2239.214580

Sum of electronic and thermal Free Energies= -2239.288203

Ni	-4.11364400	1.25171300	-1.35136700
C	-5.30943000	2.15278900	-0.05189500
C	-5.12379100	1.41771900	1.27095000
H	-5.39083100	0.35173000	1.17915700
H	-4.07610200	1.46491900	1.61216600
H	-5.75620800	1.85139700	2.07662700
C	-6.75210500	2.03935200	-0.54491100
H	-6.89621400	2.53596100	-1.51987700
H	-7.06181300	0.98624900	-0.65716300
H	-7.46358700	2.51074500	0.16894000
C	-4.88469300	3.61585700	0.07013500
H	-3.83618700	3.70884700	0.40156800
H	-4.98128000	4.15545000	-0.88780900
H	-5.50788100	4.15995300	0.81427700
C	-1.91951800	-0.43473700	-0.77691700
C	-0.59133900	-0.83128600	-0.44384100
C	0.37256900	0.10003800	-0.10652300
C	-0.00177800	1.48126100	-0.10583800
C	-1.29654900	1.81858100	-0.45309700
C	-2.98536800	-1.32921200	-1.12487800
C	-2.87841100	-2.74941200	-1.18917800
C	-3.96492000	-3.53686600	-1.52063800
C	-5.20546100	-2.88212200	-1.80354600
C	-5.25574700	-1.50197000	-1.74635900
H	-0.33246100	-1.89167700	-0.45113800
H	0.71715700	2.25835400	0.16045200
H	-1.61984100	2.86445500	-0.47338200
H	-1.92065000	-3.22417300	-0.96819200
H	-6.09774700	-3.45525900	-2.06244500

H -6.18319900 -0.96266000 -1.96463600  
N -2.23720000 0.91623900 -0.78353100  
N -4.20098500 -0.73186300 -1.42625700  
C 1.77899500 -0.29547500 0.25316800  
H 2.03975500 0.05842200 1.26556500  
H 1.92119900 -1.38575800 0.22253200  
H 2.50625700 0.16658900 -0.43672600  
C -3.87769300 -5.03726600 -1.58960400  
H -2.86973500 -5.40369900 -1.34554800  
H -4.59329800 -5.50453600 -0.89112100  
H -4.14191300 -5.40041900 -2.59786800

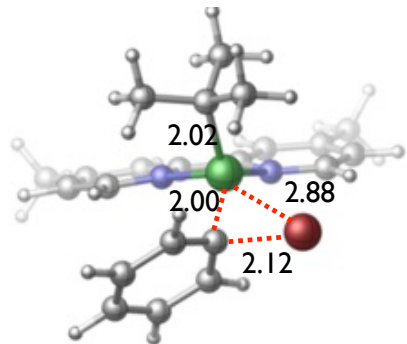
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2239.8575927

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2240.5786207

<sup>2</sup>K-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.422956 (Hartree/Particle)

Thermal correction to Energy= 0.450436

Thermal correction to Enthalpy= 0.451380

Thermal correction to Gibbs Free Energy= 0.361910

Sum of electronic and zero-point Energies= -5044.524684

Sum of electronic and thermal Energies= -5044.497204

Sum of electronic and thermal Enthalpies= -5044.496260

Sum of electronic and thermal Free Energies= -5044.585730

Ni -0.56887500 -0.10884500 0.03179700

C -2.00773300 0.73061800 -1.07367400

C -1.81809100 3.47023400 -1.69584400

C -2.92075100 1.65519000 -0.50062400

C -1.11674800 1.17633300 -2.09208400

C -1.00348100 2.54641200 -2.35490100

C -2.78829100 3.00870100 -0.78061000

H -3.67843900 1.30166500 0.19970900

H -0.51865100 0.45366200 -2.64898900

H -0.27773300 2.88665100 -3.09898500

H -3.44890000 3.72264300 -0.28033800

N 0.96824900 1.25462800 0.18035100

C 3.21273000 2.93003500 0.37307900

C 0.83008500 2.54873000 0.50132000

C 2.20297300 0.76608600 -0.07367600

C 3.34015400 1.58115100 0.02500900

C 1.91223300 3.41398800 0.60596400

N 1.02235400 -1.20962300 -0.72080700

C 3.36186200 -2.72388100 -1.08916100

C 0.95766700 -2.48795200 -1.12680000

C 2.23627000 -0.65439000 -0.49338800

C 3.41657000 -1.39275300 -0.66507600

C 2.08643100 -3.27123000 -1.32808200

H -0.19249800 2.89070500 0.66852000

H 4.32833000 1.17240700 -0.18817700

H	1.74447000	4.46123100	0.86657200
H	-0.04933300	-2.87875200	-1.29817400
H	4.38455800	-0.93335800	-0.46190100
H	1.97710300	-4.30344700	-1.66790300
Br	-2.75469400	-1.21241800	-1.48193200
C	-1.25942500	-0.30913100	1.91997700
H	-1.73267300	4.53737400	-1.91319300
C	4.60421000	-3.55062400	-1.27741800
H	5.51773900	-2.95597000	-1.13619400
H	4.62398100	-4.38736500	-0.55940500
H	4.62857200	-3.99435000	-2.28578400
C	4.40757900	3.83624900	0.49373300
H	4.53584500	4.16752900	1.53775800
H	5.33438400	3.33736200	0.17694300
H	4.27328200	4.74427900	-0.11569700
C	-0.10593600	-1.08166100	2.57143300
H	-0.28871200	-1.26294300	3.65582700
H	0.04439400	-2.07046200	2.10401900
H	0.85066700	-0.53464100	2.49801600
C	-2.54257900	-1.12746400	2.00560400
H	-3.39265400	-0.61222000	1.52877500
H	-2.44129500	-2.10488300	1.50598200
H	-2.83192800	-1.32398200	3.06326500
C	-1.44613300	1.02819800	2.62310500
H	-0.50872100	1.60684200	2.65959400
H	-2.20175500	1.65594800	2.12145600
H	-1.78013400	0.89735600	3.67747900

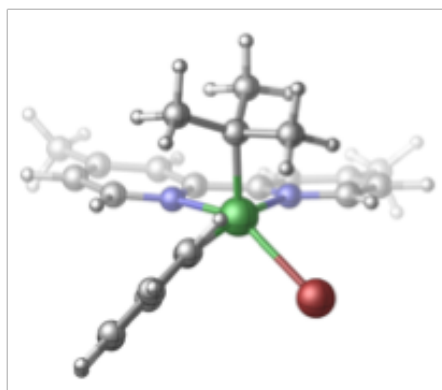
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HF=-5045.4850601

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5185177

<sup>2</sup>H



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.427053 (Hartree/Particle)

Thermal correction to Energy= 0.454642

Thermal correction to Enthalpy= 0.455586

Thermal correction to Gibbs Free Energy= 0.366581

Sum of electronic and zero-point Energies= -5044.574659

Sum of electronic and thermal Energies= -5044.547070

Sum of electronic and thermal Enthalpies= -5044.546126

Sum of electronic and thermal Free Energies= -5044.635131

Ni -0.56337000 -0.04631900 0.09841600

C -1.85804600 1.35728300 -0.24505500

C -3.52873200 3.40894600 -1.28522000

C -3.21390200 1.44004700 0.10238600

C -1.36915700 2.30647500 -1.16099100

C -2.18768500 3.32555400 -1.66969900

C -4.04026600 2.45236000 -0.40091300

H -3.65663600 0.70026200 0.77209900

H -0.33487500 2.25428800 -1.51272900

H -1.77097400 4.05105900 -2.37504500

H -5.09288800 2.48722500 -0.10395600

N 1.18278400 1.17021700 0.40025100

C 3.58866200 2.49028000 1.00364600

C 1.18840300 2.46208000 0.75018400

C 2.35080400 0.49932700 0.36745100

C 3.56597200 1.13467400 0.65549500

C 2.35389800 3.15589100 1.05841300

N 0.98567200 -1.44060000 0.01814400

C 3.18735400 -3.15570700 -0.31343600

C 0.81036500 -2.75503700 -0.17462800

C 2.24156800 -0.95071800 0.06963400

C 3.35492100 -1.78141500 -0.10891200

C 1.87055000 -3.64127200 -0.32900300

H 0.21530000 2.95470900 0.77867800

H	4.50024400	0.57419700	0.62972700
H	2.29688100	4.20930500	1.34012000
H	-0.22492700	-3.09449300	-0.22423700
H	4.36042800	-1.36177600	-0.09062700
H	1.66697900	-4.70462200	-0.47039500
Br	-1.71056300	-1.32864000	-1.79047600
C	-1.01599900	-0.24129500	2.08666000
H	-4.17092500	4.20206200	-1.67755800
C	4.35962500	-4.07147500	-0.52983600
H	5.30517400	-3.60717500	-0.21595500
H	4.22852600	-5.01710600	0.01787800
H	4.44652500	-4.32663800	-1.59967900
C	4.87320000	3.21007200	1.30667700
H	4.79438000	3.77049200	2.25134900
H	5.72470600	2.51890000	1.37659300
H	5.09240900	3.94510400	0.51426800
C	0.22707200	-0.77930600	2.79634400
H	-0.00940400	-0.91046400	3.87057900
H	0.55316300	-1.75566600	2.41804000
H	1.07501900	-0.08110100	2.74214300
C	-2.10117100	-1.30721300	2.02841900
H	-2.99088000	-0.97953000	1.47438100
H	-1.73718700	-2.23009000	1.55095900
H	-2.42666400	-1.57395900	3.05447600
C	-1.45739800	1.03009500	2.80018200
H	-0.67484600	1.80304400	2.75646200
H	-2.38075100	1.46045700	2.39947000
H	-1.63209800	0.80641000	3.87152200

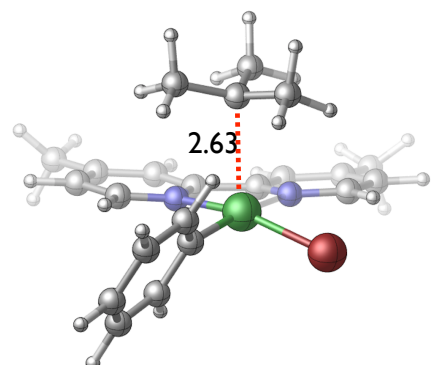
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.5318625

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.574292

<sup>2</sup>H-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.425172 (Hartree/Particle)

Thermal correction to Energy= 0.452637

Thermal correction to Enthalpy= 0.453581

Thermal correction to Gibbs Free Energy= 0.364426

Sum of electronic and zero-point Energies= -5044.568901

Sum of electronic and thermal Energies= -5044.541436

Sum of electronic and thermal Enthalpies= -5044.540492

Sum of electronic and thermal Free Energies= -5044.629648

Ni -0.46974300 0.02068700 -0.41675500

C -1.67999300 1.47643000 -0.71360400

C -3.33826900 3.67087900 -1.45007500

C -2.93718800 1.67823500 -0.11874800

C -1.29229400 2.39258900 -1.71215700

C -2.10400800 3.47523700 -2.07868300

C -3.75171000 2.76341500 -0.46731700

H -3.30482800 0.96789700 0.62514800

H -0.32764400 2.26819500 -2.21592700

H -1.76874500 4.16777600 -2.85720600

H -4.72066300 2.89393800 0.02480100

N 1.12230200 1.19482600 0.05908800

C 3.44215700 2.57769300 0.86492300

C 1.07213100 2.49079000 0.40411800

C 2.31387300 0.55519600 0.12799900

C 3.48031900 1.22225200 0.51855700

C 2.19097300 3.20998100 0.80897500

N 1.02820200 -1.38687300 -0.39265100

C 3.26845500 -3.08892800 -0.43383200

C 0.88870200 -2.70292700 -0.60348100

C 2.26766600 -0.89109600 -0.17997300

C 3.39905200 -1.71348400 -0.20709000

C 1.96801900 -3.58059600 -0.62304700

H 0.09083200 2.95996400 0.35015300

H	4.42582100	0.68308600	0.57120700
H	2.08068200	4.26147400	1.08174800
H	-0.13439700	-3.04380200	-0.77330200
H	4.38996300	-1.28850400	-0.04812700
H	1.79076100	-4.64487400	-0.79103100
Br	-2.06051700	-1.42111000	-1.56489200
C	-0.76489300	-0.35275700	2.17087200
H	-3.97441700	4.51618300	-1.72696000
C	4.46631600	-3.99507300	-0.49511200
H	5.33972900	-3.55120300	0.00358500
H	4.25037000	-4.97063000	-0.03427000
H	4.74108900	-4.18581400	-1.54668100
C	4.68127100	3.32656200	1.26822300
H	4.47905600	3.99892200	2.11576100
H	5.49923900	2.64542300	1.54252400
H	5.03122800	3.95568200	0.43210800
C	0.50842700	-0.98541500	2.66028000
H	0.42913300	-1.17567500	3.75235700
H	0.71453000	-1.95213600	2.17918200
H	1.38256900	-0.32989800	2.52195900
C	-1.93737800	-1.28175200	2.04413200
H	-2.79048300	-0.81209200	1.53520400
H	-1.67886300	-2.18943100	1.47926000
H	-2.28243400	-1.59901300	3.05304200
C	-1.03843600	1.00820600	2.74490500
H	-0.15733500	1.66519500	2.67936800
H	-1.88232000	1.51224100	2.25590000
H	-1.28787500	0.91589500	3.82446800

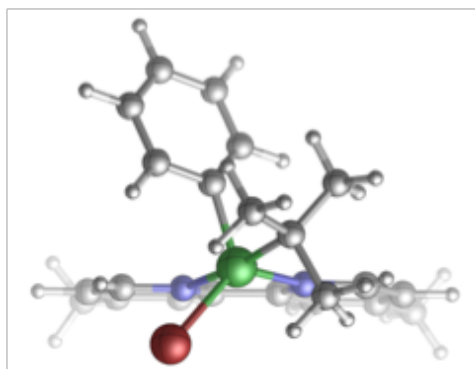
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.5308871

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5693344

<sup>2</sup>H'



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.427088 (Hartree/Particle)

Thermal correction to Energy= 0.454746

Thermal correction to Enthalpy= 0.455691

Thermal correction to Gibbs Free Energy= 0.366759

Sum of electronic and zero-point Energies= -5044.582270

Sum of electronic and thermal Energies= -5044.554611

Sum of electronic and thermal Enthalpies= -5044.553667

Sum of electronic and thermal Free Energies= -5044.642599

C 2.72335600 0.70789700 0.18970800

C 3.75298800 -0.41261300 0.27880000

C 2.97297200 1.55139000 -1.06760500

H 3.70918900 -1.07763500 -0.59404700

H 3.61723800 -1.01751200 1.18615500

H 3.11295700 0.91777900 -1.95212800

H 2.16471200 2.26014100 -1.30123400

C -1.70961900 1.57648700 -0.04686300

C -2.65033500 2.60378800 0.11017800

C -2.24023100 3.93516500 0.23146300

C -0.85950600 4.17639400 0.18311300

C 0.01525300 3.10655200 0.02942200

C -2.10728300 0.15486800 -0.20310600

C -3.43402400 -0.28925800 -0.10788500

C -3.73664600 -1.64641600 -0.26159800

C -2.66328500 -2.51620700 -0.51343200

C -1.37323800 -2.00530300 -0.59468700

H -3.71474700 2.37279300 0.12957800

H -0.46145000 5.18998600 0.26278000

H 1.08852400 3.28660900 -0.00950700

H -4.23973900 0.41575000 0.09379700

H -2.82910800 -3.58791500 -0.64141200

H -0.51253300 -2.64608700 -0.79182700

N -0.38202400 1.83068500 -0.07700800

N -1.10205500 -0.70319200 -0.44673700

Br	1.50064300	-1.34350900	-2.30830600
Ni	0.86324300	0.01931200	-0.25483600
C	0.82842000	-0.73700000	1.51091600
C	0.23013700	-0.08929800	2.59837300
C	1.21218600	-2.07747200	1.64691200
C	0.02519700	-0.76720800	3.80788300
H	-0.07727800	0.95631400	2.50930000
C	1.00946900	-2.75990800	2.85462700
H	1.67143400	-2.59586100	0.79964700
H	-0.44213800	-0.24725900	4.64978200
H	1.31577900	-3.80640300	2.94602500
C	0.41579000	-2.10526800	3.94016600
H	0.25773600	-2.63455900	4.88374000
C	-3.22289900	5.06030200	0.39531800
H	-3.13734000	5.77041300	-0.44331400
H	-4.25907000	4.69732800	0.43832600
H	-3.01158100	5.62624400	1.31685800
C	-5.14604400	-2.16100000	-0.16329600
H	-5.45402400	-2.62397900	-1.11504700
H	-5.22023800	-2.94123900	0.61121900
H	-5.85899800	-1.36057200	0.07871800
H	4.76852300	0.02874600	0.32310300
H	3.89106400	2.15487300	-0.92408400
C	2.77608100	1.55777200	1.45779800
H	1.92937700	2.24961100	1.55717100
H	2.78584400	0.93211000	2.36012000
H	3.70125000	2.16594700	1.45988300

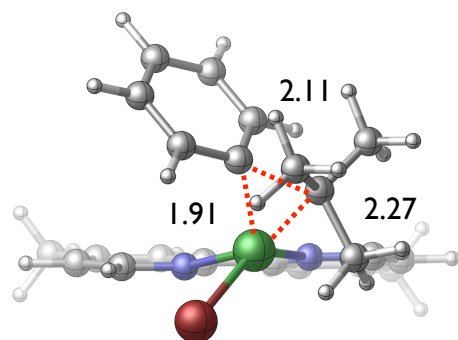
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.5387051

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5811753

<sup>2</sup>I'-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.426807 (Hartree/Particle)

Thermal correction to Energy= 0.453769

Thermal correction to Enthalpy= 0.454713

Thermal correction to Gibbs Free Energy= 0.366959

Sum of electronic and zero-point Energies= -5044.567409

Sum of electronic and thermal Energies= -5044.540447

Sum of electronic and thermal Enthalpies= -5044.539502

Sum of electronic and thermal Free Energies= -5044.627257

C 2.47756700 0.52963500 0.50619500

C 3.59074200 -0.50541300 0.62937500

C 2.69296100 1.27633000 -0.83597600

H 3.50650100 -1.27055700 -0.15537000

H 3.59625100 -0.99952600 1.61104300

H 2.89116500 0.58777800 -1.66519700

H 1.87756200 1.94478300 -1.14710300

C -2.00954800 1.50230600 -0.13993600

C -2.94484600 2.52885500 0.05036000

C -2.53057300 3.85937000 0.16696800

C -1.15101300 4.10093700 0.08094300

C -0.28187800 3.03329200 -0.11364300

C -2.40652900 0.07676900 -0.26387500

C -3.70782000 -0.38689300 -0.02447700

C -4.00335400 -1.74934200 -0.14650300

C -2.94968800 -2.60226100 -0.51360000

C -1.68261200 -2.07229100 -0.73146100

H -4.00836400 2.29638700 0.09838000

H -0.75091100 5.11370200 0.16189500

H 0.79136600 3.21327800 -0.18329100

H -4.49584300 0.30491700 0.27245700

H -3.11289100 -3.67649700 -0.62411400

H -0.83301300 -2.69688500 -1.01558600

N -0.68378500 1.75827400 -0.22123600

N -1.41708500 -0.76567600 -0.61231500

Br	1.40168200	-1.60311700	-2.29320600
Ni	0.51705700	-0.02284900	-0.48947000
C	0.89376300	-0.62924200	1.28296600
C	0.15210500	-0.03739800	2.32491800
C	1.14006200	-2.01494600	1.35357400
C	-0.36135300	-0.80750500	3.37237100
H	-0.05846300	1.03243100	2.30102800
C	0.62738700	-2.78433700	2.40279400
H	1.69961200	-2.49969600	0.55288600
H	-0.95334000	-0.32388900	4.15510200
H	0.81577700	-3.86192800	2.42004200
C	-0.12792600	-2.18770600	3.41948900
H	-0.52755500	-2.78946600	4.23983300
C	-3.50767800	4.98413300	0.36577100
H	-3.45621400	5.69224200	-0.47742700
H	-4.54096600	4.61980500	0.44926900
H	-3.26263800	5.55310700	1.27705300
C	-5.38404000	-2.28838400	0.10891100
H	-5.77387800	-2.79393300	-0.78952600
H	-5.36298100	-3.03958400	0.91507700
H	-6.08872000	-1.49437900	0.39309300
H	4.56567300	0.00058500	0.50057000
H	3.58393500	1.92378900	-0.71775000
C	2.55206500	1.54057900	1.64884500
H	1.72780400	2.26708000	1.62507400
H	2.54372100	1.05494100	2.63426500
H	3.49408500	2.11235900	1.56275600

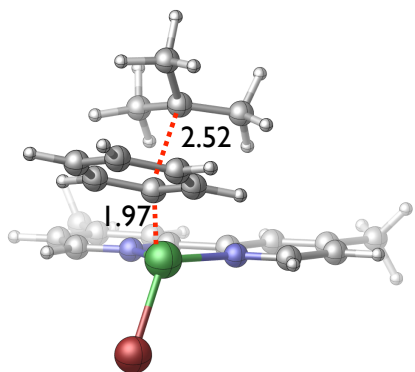
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.5303702

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5662454

<sup>2</sup>I-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.422563 (Hartree/Particle)

Thermal correction to Energy= 0.451102

Thermal correction to Enthalpy= 0.452046

Thermal correction to Gibbs Free Energy= 0.358259

Sum of electronic and zero-point Energies= -5044.535437

Sum of electronic and thermal Energies= -5044.506898

Sum of electronic and thermal Enthalpies= -5044.505954

Sum of electronic and thermal Free Energies= -5044.599741

Ni 1.15698000 0.80313900 -0.58477500

C 2.32848100 0.33814200 0.92428500

C 5.11920600 0.20086500 1.49479100

C 3.04155200 1.45126300 1.43037300

C 3.07877700 -0.84127500 0.69778300

C 4.44572700 -0.91665100 0.98020600

C 4.40990400 1.38942900 1.71551000

H 2.50862100 2.39036600 1.62008700

H 2.57443100 -1.73002300 0.30029100

H 4.99329200 -1.84687500 0.79755300

H 4.92832600 2.26924700 2.11004400

N -0.40671000 1.93537300 0.04765400

C -2.69659900 3.26224200 0.96611900

C -0.33012700 3.19445800 0.49328600

C -1.60084200 1.30443100 0.03778200

C -2.75851900 1.94552700 0.49241800

C -1.43981300 3.89018500 0.96037900

N -0.34382800 -0.53554700 -0.82382800

C -2.55499400 -2.24308600 -1.00356100

C -0.20010200 -1.78530500 -1.27722900

C -1.56646500 -0.09342900 -0.46128400

C -2.68807300 -0.92631400 -0.54461700

C -1.26934200 -2.66854900 -1.37809500

H 0.66285300 3.65020900 0.47535400

H	-3.71494700	1.42282200	0.48486800
H	-1.32455100	4.91485400	1.31930300
H	0.81230000	-2.07394900	-1.57021900
H	-3.67063500	-0.55699600	-0.25052300
H	-1.10182100	-3.68134400	-1.74991400
Br	2.04109800	1.26858800	-2.81423400
C	0.90892800	-0.32085900	2.89960800
C	2.06565600	-0.62608100	3.80419000
C	0.14137600	0.92248900	3.22797600
H	2.70749700	0.25543600	3.95657700
H	2.69217600	-1.43948000	3.40736200
H	0.79502900	1.81017100	3.24348300
H	-0.67731400	1.10772600	2.52043400
H	6.18789400	0.14581000	1.71989400
C	-3.73205600	-3.17533400	-1.08137800
H	-3.75552200	-3.69814800	-2.05004800
H	-3.65572600	-3.94915500	-0.29912400
H	-4.68451900	-2.64474300	-0.94359900
C	-3.92289200	3.98596800	1.44833600
H	-4.78181200	3.30884900	1.55526400
H	-3.73480200	4.47411400	2.41718100
H	-4.20010800	4.78067500	0.73558800
C	0.14094900	-1.50531600	2.39991100
H	-0.66437500	-1.21879700	1.71091100
H	0.79774500	-2.22523500	1.88452900
H	-0.33165900	-2.05264400	3.24444900
H	-0.31389900	0.84612700	4.23935000
H	1.69877500	-0.94589100	4.80346700

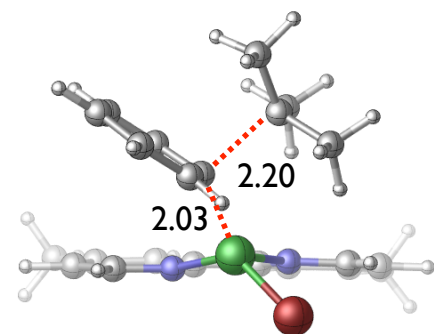
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.4957062

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5367812

<sup>4</sup>I-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.423090 (Hartree/Particle)

Thermal correction to Energy= 0.450907

Thermal correction to Enthalpy= 0.451851

Thermal correction to Gibbs Free Energy= 0.361396

Sum of electronic and zero-point Energies= -5044.542031

Sum of electronic and thermal Energies= -5044.514215

Sum of electronic and thermal Enthalpies= -5044.513270

Sum of electronic and thermal Free Energies= -5044.603726

Ni 1.00135600 1.42580000 0.04129900

C 2.02000000 0.07814700 1.17453600

C 3.93315500 -2.07728400 1.15319100

C 3.43288500 0.29828400 1.02085000

C 1.63092500 -1.30446900 1.21498900

C 2.55286200 -2.34444500 1.23634800

C 4.35898900 -0.74276500 1.03278300

H 3.79393400 1.32489200 0.90185400

H 0.56512100 -1.55255200 1.24828500

H 2.20252300 -3.38075100 1.29090300

H 5.42643100 -0.52063300 0.92890300

N -0.89401700 2.03940500 0.51380700

C -3.56934600 2.63784800 1.09908700

C -1.21167500 3.15802500 1.18217100

C -1.87998700 1.21421900 0.09068800

C -3.22327800 1.48924400 0.38009700

C -2.52195100 3.49173000 1.49550500

N -0.09733200 0.04534000 -0.98052000

C -1.72321000 -2.06579300 -1.84944700

C 0.42587200 -0.97644000 -1.67241200

C -1.41869300 0.04577700 -0.68977000

C -2.24981100 -0.99990400 -1.11028500

C -0.34489300 -2.03680300 -2.13142800

H -0.37158200 3.79948600 1.45667900

H -4.00678500 0.81553400 0.03241500

H	-2.72889600	4.41386500	2.04218700
H	1.50289900	-0.93420300	-1.84772000
H	-3.31006400	-0.99346900	-0.85626500
H	0.12601800	-2.84155600	-2.69960600
Br	2.10369000	3.55939100	-0.38470900
C	1.60926300	0.73058300	3.23513900
C	2.54956500	-0.10886100	4.05616600
C	1.93791500	2.19806400	3.21829800
H	3.60092400	0.08623000	3.79444600
H	2.35480700	-1.18411500	3.92224600
H	2.99142400	2.37533000	2.95440400
H	1.32098100	2.75065800	2.49722400
H	4.65885800	-2.89498300	1.15903700
C	-2.58235200	-3.20804900	-2.31633900
H	-3.63844800	-3.06244600	-2.04961600
H	-2.51152600	-3.32503800	-3.40984200
H	-2.23976200	-4.15626900	-1.87026900
C	-4.99807700	2.96615500	1.43148300
H	-5.69656700	2.22517400	1.01846500
H	-5.13933800	3.00560100	2.52409000
H	-5.26879600	3.95941000	1.03818500
C	0.15182100	0.40560400	3.42019600
H	-0.46414900	0.87526500	2.64206100
H	-0.02996500	-0.67959200	3.39789400
H	-0.20509700	0.78000400	4.40156300
H	1.76418400	2.64050000	4.22078000
H	2.42471500	0.12233500	5.13414200

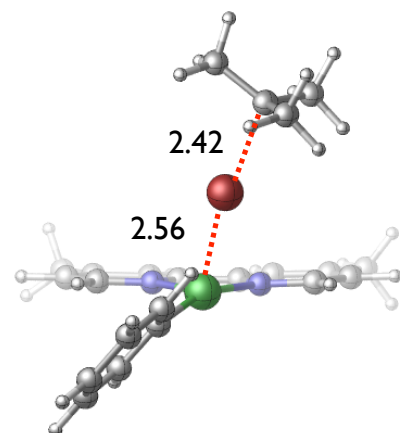
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.4932847

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5389644

<sup>2</sup>L-TS



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.422923 (Hartree/Particle)

Thermal correction to Energy= 0.451544

Thermal correction to Enthalpy= 0.452488

Thermal correction to Gibbs Free Energy= 0.356259

Sum of electronic and zero-point Energies= -5044.541704

Sum of electronic and thermal Energies= -5044.513083

Sum of electronic and thermal Enthalpies= -5044.512138

Sum of electronic and thermal Free Energies= -5044.608368

Ni -0.42182000 0.28394900 0.52338800

C -1.80483100 1.59839400 0.08222800

C -3.75127500 3.55126000 -0.71046800

C -3.11085500 1.62512200 0.62313500

C -1.52534800 2.60154000 -0.87716700

C -2.46877300 3.56119400 -1.27153000

C -4.06879200 2.57418800 0.24189900

H -3.38188200 0.88078600 1.38049600

H -0.52897400 2.64102100 -1.33505300

H -2.20361700 4.31984100 -2.01585000

H -5.06786900 2.55682500 0.69094200

N 1.38089900 1.24846600 0.65316400

C 3.94018900 2.39158400 0.67012700

C 1.53603800 2.56455300 0.83111200

C 2.47049700 0.46895700 0.48739400

C 3.75881600 1.01414900 0.48322500

C 2.78796100 3.17384400 0.85031300

N 0.86296800 -1.29176100 0.35058000

C 2.78615500 -3.31968700 0.11769000

C 0.49414600 -2.57644400 0.27451100

C 2.17591600 -0.98024300 0.33754200

C 3.15617800 -1.97280700 0.21740800

C 1.41223400 -3.61388500 0.15535900

H	0.61307200	3.13688900	0.95197000
H	4.62822500	0.37261300	0.33671800
H	2.86288300	4.25292500	1.00057900
H	-0.58091200	-2.76685800	0.32347700
H	4.21293800	-1.70438900	0.20897700
H	1.05973000	-4.64567000	0.09425500
C	-1.70339200	-1.51724700	4.95922900
H	-4.49368300	4.29747100	-1.00910500
C	3.80773700	-4.41465600	-0.01957800
H	4.83431000	-4.02383000	0.01188900
H	3.69193600	-5.15735300	0.78619000
H	3.67032500	-4.95176500	-0.97237900
C	5.31233900	3.00659900	0.70318300
H	5.61477600	3.20019900	1.74652000
H	6.06663800	2.34633500	0.25205200
H	5.32697200	3.97333400	0.17749400
C	-3.17908900	-1.74559500	4.77101000
H	-3.72436000	-0.79764000	4.64426200
H	-3.38258300	-2.38384100	3.89725400
H	-3.59803200	-2.25610000	5.66316200
C	-0.85347500	-2.75976100	4.94519400
H	0.22032000	-2.51700700	4.94208600
H	-1.05765300	-3.36614900	5.85213700
H	-1.07002500	-3.38925900	4.06831700
C	-1.33700500	-0.49842700	6.00492700
H	-1.58017900	-0.89104400	7.01440800
H	-0.26032500	-0.26896400	5.98909400
H	-1.89570700	0.44068200	5.87117300
Br	-1.07018600	-0.42224200	2.89432600

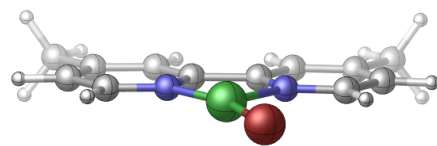
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5045.5065849

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-5046.5454233

<sup>2</sup>P<sub>B</sub>



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.214708 (Hartree/Particle)

Thermal correction to Energy= 0.230876

Thermal correction to Enthalpy= 0.231820

Thermal correction to Gibbs Free Energy= 0.166322

Sum of electronic and zero-point Energies= -4655.579462

Sum of electronic and thermal Energies= -4655.563294

Sum of electronic and thermal Enthalpies= -4655.562349

Sum of electronic and thermal Free Energies= -4655.627848

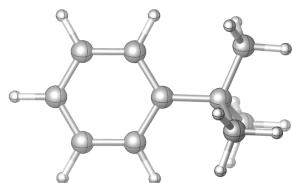
Ni	-3.99183200	1.30009300	-1.54653000
C	-1.87359800	-0.39708200	-0.83451100
C	-0.60463100	-0.81497600	-0.41773600
C	0.38038200	0.12429500	-0.09164500
C	0.02971800	1.48282700	-0.20377800
C	-1.24637600	1.83242200	-0.62458000
C	-2.97912600	-1.31121800	-1.19958100
C	-2.90514100	-2.70937500	-1.17420400
C	-4.01531200	-3.47922700	-1.54213400
C	-5.17917400	-2.78948900	-1.92940000
C	-5.18373000	-1.39947400	-1.93172800
H	-0.37768600	-1.87943400	-0.34653600
H	0.75171100	2.26661600	0.03582700
H	-1.54668200	2.87837300	-0.72245500
H	-1.98301600	-3.20525100	-0.86858000
H	-6.07708900	-3.33486700	-2.22820500
H	-6.06484300	-0.82279400	-2.22618200
N	-2.18757800	0.92199600	-0.93709400
N	-4.11313300	-0.67488700	-1.57617200
C	1.75486100	-0.28579000	0.36089600
H	1.95781900	0.09097300	1.37696900
H	1.87445700	-1.37835300	0.36807700
H	2.52660300	0.14347000	-0.29878000
C	-3.97959500	-4.98297800	-1.52479800
H	-2.98906900	-5.36658100	-1.24261000
H	-4.71990400	-5.37924000	-0.81035300
H	-4.24149600	-5.38984900	-2.51492300
Br	-5.83964900	2.56755300	-2.20535500

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4656.2756634

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF=-4656.9610627

**tBu-Ph**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.212552 (Hartree/Particle)

Thermal correction to Energy= 0.222172

Thermal correction to Enthalpy= 0.223116

Thermal correction to Gibbs Free Energy= 0.178247

Sum of electronic and zero-point Energies= -389.034504

Sum of electronic and thermal Energies= -389.024885

Sum of electronic and thermal Enthalpies= -389.023940

Sum of electronic and thermal Free Energies= -389.068810

C 0.04421000 1.70961600 1.14572500

C 1.44490300 1.73358100 1.12055900

C 2.12656700 2.88889000 0.73341700

C 1.38993500 4.02254300 0.37071900

C -0.00591100 3.99264800 0.39798600

C -0.71318100 2.83690300 0.78576000

H -0.45028400 0.78804600 1.45338500

H 2.00284900 0.83794100 1.40728300

H 3.21924500 2.90824800 0.71359200

H 1.90528600 4.93693000 0.06433300

H -0.55299400 4.89314100 0.10924100

C -2.25333700 2.84939200 0.79950700

C -2.74935100 3.94948800 1.76551900

H -2.40180400 4.94892000 1.46299200

H -3.85134000 3.97127700 1.78683700

H -2.39094800 3.76208800 2.79042100

C -2.84401600 1.50472700 1.25836000

H -3.94379100 1.56138300 1.25324000

H -2.55073000 0.67977000 0.59043900

H -2.52933600 1.24749100 2.28192300

C -2.77455500 3.14452300 -0.62558100

H -3.87683800 3.15946900 -0.63495500

H -2.42445300 4.11964200 -0.99703000

H -2.43723000 2.37132500 -1.33443100

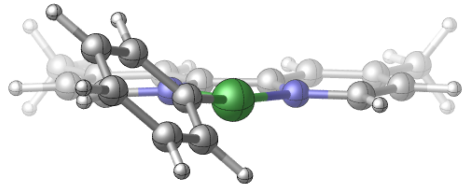
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-389.3290308

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-389.6748001

<sup>2</sup>P<sub>C</sub>



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.303134 (Hartree/Particle)

Thermal correction to Energy= 0.322860

Thermal correction to Enthalpy= 0.323805

Thermal correction to Gibbs Free Energy= 0.250611

Sum of electronic and zero-point Energies= -2313.026966

Sum of electronic and thermal Energies= -2313.007240

Sum of electronic and thermal Enthalpies= -2313.006296

Sum of electronic and thermal Free Energies= -2313.079489

Ni	-3.96398300	1.44675100	-1.21509900
C	-1.83575500	-0.36900700	-0.73179300
C	-0.55130400	-0.83036800	-0.42006200
C	0.43164200	0.05760500	0.03112800
C	0.06264200	1.41025300	0.15257000
C	-1.22735300	1.80375600	-0.17808400
C	-2.94328000	-1.23262600	-1.20069300
C	-2.85927000	-2.62301000	-1.34872800
C	-3.97607300	-3.35335800	-1.77274900
C	-5.15577000	-2.63260900	-2.03077500
C	-5.16546600	-1.25120500	-1.86635600
H	-0.31078600	-1.88864300	-0.52848300
H	0.78098300	2.15531600	0.50175800
H	-1.53996000	2.84781500	-0.09948800
H	-1.92710500	-3.14475700	-1.12852900
H	-6.06285900	-3.14566800	-2.35800600
H	-6.06078500	-0.65514500	-2.05962300
N	-2.16615900	0.94343400	-0.61445400
N	-4.08728100	-0.56287300	-1.46861500
C	1.82261900	-0.39985100	0.37491200
H	2.05670900	-0.17550000	1.42863300
H	1.94941500	-1.48000600	0.21584000
H	2.56967500	0.13060300	-0.23804800
C	-3.92851700	-4.84679700	-1.94782400
H	-2.95001600	-5.26161200	-1.66768600
H	-4.70238900	-5.33564400	-1.33408100
H	-4.13207500	-5.11973300	-2.99640100
C	-5.53228400	2.36315800	-1.90331200
C	-6.16411500	3.46923000	-1.27984400

C -6.13057200 1.95132100 -3.12227500  
C -7.29512300 4.10503900 -1.81103500  
H -5.75841000 3.85373800 -0.33612700  
C -7.25947700 2.57286700 -3.67293800  
H -5.69267000 1.10759400 -3.66945300  
C -7.85243900 3.65805100 -3.01549200  
H -7.74560100 4.95361600 -1.28450500  
H -7.67970900 2.21192800 -4.61812500  
H -8.73512500 4.14906300 -3.43623300

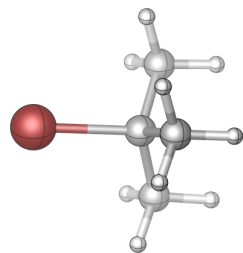
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HF=-2313.662927

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2314.4257224

**tBu-Br**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.121249 (Hartree/Particle)

Thermal correction to Energy= 0.127905

Thermal correction to Enthalpy= 0.128849

Thermal correction to Gibbs Free Energy= 0.090653

Sum of electronic and zero-point Energies= -2731.509462

Sum of electronic and thermal Energies= -2731.502806

Sum of electronic and thermal Enthalpies= -2731.501862

Sum of electronic and thermal Free Energies= -2731.540058

Br 0.66845300 -3.32582400 -0.02012300

C -0.35404100 -1.55353000 -0.02192100

C 0.09890600 -0.79012700 -1.26110700

H -0.13485200 -1.34337700 -2.18230000

H -0.43396200 0.17601100 -1.29443400

H 1.17917200 -0.58596100 -1.23527900

C -1.83589000 -1.90706100 -0.07308600

H -2.13789300 -2.49998800 0.80239100

H -2.42204700 -0.97172100 -0.07532100

H -2.08349000 -2.47031100 -0.98450400

C 0.02110100 -0.83233800 1.26757900

H -0.51694000 0.13087300 1.30189300

H -0.26517900 -1.41745400 2.15359400

H 1.10013000 -0.62515900 1.31345500

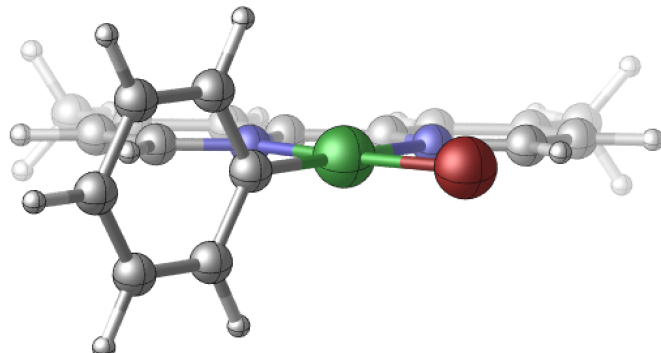
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2731.8563155

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-2732.1242505

**Figure S9 (with Becke-Johnson damping)**



UB3LYP-D3BJ/def2-SVP-CPCM(THF)

Zero-point correction= 0.306461 (Hartree/Particle)

Thermal correction to Energy= 0.326798

Thermal correction to Enthalpy= 0.327742

Thermal correction to Gibbs Free Energy= 0.253612

Sum of electronic and zero-point Energies= -4887.034664

Sum of electronic and thermal Energies= -4887.014327

Sum of electronic and thermal Enthalpies= -4887.013383

Sum of electronic and thermal Free Energies= -4887.087513

Ni -4.61455800 0.39141800 -0.38378100

C -6.47995100 0.32752600 -0.64642700

C -7.06929100 0.72563800 -1.85950000

C -7.32573900 -0.19722300 0.34733000

C -8.44464000 0.58283800 -2.08288600

H -6.44788200 1.15381400 -2.65241400

C -8.70232500 -0.34109800 0.13197000

H -6.90771400 -0.51094000 1.30937600

C -9.26861600 0.04583400 -1.08749400

H -8.87587500 0.89543900 -3.03877400

H -9.33600100 -0.75704500 0.92114600

H -10.34235900 -0.06639400 -1.25931300

Br -4.83954100 2.61489600 0.34890300

C -2.08006400 -0.92116800 -0.45758300

C -0.71658600 -1.19386200 -0.31567800

C 0.14713300 -0.21566200 0.18983700

C -0.42098800 1.02026200 0.53406200

C -1.78653700 1.22356800 0.36804800

C -3.06510700 -1.88765800 -0.98117200

C -2.72565700 -3.17109000 -1.41575200

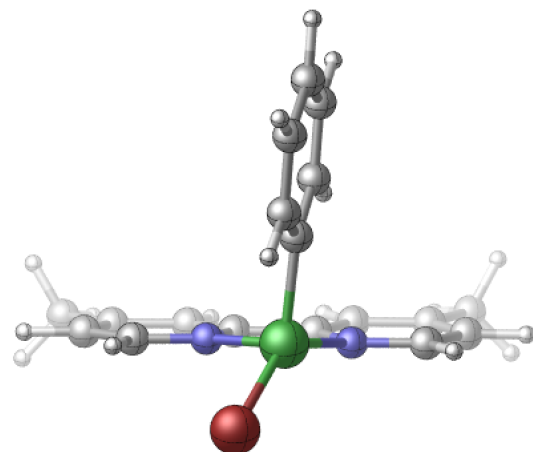
C -3.70626600 -4.03929000 -1.90596700

C -5.02125000 -3.55308100 -1.93323700

C -5.29618900 -2.26681600 -1.48742800

H -0.31865600 -2.16867700 -0.59552600

H	0.19699500	1.82707600	0.93279300
H	-2.26735000	2.17027400	0.62492300
H	-1.68801300	-3.50015800	-1.37533500
H	-5.84024600	-4.17312400	-2.30241600
H	-6.31123600	-1.87427200	-1.50210400
N	-2.60198000	0.27710900	-0.11591900
N	-4.34663200	-1.43785100	-1.01661200
C	1.61663500	-0.47344500	0.36269600
H	1.89776400	-0.40195800	1.42590300
H	1.90088100	-1.46896700	-0.00428500
H	2.20936800	0.28185300	-0.17712700
C	-3.37343200	-5.42157500	-2.38792100
H	-3.59867400	-5.51805300	-3.46249700
H	-2.31286900	-5.66220400	-2.23393200
H	-3.98429000	-6.17334000	-1.86370600



UB3LYP-D3BJ/def2-SVP-CPCM(THF)

Zero-point correction= 0.305082 (Hartree/Particle)

Thermal correction to Energy= 0.327019

Thermal correction to Enthalpy= 0.327963

Thermal correction to Gibbs Free Energy= 0.246939

Sum of electronic and zero-point Energies= -4887.032375

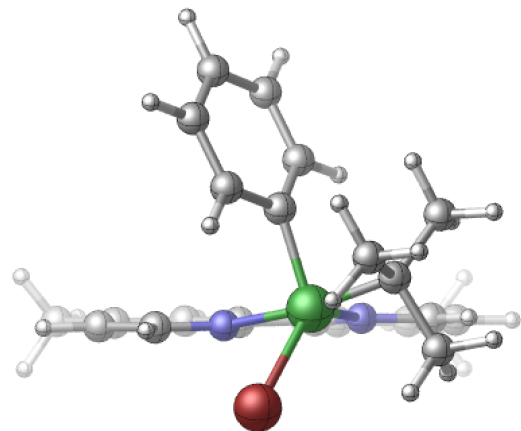
Sum of electronic and thermal Energies= -4887.010439

Sum of electronic and thermal Enthalpies= -4887.009494

Sum of electronic and thermal Free Energies= -4887.090519

C	-1.67439600	0.90308300	-0.64637300
C	-1.84605600	0.00921700	0.42800600
C	-3.07766900	-0.67227900	0.49143500
C	-4.08768700	-0.47102500	-0.45923300
C	-3.88787800	0.42453000	-1.51520600
C	-2.67264600	1.11236300	-1.60688000
Br	-0.90109600	-2.16723900	3.30580200
H	-3.25387400	-1.38515300	1.30419200
H	-0.73615300	1.46023800	-0.75016000
H	-2.50304800	1.81374400	-2.43003000
H	-4.67130500	0.58428500	-2.26125500
H	-5.03298000	-1.01689000	-0.37771600
C	1.99355600	0.06542900	0.28566900
C	1.37074100	-2.17251900	0.20732800
C	2.44074900	-2.48947000	-0.62142100
C	3.33200200	-1.47888000	-1.01880900
C	3.08937000	-0.18348100	-0.54637800
C	1.65209400	1.39945800	0.83712100
C	0.17967400	2.57976100	2.18952000
C	0.83585100	3.77934100	1.94024000
C	1.95525000	3.79245000	1.09192400
C	2.35588900	2.57050500	0.53879900
H	0.65892000	-2.92818700	0.54830100

H	2.58098600	-3.51968000	-0.95371900
H	3.76239000	0.62605000	-0.82761700
H	-0.70242300	2.53971100	2.83335800
H	0.47388300	4.69967200	2.40212200
H	3.21583000	2.53874300	-0.12966200
N	1.15583200	-0.92758900	0.64512400
N	0.58069100	1.41995500	1.65742100
Ni	-0.43284100	-0.35400600	1.76916300
C	2.67656200	5.07255000	0.78089400
H	3.61042600	4.89040300	0.23203700
H	2.03937000	5.72984800	0.16640500
H	2.91062900	5.62385500	1.70458600
C	4.49472400	-1.78212000	-1.92015900
H	5.15539700	-0.91255100	-2.03768400
H	5.08551300	-2.62309900	-1.52506400
H	4.13702000	-2.08132800	-2.91901300



UB3LYP-D3BJ/def2-SVP-CPCM(THF)

Zero-point correction= 0.426844 (Hartree/Particle)

Thermal correction to Energy= 0.454721

Thermal correction to Enthalpy= 0.455665

Thermal correction to Gibbs Free Energy= 0.365748

Sum of electronic and zero-point Energies= -5044.644159

Sum of electronic and thermal Energies= -5044.616282

Sum of electronic and thermal Enthalpies= -5044.615338

Sum of electronic and thermal Free Energies= -5044.705255

C 2.71091400 0.70133000 0.17596900

C 3.72349900 -0.43154200 0.28402500

C 2.98998400 1.53526600 -1.08049000

H 3.67687400 -1.10725700 -0.58063800

H 3.57774000 -1.02143400 1.19949200

H 3.15682700 0.89559100 -1.95584100

H 2.18550900 2.23884300 -1.34152700

C -1.69402700 1.55618600 -0.04726500

C -2.63173800 2.57677800 0.15141200

C -2.22064200 3.90684800 0.27641200

C -0.84314500 4.15306100 0.18713200

C 0.03023000 3.08910500 -0.00498100

C -2.09021700 0.13838900 -0.21383300

C -3.41303500 -0.30945800 -0.10707200

C -3.71271400 -1.66483200 -0.27611500

C -2.64112400 -2.52694500 -0.55775500

C -1.35382900 -2.01254400 -0.65025000

H -3.69370500 2.34105200 0.20160700

H -0.44643100 5.16687700 0.26539300

H 1.10083200 3.27379300 -0.07558200

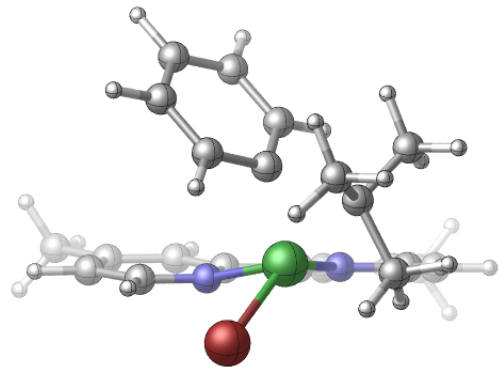
H -4.21723300 0.39093900 0.11374100

H -2.80673000 -3.59632400 -0.70153300

H -0.49597000 -2.64747600 -0.87629200

N -0.36945900 1.81503100 -0.11266300

N	-1.08673700	-0.71263200	-0.48381700
Br	1.50821600	-1.31897800	-2.33001600
Ni	0.85259800	0.02307700	-0.28079500
C	0.79730100	-0.69864000	1.49793900
C	0.19519200	-0.03263100	2.57163300
C	1.17579900	-2.03728600	1.66018800
C	-0.01918000	-0.68997200	3.79003700
H	-0.10878200	1.01265000	2.47201700
C	0.96369600	-2.69875500	2.87698900
H	1.63990400	-2.57691200	0.82945200
H	-0.48937000	-0.15415600	4.61983500
H	1.26665300	-3.74406700	2.98682800
C	0.36551400	-2.02585000	3.94750800
H	0.19984000	-2.53889000	4.89826300
C	-3.20164200	5.02385800	0.48646100
H	-3.13626200	5.75429200	-0.33587700
H	-4.23415500	4.65350500	0.54162900
H	-2.97452000	5.56770700	1.41724000
C	-5.11835800	-2.18248600	-0.16317700
H	-5.43728700	-2.64475300	-1.11124400
H	-5.18275500	-2.96280500	0.61179300
H	-5.82734500	-1.38202800	0.08820300
H	4.74437300	-0.00292000	0.32699200
H	3.90056500	2.14439200	-0.91590400
C	2.76897100	1.55791900	1.43739900
H	1.95083600	2.28612000	1.51619100
H	2.74202400	0.94099800	2.34489300
H	3.71643100	2.13055100	1.44891200



UB3LYP-D3BJ/def2-SVP-CPCM(THF)

Zero-point correction= 0.426726 (Hartree/Particle)

Thermal correction to Energy= 0.453765

Thermal correction to Enthalpy= 0.454709

Thermal correction to Gibbs Free Energy= 0.366635

Sum of electronic and zero-point Energies= -5044.630260

Sum of electronic and thermal Energies= -5044.603220

Sum of electronic and thermal Enthalpies= -5044.602276

Sum of electronic and thermal Free Energies= -5044.690350

C 2.44567300 0.53879700 0.51789700

C 3.55988700 -0.48320800 0.69515100

C 2.69124800 1.26143600 -0.82820000

H 3.51406200 -1.25983000 -0.08110200

H 3.53199200 -0.96353900 1.68286000

H 2.97872300 0.55874900 -1.61770900

H 1.84801100 1.85518400 -1.21171100

C -1.97392900 1.50906500 -0.14440500

C -2.90744400 2.53108100 0.06641500

C -2.49476400 3.86306000 0.16272800

C -1.11973300 4.10986300 0.03412600

C -0.25100100 3.04575600 -0.17615500

C -2.36794500 0.08522200 -0.26140900

C -3.65266100 -0.38646800 0.03509500

C -3.94659600 -1.74822600 -0.09126900

C -2.90999100 -2.58957500 -0.52593500

C -1.65662000 -2.05212900 -0.79547800

H -3.96773700 2.29311700 0.14331500

H -0.72282900 5.12494000 0.09412100

H 0.81856500 3.23114600 -0.27955900

H -4.42648800 0.29800700 0.38125300

H -3.07541100 -3.66202900 -0.64685900

H -0.81926500 -2.66854200 -1.13014400

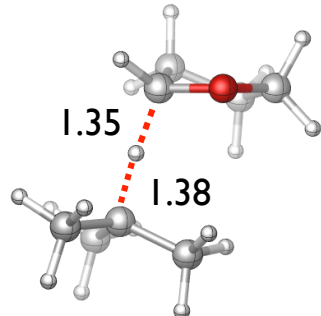
N -0.65260300 1.76954400 -0.26060600

N -1.39242000 -0.74687800 -0.66659600

Br 1.43767500 -1.57408300 -2.31924700

Ni	0.52937800	0.00131100	-0.52962500
C	0.85280700	-0.63865600	1.24830300
C	0.09354700	-0.06193700	2.28413400
C	1.10655300	-2.02232200	1.30784000
C	-0.43862000	-0.84744200	3.31019100
H	-0.11698500	1.00807500	2.27819200
C	0.57567900	-2.80689800	2.33566600
H	1.68758200	-2.49660000	0.51628400
H	-1.04554100	-0.37555300	4.08810000
H	0.76942900	-3.88326900	2.34387300
C	-0.20357300	-2.22681000	3.34286400
H	-0.61848600	-2.84046300	4.14613200
C	-3.47141200	4.98238300	0.38274900
H	-3.43766600	5.69516600	-0.45690500
H	-4.50023900	4.61061200	0.48136200
H	-3.21471500	5.54710100	1.29320600
C	-5.30712300	-2.29630900	0.23476800
H	-5.71999000	-2.85009600	-0.62328000
H	-5.24523600	-3.00657000	1.07528700
H	-6.01222800	-1.49930600	0.50773000
H	4.53399100	0.03065800	0.59548900
H	3.52402500	1.97625100	-0.68136900
C	2.46599700	1.56065500	1.64871200
H	1.63202800	2.27372300	1.59353400
H	2.43495200	1.08311300	2.63716600
H	3.40078300	2.14669100	1.58526300

**Figure S11**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.229357 (Hartree/Particle)

Thermal correction to Energy= 0.240955

Thermal correction to Enthalpy= 0.241899

Thermal correction to Gibbs Free Energy= 0.191011

Sum of electronic and zero-point Energies= -389.737578

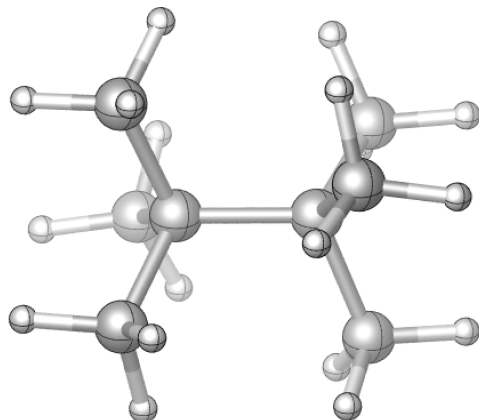
Sum of electronic and thermal Energies= -389.725980

Sum of electronic and thermal Enthalpies= -389.725036

Sum of electronic and thermal Free Energies= -389.775924

C	-1.44189600	-5.62124800	-0.67071100
C	-0.61480900	-5.32477500	-1.90511300
H	0.42581800	-5.67253000	-1.79015200
H	-1.03222900	-5.82984500	-2.79962000
H	-0.58926400	-4.24371200	-2.12344100
C	-2.83311600	-5.02126000	-0.70209400
H	-3.44082800	-5.45479900	-1.52205400
H	-3.37638700	-5.20671800	0.23949800
H	-2.79434600	-3.93056700	-0.85593700
C	-1.39181600	-7.06756400	-0.22146500
H	-0.35337000	-7.42058800	-0.10399400
H	-1.91094600	-7.21344800	0.74069000
H	-1.88170500	-7.73489100	-0.95970900
H	-0.81877200	-4.90191300	0.32518700
C	-0.33452700	-4.09984500	1.30392300
O	-1.24547200	-3.04426000	1.45253600
C	-0.33630300	-4.89587900	2.59470000
H	0.63069200	-3.73694500	0.91889100
C	-2.10155200	-3.29099300	2.57721000
C	-1.79547200	-4.71958900	3.04322000
H	0.35604400	-4.45544600	3.33635100
H	-0.04777600	-5.94793800	2.44839100
H	-1.87162600	-2.55110400	3.36634100
H	-3.15091700	-3.14936800	2.27110800
H	-1.94582300	-4.85268400	4.12428800

H -2.44122600 -5.44128100 2.51794600  
UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -390.4090221  
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -390.0778207



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.243588 (Hartree/Particle)  
 Thermal correction to Energy= 0.253800  
 Thermal correction to Enthalpy= 0.254744  
 Thermal correction to Gibbs Free Energy= 0.210503  
 Sum of electronic and zero-point Energies= -315.255129  
 Sum of electronic and thermal Energies= -315.244917  
 Sum of electronic and thermal Enthalpies= -315.243973  
 Sum of electronic and thermal Free Energies= -315.288214  
 C -2.41708000 -0.14457300 0.06622800  
 C -0.88402500 -0.24743600 -0.08529400  
 H -2.74165800 0.88150100 0.29535000  
 H -2.94544800 -0.47308000 -0.84217800  
 H -2.75610000 -0.78736300 0.89495100  
 C -0.51590700 -1.74621100 -0.13042400  
 C -0.24575200 0.35513700 1.18492900  
 H 0.55546200 -1.90306300 -0.32988500  
 H -0.74310900 -2.21871100 0.83915500  
 H -1.08705800 -2.28817800 -0.89894500  
 H -0.70779400 -0.08687200 2.08277100  
 H 0.83422400 0.15250100 1.24049600  
 H -0.39027400 1.44511100 1.24399800  
 C -0.38354500 0.50698200 -1.39515200  
 C -1.02342000 1.90725400 -1.51033200  
 C 1.14931300 0.69016800 -1.38276200  
 C -0.75168200 -0.28406100 -2.66911300  
 H -2.10296400 1.85183000 -1.71534600  
 H -0.88128800 2.50327700 -0.59565000  
 H -0.56103700 2.46448200 -2.34135800  
 H 1.67895000 -0.26049400 -1.21503400  
 H 1.48707500 1.08812900 -2.35358700  
 H 1.47371500 1.40115000 -0.60831800  
 H -0.52342700 0.31783100 -3.56385600

H -0.18082000 -1.22083700 -2.75183900  
H -1.82331800 -0.53448200 -2.70559900

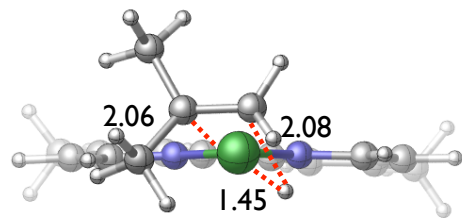
UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -315.8529126

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -315.5614974

**Figure S12**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.328127 (Hartree/Particle)

Thermal correction to Energy= 0.347968

Thermal correction to Enthalpy= 0.348913

Thermal correction to Gibbs Free Energy= 0.279623

Sum of electronic and zero-point Energies= -2239.220375

Sum of electronic and thermal Energies= -2239.200534

Sum of electronic and thermal Enthalpies= -2239.199590

Sum of electronic and thermal Free Energies= -2239.268880

Ni -4.61321300 0.86810700 -0.83470900

C -5.09920600 2.52182300 0.29792100

C -4.55216600 2.36939500 1.70282700

H -4.74404000 1.36415800 2.10736000

H -3.46927900 2.55485400 1.75648600

H -5.03615100 3.10331500 2.37489000

C -6.24127500 1.79571800 -0.07944700

H -5.62378300 0.94464700 -1.87805000

H -6.68945700 1.07302700 0.61440600

H -6.94016700 2.20648100 -0.81286700

C -4.78498200 3.83112300 -0.39198000

H -3.70030100 4.01587900 -0.45027500

H -5.18660100 3.85527800 -1.41595400

H -5.21989000 4.68057300 0.16788600

C -2.20522700 -0.58647000 -0.60403900

C -0.85255000 -0.89231600 -0.31440800

C 0.03915300 0.09547400 0.07013200

C -0.46446300 1.42555500 0.15669300

C -1.79703900 1.66488700 -0.11083000

C -3.17470400 -1.53453700 -1.08989400

C -2.90900100 -2.88733300 -1.42387700

C -3.92227300 -3.72644200 -1.85676900

C -5.23232100 -3.17313400 -1.96101900

C -5.42983800 -1.83926700 -1.66462100

H -0.50950600 -1.92534500 -0.39587200

H 0.19020400 2.25523600 0.43172100

H -2.19733800 2.67733000 -0.04657800

H -1.88994200 -3.26940700 -1.34127500

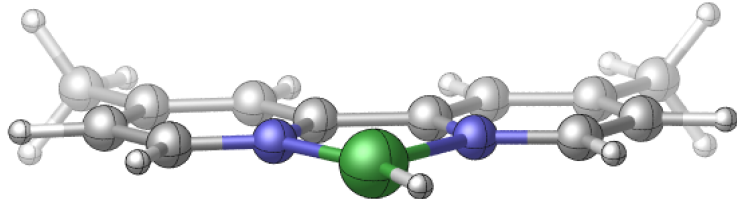
H -6.07577700 -3.79000100 -2.27829500  
H -6.41337500 -1.37414400 -1.76261000  
N -2.67878000 0.70624200 -0.47133900  
N -4.44404300 -1.02209100 -1.23617000  
C 1.48007800 -0.19579000 0.38774600  
H 1.72097600 0.09427000 1.42487200  
H 1.72000200 -1.26210300 0.26533200  
H 2.15178000 0.38656200 -0.26590500  
C -3.67659100 -5.16537400 -2.22023400  
H -4.31313100 -5.83586200 -1.61799400  
H -3.93272000 -5.35359100 -3.27696500  
H -2.62705600 -5.45514600 -2.06506800

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2240.5636168

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2239.8498736



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.218542 (Hartree/Particle)

Thermal correction to Energy= 0.233345

Thermal correction to Enthalpy= 0.234289

Thermal correction to Gibbs Free Energy= 0.174716

Sum of electronic and zero-point Energies= -2082.200898

Sum of electronic and thermal Energies= -2082.186094

Sum of electronic and thermal Enthalpies= -2082.185150

Sum of electronic and thermal Free Energies= -2082.244723

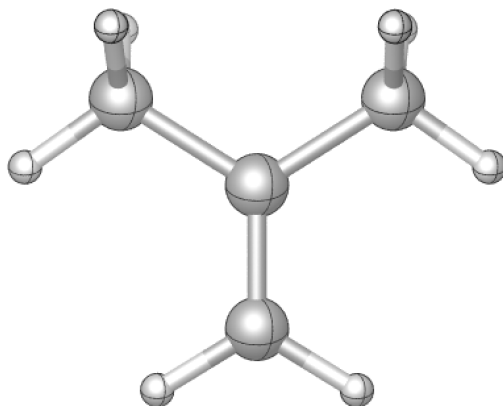
Ni	-4.14128400	1.35473100	-1.69261700
H	-5.44683100	2.14366100	-2.15615700
C	-1.98417500	-0.28633800	-0.86820800
C	-0.70498300	-0.64872200	-0.42793000
C	0.27134100	0.32654400	-0.20166200
C	-0.09836300	1.66595300	-0.43842900
C	-1.38204500	1.95976800	-0.87425400
C	-3.07986300	-1.24022000	-1.13348300
C	-2.98621500	-2.63051500	-0.98125800
C	-4.08889600	-3.44510900	-1.26067400
C	-5.26798000	-2.80868900	-1.69242600
C	-5.29004200	-1.42466400	-1.82069600
H	-0.46479800	-1.69941600	-0.25909600
H	0.61651500	2.47707900	-0.28247700
H	-1.69463300	2.98938000	-1.06507000
H	-2.05320000	-3.08385600	-0.64411100
H	-6.16263600	-3.39015800	-1.92588200
H	-6.18629700	-0.89328500	-2.15166600
N	-2.31931600	1.01465600	-1.08971900
N	-4.22769500	-0.65399600	-1.54992000
C	1.65392700	-0.02271100	0.27689900
H	1.84919000	0.42643700	1.26505500
H	1.79507800	-1.10967900	0.36065200
H	2.41867300	0.37455100	-0.41046400
C	-4.02947500	-4.94106600	-1.11192900
H	-3.05325800	-5.27716600	-0.73493400
H	-4.81014400	-5.29435300	-0.41873800
H	-4.21487300	-5.43465100	-2.08017100

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2083.2746658

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2082.7071176



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.107040 (Hartree/Particle)

Thermal correction to Energy= 0.112322

Thermal correction to Enthalpy= 0.113267

Thermal correction to Gibbs Free Energy= 0.079765

Sum of electronic and zero-point Energies= -157.012996

Sum of electronic and thermal Energies= -157.007713

Sum of electronic and thermal Enthalpies= -157.006769

Sum of electronic and thermal Free Energies= -157.040270

C -5.45780700 2.07262200 0.23723400

C -5.52946100 0.85899400 1.12932400

H -6.18072200 0.07439500 0.71608900

H -4.52384500 0.42950900 1.28546300

H -5.90716300 1.13242700 2.13066000

C -6.12859400 2.15207900 -0.92172300

H -6.76266200 1.33174500 -1.27263500

H -6.06610800 3.04018900 -1.55848800

C -4.57879200 3.19055500 0.73860400

H -3.54036500 2.83933200 0.87450700

H -4.56607400 4.05129000 0.05348300

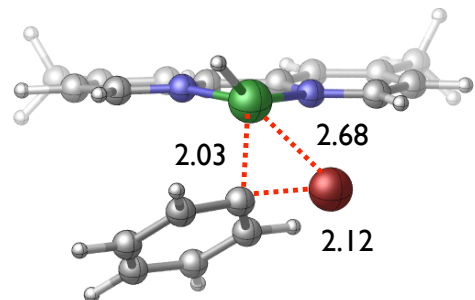
H -4.91872000 3.53998000 1.72977100

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -157.3001298

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -157.1516226



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.309716 (Hartree/Particle)

Thermal correction to Energy= 0.331219

Thermal correction to Enthalpy= 0.332163

Thermal correction to Gibbs Free Energy= 0.255765

Sum of electronic and zero-point Energies= -4887.483265

Sum of electronic and thermal Energies= -4887.461762

Sum of electronic and thermal Enthalpies= -4887.460818

Sum of electronic and thermal Free Energies= -4887.537216

Ni -0.79917400 -0.18317200 0.37337300

C -1.59750800 0.64027300 -1.29816000

C -0.96341900 3.22629700 -2.16800300

C -2.36114100 1.73785700 -0.84662200

C -0.61544700 0.82411200 -2.29669300

C -0.27976900 2.11796100 -2.69156100

C -2.01349800 3.02495500 -1.26137200

H -3.17246700 1.57386600 -0.13808600

H -0.08267700 -0.03906500 -2.69625000

H 0.52534700 2.26263500 -3.41731900

H -2.56944300 3.88025800 -0.86738400

N 0.73831700 1.17004400 0.61386500

C 2.88889900 2.96658200 0.63201800

C 0.57718400 2.41769300 1.06391300

C 1.94612800 0.78038200 0.15858700

C 3.03910900 1.65625900 0.16120700

C 1.61503200 3.34445000 1.08820000

N 0.80701100 -1.25157900 -0.39748800

C 3.15053000 -2.56636100 -1.22288500

C 0.76597600 -2.51433900 -0.84472100

C 1.99870400 -0.61427200 -0.34700400

C 3.17962600 -1.24773700 -0.75376000

C 1.89886900 -3.20297600 -1.26433900

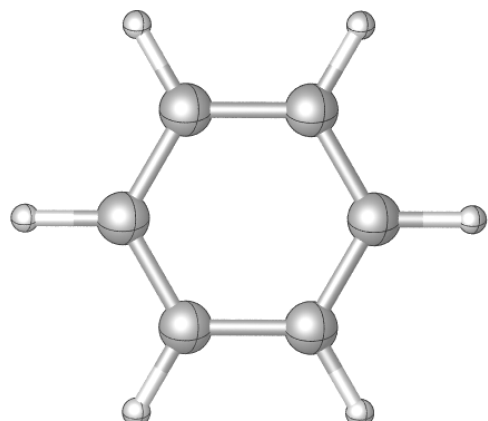
H -0.42788700 2.66934600 1.41062100

H 4.00894200 1.32914000 -0.21439100

H 1.43146900 4.35481000 1.45993100

H -0.22342900 -2.97575700 -0.87505900

H 4.13164700 -0.71891900 -0.70211900  
H 1.80530400 -4.23033100 -1.62269200  
Br -2.58780400 -1.23666900 -1.32829200  
H -0.69604400 4.23708700 -2.48470300  
C 4.40185400 -3.28332100 -1.64903900  
H 4.66136500 -4.06629700 -0.91677400  
H 4.25605200 -3.78560800 -2.61812300  
H 5.25751200 -2.59854300 -1.73256300  
C 4.04019800 3.93441300 0.65931400  
H 3.75413400 4.89726400 0.20758700  
H 4.33939300 4.14378400 1.69999600  
H 4.91734100 3.54304800 0.12497400  
H -1.77995400 0.05875200 1.64063100  
UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4889.1966238  
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4888.3092754



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.100501 (Hartree/Particle)

Thermal correction to Energy= 0.104869

Thermal correction to Enthalpy= 0.105813

Thermal correction to Gibbs Free Energy= 0.073041

Sum of electronic and zero-point Energies= -231.991414

Sum of electronic and thermal Energies= -231.987046

Sum of electronic and thermal Enthalpies= -231.986102

Sum of electronic and thermal Free Energies= -232.018875

C -1.78619400 0.38638600 -1.50220400

C -1.08232300 3.09483000 -1.40580500

C -2.51675000 1.27142400 -0.70025300

C -0.70376700 0.85565300 -2.25604600

C -0.35185200 2.20986100 -2.20777100

C -2.16479500 2.62562500 -0.65210600

H -3.36188700 0.90503200 -0.11142400

H -0.13344100 0.16460300 -2.88246100

H 0.49324200 2.57637100 -2.79661600

H -2.73519400 3.31669300 -0.02575100

H -0.80755600 4.15235000 -1.36828900

H -2.06099000 -0.67106600 -1.53988700

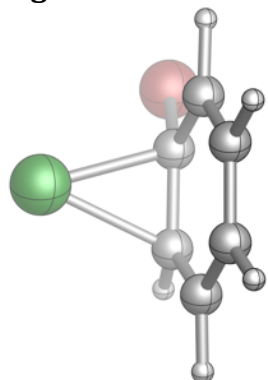
UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -232.3465283

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -232.1429457

**Figure S14**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.090677 (Hartree/Particle)

Thermal correction to Energy= 0.098399

Thermal correction to Enthalpy= 0.099343

Thermal correction to Gibbs Free Energy= 0.055641

Sum of electronic and zero-point Energies= -4313.276593

Sum of electronic and thermal Energies= -4313.268871

Sum of electronic and thermal Enthalpies= -4313.267927

Sum of electronic and thermal Free Energies= -4313.311630

C -3.04310700 0.16108800 1.66008100

C -2.34670500 -0.85049300 0.93821400

C -2.07481300 -0.67622800 -0.45526600

C -2.52185500 0.51012300 -1.09572800

C -3.19602600 1.48917000 -0.38006500

C -3.44095600 1.31673200 1.00062000

Br -2.20662100 -2.61196300 1.72318600

H -1.71985100 -1.51667200 -1.05788600

H -3.26327800 0.01302200 2.71784300

H -3.96559900 2.09429900 1.56028000

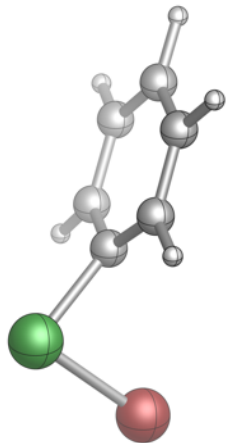
H -3.54390700 2.39323100 -0.88348800

H -2.34869300 0.62453200 -2.16777600

Ni -0.45364000 -0.16362000 0.82203800

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4314.14952807



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.090158 (Hartree/Particle)

Thermal correction to Energy= 0.098267

Thermal correction to Enthalpy= 0.099211

Thermal correction to Gibbs Free Energy= 0.054044

Sum of electronic and zero-point Energies= -4313.273303

Sum of electronic and thermal Energies= -4313.265194

Sum of electronic and thermal Enthalpies= -4313.264250

Sum of electronic and thermal Free Energies= -4313.309417

C -2.63732100 0.92656200 1.28790500

C -1.85059900 -0.10489600 0.78343200

C -1.92216100 -0.59087000 -0.51925500

C -2.75164300 0.10387300 -1.41014300

C -3.51157500 1.19149400 -0.96224800

C -3.45850300 1.60214200 0.37557400

Br -1.50891500 -2.09449000 2.65210700

H -1.34389100 -1.45894000 -0.84267800

H -2.60359000 1.21394500 2.34051600

H -4.06526100 2.44163400 0.72291300

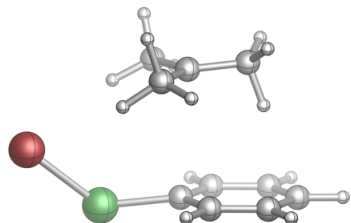
H -4.17022700 1.71302600 -1.66029900

H -2.80843700 -0.22077900 -2.45173900

Ni -0.21539600 -0.46360600 1.69357600

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4314.14216150



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.208392 (Hartree/Particle)

Thermal correction to Energy= 0.223964

Thermal correction to Enthalpy= 0.224908

Thermal correction to Gibbs Free Energy= 0.162774

Sum of electronic and zero-point Energies= -4470.909091

Sum of electronic and thermal Energies= -4470.893518

Sum of electronic and thermal Enthalpies= -4470.892574

Sum of electronic and thermal Free Energies= -4470.954708

Ni -5.69003100 0.64818100 0.19683600

C -5.31262800 2.49967100 0.66139700

C -6.23448800 3.52426700 0.36697900

C -4.17964800 2.83013800 1.43170000

C -6.03777200 4.83176200 0.82555600

H -7.12224700 3.30081900 -0.23322700

C -3.97330700 4.13463800 1.89512600

H -3.44227800 2.05887000 1.67670000

C -4.90263100 5.13978700 1.58920300

H -6.76310200 5.61325100 0.58422500

H -3.08705600 4.37265000 2.48928000

H -4.74153100 6.16071500 1.94406100

Br -5.68614100 -0.46437700 -1.89372500

C -3.77679300 3.39391200 -1.98984200

C -2.89096800 2.24435300 -1.77541200

H -3.42256200 1.28372100 -1.75682300

H -2.23827200 2.24155400 -2.67717100

H -2.22444500 2.38095000 -0.91340100

C -3.36068800 4.72070500 -1.55843600

H -3.85202000 4.85510400 -0.56365600

H -2.28012400 4.80823100 -1.38558500

H -3.76455200 5.52248000 -2.19272700

C -5.01632300 3.22690400 -2.75395700

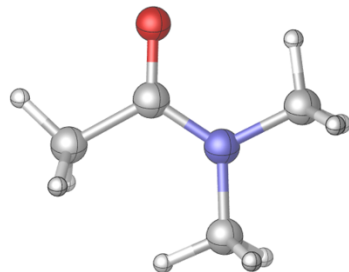
H -5.49823400 2.25608700 -2.56766100

H -5.70555800 4.07386700 -2.65002300

H -4.67682000 3.20337300 -3.81377000

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4472.07517251

**DMA**

UB3LYP-D3/def2-SVP-CPCM(THF)

 $\langle S^2 \rangle = 0.0000$ 

Zero-point correction= 0.129574 (Hartree/Particle)

Thermal correction to Energy= 0.137214

Thermal correction to Enthalpy= 0.138159

Thermal correction to Gibbs Free Energy= 0.097818

Sum of electronic and zero-point Energies= -287.506478

Sum of electronic and thermal Energies= -287.498838

Sum of electronic and thermal Enthalpies= -287.497894

Sum of electronic and thermal Free Energies= -287.538234

C -2.77549400 1.10576800 0.17344400

O -3.37576400 0.04098600 0.30285000

N -1.42066200 1.15712500 -0.02483700

C -3.53169900 2.42604000 0.23195600

H -3.17999200 3.06144900 1.05983000

H -3.42570500 3.00001200 -0.70177600

H -4.59143000 2.19194000 0.38873300

C -0.64344600 2.37040000 -0.20318800

H -1.25639800 3.26981200 -0.09049600

H 0.17034500 2.41803800 0.54165400

H -0.18014800 2.39406000 -1.20606400

C -0.65581000 -0.07661500 -0.10272300

H -1.32981400 -0.92785900 0.04169300

H -0.16180500 -0.16647800 -1.08632700

H 0.12810200 -0.09653400 0.67449200

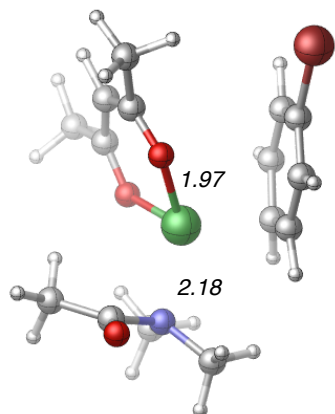
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -287.968183

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -287.7556375

**<sup>2</sup>A-DMA-N**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.334610 (Hartree/Particle)

Thermal correction to Energy= 0.359997

Thermal correction to Enthalpy= 0.360941

Thermal correction to Gibbs Free Energy= 0.276130

Sum of electronic and zero-point Energies= -4945.881912

Sum of electronic and thermal Energies= -4945.856525

Sum of electronic and thermal Enthalpies= -4945.855581

Sum of electronic and thermal Free Energies= -4945.940391

C -2.38385200 1.54700000 -1.19046200

O -2.56104700 1.50095400 -2.39614100

N -1.37287000 0.74453500 -0.58367600

C -3.17663100 2.47036200 -0.29399500

H -3.55728500 1.94507700 0.59047500

H -2.53641800 3.30063700 0.04600600

H -4.00683300 2.87792300 -0.88264400

C -0.33484600 0.28781600 -1.52187900

H 0.34963800 1.11482100 -1.77797300

H 0.24348600 -0.51919100 -1.05542400

H -0.80658500 -0.07658600 -2.43881000

C -0.81492300 1.17330700 0.71106500

H -0.21818800 2.09533400 0.60455600

H -1.61496000 1.31223300 1.44292300

H -0.16279700 0.37309800 1.08604200

Ni -2.85937700 -0.81031400 -0.20266100

C -1.59741300 -2.49138000 0.12393600

C -1.82822800 -2.75913400 -1.24900300

C -2.83693300 -3.65734500 -1.64005400

C -3.60617300 -4.28244600 -0.66447200

C -3.38749600 -4.04805600 0.70562100

C -2.38182400 -3.17159900 1.09077500

H -0.68264900 -1.98414900 0.44480200

H -1.16989500 -2.34103900 -2.01103100

H -4.00662500 -4.54478200 1.45386400  
H -2.19922200 -2.99115700 2.15173500  
H -3.01374000 -3.85569400 -2.69803300  
Br -5.00080200 -5.48418200 -1.18944900  
C -5.66791900 -1.42806900 0.14358400  
O -4.68295600 -1.40061200 -0.65387600  
C -5.64863800 -0.98083500 1.48607200  
H -6.56505600 -1.09680800 2.06563100  
C -4.56163700 -0.34030500 2.11684600  
O -3.43375200 -0.09849100 1.58088900  
C -6.94534600 -2.01407900 -0.41916700  
H -6.76202500 -3.06891200 -0.68247200  
H -7.79456700 -1.95707900 0.27521100  
H -7.20169600 -1.49108000 -1.35446600  
C -4.71613900 0.13944400 3.54551000  
H -5.67601600 -0.14327700 3.99848000  
H -3.89140200 -0.26777100 4.15242000  
H -4.61752500 1.23764400 3.56964300

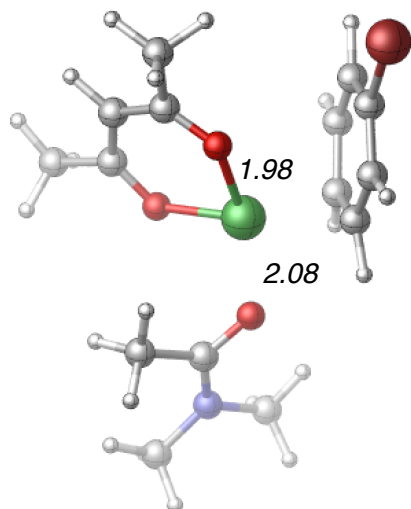
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.7109019

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.8406731

**<sup>2</sup>A-DMA-O**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.333954 (Hartree/Particle)

Thermal correction to Energy= 0.359909

Thermal correction to Enthalpy= 0.360854

Thermal correction to Gibbs Free Energy= 0.272367

Sum of electronic and zero-point Energies= -4945.890866

Sum of electronic and thermal Energies= -4945.864911

Sum of electronic and thermal Enthalpies= -4945.863966

Sum of electronic and thermal Free Energies= -4945.952453

C -0.86706700 1.17746200 -0.95642900

O -1.02443200 -0.05873500 -1.01941100

N 0.37547300 1.70557900 -0.88856000

C -2.05802200 2.11220400 -0.97006500

H -2.12290700 2.68122800 -0.03129400

H -1.99841800 2.82320400 -1.80761900

H -2.96704100 1.50923000 -1.06437000

C 1.54849100 0.84014400 -0.89418900

H 2.19553900 1.08629000 -1.75258200

H 2.13121500 0.98605900 0.03045300

H 1.23403000 -0.20563700 -0.96489100

C 0.66981200 3.13028300 -0.83989000

H 1.19171200 3.44869700 -1.75861100

H -0.23689200 3.73163700 -0.72865300

H 1.32934300 3.34351900 0.01719400

Ni -2.70280800 -1.11931500 -0.39915300

C -1.58501700 -2.90884700 -0.31627200

C -2.03253700 -2.81770200 -1.66571600

C -3.15090700 -3.57289500 -2.09104400

C -3.81874500 -4.37156000 -1.17714900

C -3.38044100 -4.49446100 0.15961000

C	-2.26391100	-3.78563400	0.57266500
H	-0.60565900	-2.51218800	-0.03522000
H	-1.42548700	-2.29945900	-2.41173600
H	-3.91793600	-5.14219300	0.85425500
H	-1.90366000	-3.89625500	1.59837400
H	-3.49011500	-3.51044800	-3.12624800
Br	-5.38985200	-5.32520000	-1.72296400
C	-5.45629800	-1.19052300	0.55674200
O	-4.63582000	-1.55383600	-0.33846700
C	-5.15679300	-0.35561800	1.65824300
H	-5.96562100	-0.13106600	2.35471000
C	-3.90139900	0.24895600	1.89761700
O	-2.86127100	0.09400400	1.18780900
C	-6.86784600	-1.71878700	0.39092700
H	-6.83210900	-2.81883500	0.33026100
H	-7.54664700	-1.41808400	1.20076700
H	-7.27395600	-1.36419700	-0.57092500
C	-3.74925500	1.18362100	3.08258300
H	-4.64679800	1.23904000	3.71390000
H	-2.89252300	0.85729700	3.69413900
H	-3.50941300	2.19501100	2.71330100

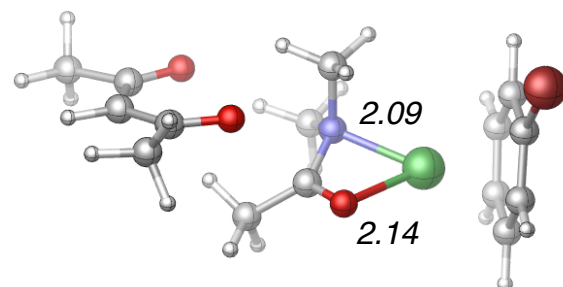
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.7206989

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.8519092

**<sup>2</sup>A-DMA'-conf1**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.333515 (Hartree/Particle)

Thermal correction to Energy= 0.359366

Thermal correction to Enthalpy= 0.360310

Thermal correction to Gibbs Free Energy= 0.272439

Sum of electronic and zero-point Energies= -4945.808783

Sum of electronic and thermal Energies= -4945.782931

Sum of electronic and thermal Enthalpies= -4945.781987

Sum of electronic and thermal Free Energies= -4945.869858

C -1.76896800 -1.05834000 -1.69978000

O -2.82325200 -1.09231900 -1.06125600

N -0.58437700 -0.92091500 -0.90100900

C -1.66946700 -1.35361400 -3.15555400

H -0.93675900 -0.68494500 -3.62107200

H -1.34230900 -2.39886300 -3.28715000

H -2.65950300 -1.23344400 -3.60939200

C 0.66491700 -1.47728000 -1.45110200

H 1.03663800 -0.85964200 -2.28180700

H 1.40664000 -1.48964300 -0.64016800

H 0.49895100 -2.51356400 -1.77291100

C -0.37715200 0.44648200 -0.33981100

H 0.04488100 1.08654600 -1.12503300

H -1.34179300 0.85730100 -0.02381000

H 0.29271400 0.35645500 0.52685200

Ni -1.69568400 -1.88151400 0.58388000

C -1.09472400 -3.69294800 1.63628900

C -2.50423200 -3.59613200 1.66638100

C -3.12291600 -2.47049000 2.25486800

C -2.33578900 -1.46444700 2.85918700

C -0.93345900 -1.54421600 2.82530700

C -0.32220300 -2.64999500 2.19317900

H -0.60804800 -4.55054200 1.17173200

H -3.12287000 -4.37377000 1.21740000

H -0.32313000 -0.75156700 3.25781700

H 0.76606200 -2.68901400 2.12960900

H -4.20838700 -2.37567100 2.23749800

Br -3.18322100 0.01458400 3.69556100

C	-2.17004800	1.92339200	-3.58430000
O	-2.10679600	1.43491100	-2.43352100
C	-1.12117800	1.99559600	-4.54397800
H	-1.36831500	2.44927000	-5.50674100
C	0.22430400	1.56901900	-4.36536700
O	0.71147400	1.04080100	-3.33817400
C	-3.53208000	2.49251800	-4.00121100
H	-3.85742000	3.22898500	-3.24757500
H	-3.54184800	2.96506900	-4.99459900
H	-4.27716100	1.67856400	-3.98559300
C	1.18034500	1.76458300	-5.54801300
H	0.72798600	2.26269900	-6.41825500
H	2.05205800	2.34956500	-5.20985000
H	1.56650400	0.77976400	-5.86168700

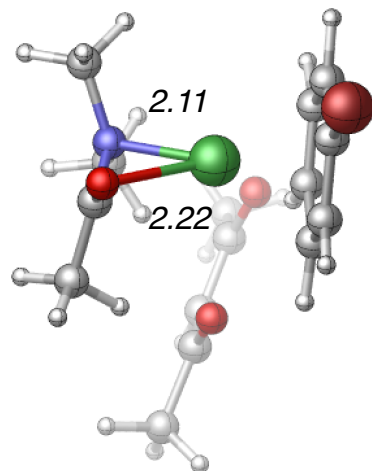
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.6428298

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.781502

<sup>2</sup>A-DMA'-conf2



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.333505 (Hartree/Particle)

Thermal correction to Energy= 0.359314

Thermal correction to Enthalpy= 0.360258

Thermal correction to Gibbs Free Energy= 0.274476

Sum of electronic and zero-point Energies= -4945.816723

Sum of electronic and thermal Energies= -4945.790914

Sum of electronic and thermal Enthalpies= -4945.789970

Sum of electronic and thermal Free Energies= -4945.875752

C -1.88767300 -0.84033800 -1.04841300

O -2.60783000 -0.12550100 -0.32523300

N -0.51147500 -0.89937600 -0.73734300

C -2.41367300 -1.63218500 -2.20696500

H -1.85094300 -1.39745200 -3.12329500

H -2.29278700 -2.71307000 -2.00725900

H -3.47479500 -1.38929500 -2.33665700

C 0.36458600 -1.82564000 -1.47047100

H 0.60359700 -1.43086600 -2.47193300

H 1.29135100 -1.94890000 -0.89699700

H -0.10477800 -2.81395900 -1.52797000

C 0.14916000 0.33936700 -0.28466600

H 0.59081300 0.86485400 -1.14804000

H -0.58281200 0.98695400 0.20944900

H 0.94454500 0.08248800 0.42718100

Ni -1.57854600 -1.64404200 0.92855200

C -0.79633600 -3.62215300 1.45757300

C -2.20485100 -3.71757600 1.42065400

C -2.99164900 -2.89024200 2.24863300

C -2.36196500 -1.98314200 3.13235800

C -0.95844600 -1.88392400 3.19573600

C -0.18778600 -2.70092100 2.33966500

H	-0.18540400	-4.18833100	0.74267700
H	-2.66798900	-4.38845500	0.69781200
H	-0.48285100	-1.17070700	3.86834800
H	0.89873400	-2.59778600	2.33882400
H	-4.07943500	-2.92305100	2.19034500
Br	-3.43403700	-0.86500700	4.23749400
C	-0.95743000	-5.24687700	-2.30010800
O	-1.46421300	-4.53735600	-1.39076400
C	0.40181000	-5.63448900	-2.40272300
H	0.67614500	-6.22996200	-3.27663100
C	1.44981000	-5.33198600	-1.48222200
O	1.35003400	-4.70111100	-0.40715000
C	-1.90361400	-5.68993500	-3.41763500
H	-2.24358200	-4.79740500	-3.97143400
H	-1.45719000	-6.39744000	-4.13178100
H	-2.80241400	-6.14381400	-2.96898600
C	2.85423500	-5.82774200	-1.84744000
H	2.90314100	-6.40103600	-2.78479600
H	3.52921700	-4.95856300	-1.92525300
H	3.23930700	-6.45079100	-1.02296600

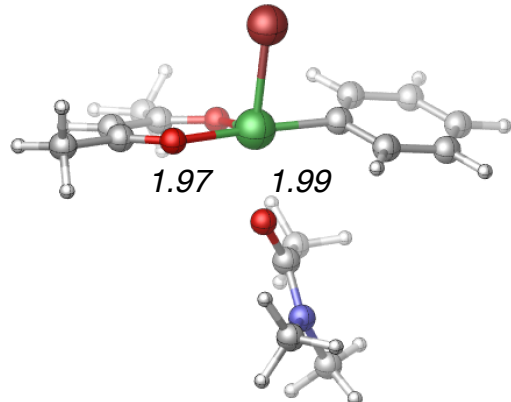
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.6480744

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.7844496

## <sup>2</sup>C-DMA-O



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.335220 (Hartree/Particle)

Thermal correction to Energy= 0.360904

Thermal correction to Enthalpy= 0.361848

Thermal correction to Gibbs Free Energy= 0.275250

Sum of electronic and zero-point Energies= -4945.908362

Sum of electronic and thermal Energies= -4945.882678

Sum of electronic and thermal Enthalpies= -4945.881734

Sum of electronic and thermal Free Energies= -4945.968331

C	-2.72312300	0.16404400	-0.21128400
O	-1.94642200	-0.57518600	-0.86605100
N	-2.21526200	1.05771300	0.65247600
C	-4.21765800	0.07104800	-0.37844300
H	-4.65738900	-0.43891400	0.49349000
H	-4.68111300	1.06314300	-0.47105600
H	-4.42823600	-0.52882300	-1.26950100
C	-3.05005100	1.95397900	1.44806100
H	-2.48680000	2.25413600	2.34170300
H	-3.31813700	2.86365200	0.88478000
H	-3.96799200	1.45619400	1.78328200
C	-0.77110100	1.22317500	0.79533900
H	-0.49927800	2.27675700	0.62342000
H	-0.45500500	0.94083600	1.81257700
H	-0.25444200	0.59111300	0.06665600
Ni	-2.13583600	-2.30617300	-1.82492900
C	-3.38599700	-3.35898200	1.99830200
C	-2.61469400	-3.00223900	0.88261600
C	-3.20029800	-3.02036800	-0.38301800
C	-4.52367500	-3.41253300	-0.56565600
C	-5.28795800	-3.76754100	0.55458700
C	-4.72306500	-3.73915900	1.83541000
H	-2.93628800	-3.33712600	2.99500900
H	-1.57211100	-2.69925400	1.01066300

H -4.95718000 -3.41711400 -1.56654900  
H -6.33158600 -4.06686500 0.42185700  
H -5.32322900 -4.01714000 2.70561000  
Br -1.57001500 -4.55993400 -2.32136400  
C -3.62262900 -1.35164500 -4.12517800  
O -3.69114100 -1.86408700 -2.95889500  
C -2.44511800 -0.99835600 -4.80590200  
H -2.54811400 -0.57952300 -5.80649000  
C -1.13753900 -1.12968700 -4.28123200  
O -0.85029900 -1.59082900 -3.13839600  
C -4.96290800 -1.11180700 -4.78276300  
H -5.53193500 -0.38740400 -4.17682500  
H -4.87724100 -0.73298200 -5.80965800  
H -5.53811100 -2.05118400 -4.78337100  
C 0.04290300 -0.70039700 -5.12368000  
H -0.25102200 -0.25180500 -6.08169000  
H 0.65229600 0.01769300 -4.55237700  
H 0.67916200 -1.57992700 -5.31529200

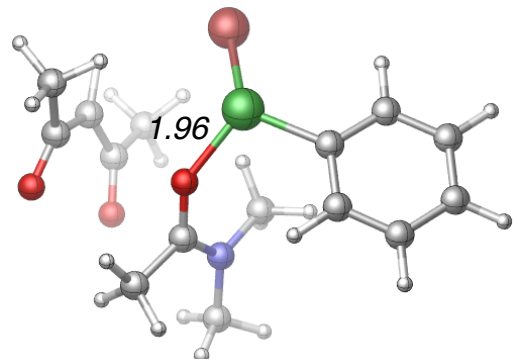
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.7369949

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.8553074

<sup>2</sup>C-DMA'-conf1



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.332583 (Hartree/Particle)

Thermal correction to Energy= 0.359318

Thermal correction to Enthalpy= 0.360262

Thermal correction to Gibbs Free Energy= 0.270529

Sum of electronic and zero-point Energies= -4945.848754

Sum of electronic and thermal Energies= -4945.822019

Sum of electronic and thermal Enthalpies= -4945.821075

Sum of electronic and thermal Free Energies= -4945.910808

C	-0.46249000	-1.46454300	-4.21905300
O	-1.34334000	-0.66262100	-3.80734000
N	0.52565700	-1.90817000	-3.43606900
C	-0.52213100	-1.90221900	-5.65882300
H	0.39506000	-1.59029400	-6.17737700
H	-0.63077100	-2.99403300	-5.74388900
H	-1.37721000	-1.40913600	-6.13144800
C	1.58082300	-2.79211400	-3.92336500
H	2.43412700	-2.19163500	-4.27828900
H	1.91214400	-3.43893300	-3.09925300
H	1.22907500	-3.43654500	-4.73540200
C	0.73339300	-1.37632900	-2.09107200
H	1.81332300	-1.30158400	-1.90572000
H	0.29755800	-0.37667300	-1.99651800
H	0.27365300	-2.02967400	-1.33424600
Ni	-2.33094100	-0.14263900	-2.20039400
C	-2.74566800	-2.84195600	1.08050100
C	-2.61783000	-1.68172200	0.30469400
C	-2.54896800	-1.74000000	-1.10003100
C	-2.61071300	-3.01274100	-1.70061100
C	-2.73837300	-4.17978900	-0.93515800
C	-2.80601000	-4.09614500	0.46069300
H	-2.79457000	-2.76899800	2.17145900
H	-2.56040800	-0.70921100	0.80529100
H	-2.55297800	-3.10178500	-2.79133300

H	-2.78259600	-5.15644200	-1.42725500
H	-2.90224800	-5.00408200	1.06279200
Br	-1.43760100	1.85644400	-1.22776000
C	-0.69949200	1.75083400	-5.43385200
O	-0.35133500	1.07006000	-6.39037900
C	0.21237100	2.09348100	-4.33103900
H	-0.07679100	2.92195000	-3.68052400
C	1.45703600	1.39228200	-4.00544900
O	1.83329600	0.39214200	-4.61319600
C	-2.09775600	2.32091600	-5.32193000
H	-2.73897300	1.88686700	-6.10004300
H	-2.51633400	2.12603700	-4.32275000
H	-2.06112400	3.41653800	-5.44755000
C	2.23643700	1.94952500	-2.83293300
H	1.59769100	1.94290200	-1.93415400
H	3.13808400	1.34726900	-2.66201300
H	2.51748700	2.99881500	-3.02071500

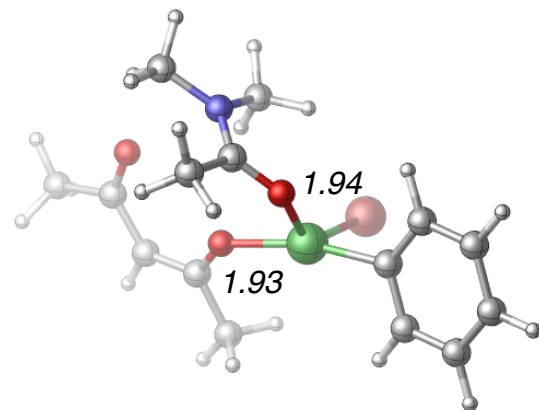
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4947.6653381

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.7828069

<sup>2</sup>C-DMA'-conf2



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.334582 (Hartree/Particle)

Thermal correction to Energy= 0.360359

Thermal correction to Enthalpy= 0.361303

Thermal correction to Gibbs Free Energy= 0.275123

Sum of electronic and zero-point Energies= -4945.869422

Sum of electronic and thermal Energies= -4945.843645

Sum of electronic and thermal Enthalpies= -4945.842701

Sum of electronic and thermal Free Energies= -4945.928881

C	-2.01216100	0.86910700	-0.04072900
O	-2.27815700	-0.35815900	-0.22735100
N	-1.12610100	1.26725300	0.86415700
C	-2.77296000	1.87403300	-0.86856500
H	-3.23419700	2.66188900	-0.25677200
H	-2.08983800	2.35148400	-1.58917400
H	-3.55218800	1.33887400	-1.42306500
C	-0.72519000	2.65795000	1.06780000
H	-0.90266600	2.94235000	2.11693000
H	0.34350700	2.74853500	0.83116300
H	-1.28654200	3.33786700	0.42091000
C	-0.35376100	0.28037700	1.61671900
H	-0.02560500	0.73412200	2.56082000
H	-0.97005500	-0.59962400	1.83459000
H	0.51945900	-0.01674600	1.02005000
Ni	-1.14441600	-1.85866500	-0.70746700
C	-4.75390500	-3.68551400	0.41638200
C	-3.58731600	-2.91548200	0.50658100
C	-2.60890600	-3.08051400	-0.46876600
C	-2.72666600	-4.01280000	-1.49678300
C	-3.90637800	-4.76990100	-1.58229500
C	-4.91387700	-4.60767500	-0.62664800
H	-5.54179800	-3.55949200	1.16398500
H	-3.45768800	-2.18225800	1.30413600

H -1.92404700 -4.16192300 -2.22475700  
H -4.02272900 -5.49525800 -2.39200600  
H -5.82484300 -5.20822200 -0.68808800  
Br 0.08631200 -3.32753100 0.60460900  
C 1.27103400 -0.89155200 -2.07248900  
O 0.19192900 -0.64673100 -1.40498000  
C 2.37996900 -0.05640200 -2.07907800  
H 3.20646300 -0.33282200 -2.73794600  
C 2.55225900 1.14703500 -1.29164100  
O 1.74485000 1.58272700 -0.46372800  
C 1.28738000 -2.16325800 -2.89682200  
H 0.39033100 -2.19207500 -3.53830900  
H 2.18589500 -2.27002900 -3.51926300  
H 1.21862300 -3.02708800 -2.21395700  
C 3.86097600 1.90096200 -1.52779500  
H 3.93606300 2.22819800 -2.57835100  
H 3.90766400 2.77801800 -0.86770600  
H 4.72821700 1.24831600 -1.33297200

UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

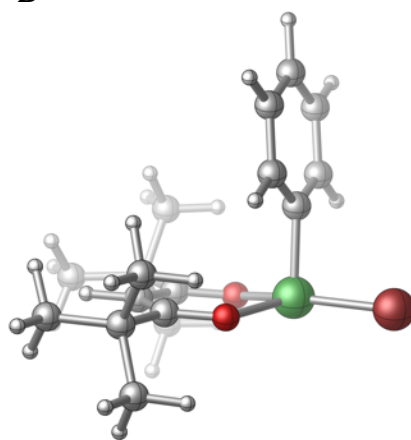
HF= -4947.6993646

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4946.8143409

**Figure S15**

$^2\text{B}^{TMHD}$



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.372835 (Hartree/Particle)

Thermal correction to Energy= 0.396936

Thermal correction to Enthalpy= 0.397880

Thermal correction to Gibbs Free Energy= 0.316227

Sum of electronic and zero-point Energies= -4893.938481

Sum of electronic and thermal Energies= -4893.914380

Sum of electronic and thermal Enthalpies= -4893.913436

Sum of electronic and thermal Free Energies= -4893.995089

C -2.56869900 1.06653100 1.31644600

C -1.72660600 0.13435400 0.72201700

C -1.72732600 -0.12971300 -0.64220700

C -2.61211200 0.59908600 -1.45179000

C -3.46751600 1.55172000 -0.88821000

C -3.44801300 1.78355000 0.49177300

Br -1.75701700 -2.55072000 2.20848200

H -1.05887500 -0.87635900 -1.06983800

H -2.54326000 1.24830600 2.39274000

H -4.11359100 2.52760500 0.93735000

H -4.15375800 2.11420600 -1.52591300

H -2.62720700 0.41151600 -2.52880500

Ni -0.42983800 -0.68760000 1.84004300

O 0.93295200 -1.16607200 0.54263200

O 0.39100300 0.96684900 2.23863900

C 1.28316400 1.56067900 1.53832600

C 1.79673800 -0.37119200 0.04939100

C 2.65607100 -0.99141700 -1.06922800

C 1.52176100 3.02424000 1.95026700

C 1.97810800 0.96352900 0.47281000

H 2.71755000 1.56234200 -0.04319500

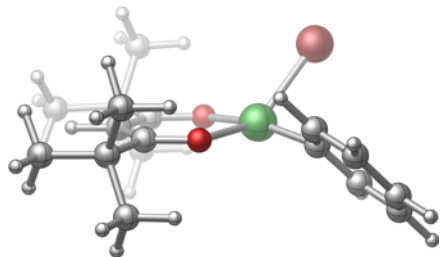
C 1.69380800 -1.49935100 -2.16566200

H	0.98004100	-2.22654400	-1.75315500
H	1.12391500	-0.66664400	-2.60842600
H	2.26532700	-1.98824600	-2.97009600
C	3.66344500	-0.01191800	-1.68855700
H	4.38344600	0.36223100	-0.94467900
H	4.23705700	-0.52670700	-2.47498400
H	3.16265400	0.85147200	-2.15325400
C	3.41496200	-2.18645700	-0.45034800
H	3.99998400	-2.70370900	-1.22715900
H	4.11160300	-1.84858400	0.33394300
H	2.71297700	-2.90450200	-0.00295700
C	2.69077000	3.68788200	1.20795500
H	2.81275200	4.72079600	1.56925000
H	3.63951800	3.15753900	1.38389700
H	2.51530400	3.73733500	0.12249100
C	1.79255100	3.06090600	3.46923300
H	1.90301400	4.10371300	3.80558200
H	0.96580200	2.59911200	4.02657100
H	2.72094900	2.52151300	3.71750400
C	0.21249400	3.78414400	1.63441400
H	-0.00553100	3.76202100	0.55459400
H	-0.63914600	3.33245900	2.16268100
H	0.30437000	4.83669000	1.94595400

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4895.7394255

<sup>2</sup>C<sup>TMHD</sup>



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.372626 (Hartree/Particle)

Thermal correction to Energy= 0.396808

Thermal correction to Enthalpy= 0.397752

Thermal correction to Gibbs Free Energy= 0.315737

Sum of electronic and zero-point Energies= -4893.929733

Sum of electronic and thermal Energies= -4893.905551

Sum of electronic and thermal Enthalpies= -4893.904607

Sum of electronic and thermal Free Energies= -4893.986622

C -3.03615900 -1.47922900 -1.00536000

O -3.58360700 -0.31983100 -1.08381500

C -3.72718600 -2.69444500 -0.92015000

H -3.14584600 -3.60673300 -0.88326300

C -5.13538500 -2.79891800 -0.83990500

O -5.91740500 -1.79941500 -0.86169000

C -5.84418000 -4.16080100 -0.71787200

C -1.49617300 -1.41323600 -1.00946800

Ni -5.43455200 0.05667100 -1.16028600

C -5.07611800 1.92976300 -1.04472300

C -4.02526300 2.51299000 -1.74720500

C -5.81228300 2.63386400 -0.09387100

C -3.67941300 3.84006300 -1.46191200

H -3.47149700 1.93696700 -2.49080200

C -5.44910000 3.95961700 0.19256000

H -6.66031600 2.17381100 0.42047400

C -4.38776800 4.56061800 -0.49129000

H -2.85118600 4.30956500 -1.99978500

H -6.00953300 4.52070500 0.94532200

H -4.11536900 5.59643300 -0.27374000

Br -7.01082900 0.53755800 -2.78894400

C -0.82487900 -2.79026300 -0.90392000

H 0.26880700 -2.66309600 -0.90844200

H -1.09603800 -3.30833600 0.02879000

H -1.08623300 -3.44054200 -1.75290200

C -1.06736500 -0.53931500 0.19008600

H -1.52928500 0.45628300 0.13242800

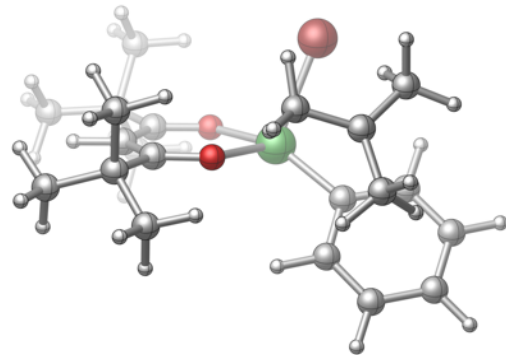
H -1.36436100 -1.00500600 1.14367200

H 0.02751600 -0.42002100 0.19436000  
C -1.06446000 -0.73322900 -2.32785900  
H -1.37069900 -1.33269600 -3.20032400  
H -1.51391800 0.26561900 -2.41892200  
H 0.03150500 -0.62708200 -2.35371200  
C -4.88062500 -5.35397100 -0.64152600  
H -4.25497900 -5.43533600 -1.54363900  
H -4.21904700 -5.29123900 0.23639500  
H -5.45993100 -6.28646500 -0.55518600  
C -6.74880700 -4.30624300 -1.96245300  
H -7.43935700 -3.45456000 -2.04257700  
H -6.14709900 -4.34976800 -2.88470800  
H -7.33724200 -5.23482400 -1.89380000  
C -6.71490500 -4.12101300 0.55675900  
H -7.28836300 -5.05672800 0.64971000  
H -6.09018400 -4.01201200 1.45819700  
H -7.42044400 -3.27901100 0.52120300

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4895.73076529

**<sup>3</sup>C-complex<sup>TMHD</sup>**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.489906 (Hartree/Particle)

Thermal correction to Energy= 0.521374

Thermal correction to Enthalpy= 0.522318

Thermal correction to Gibbs Free Energy= 0.425218

Sum of electronic and zero-point Energies= -5051.534392

Sum of electronic and thermal Energies= -5051.502925

Sum of electronic and thermal Enthalpies= -5051.501980

Sum of electronic and thermal Free Energies= -5051.599081

C -4.19805300 -1.47087900 2.19745700

O -4.50713200 -0.26041500 2.40633100

C -3.64201300 -1.95701600 0.98997300

H -3.38548300 -3.00792600 0.94331000

C -3.40732200 -1.18306300 -0.16154400

O -3.67565300 0.06084000 -0.26625000

C -2.82101300 -1.83063900 -1.43715800

C -4.49347400 -2.42012600 3.38050200

Ni -4.38692600 1.27767400 1.14689900

C -3.53261500 2.98911100 0.63333600

C -4.09190300 4.26960500 0.82963200

C -2.20030100 2.95478800 0.16314500

C -3.37483200 5.44670800 0.57720100

H -5.12660000 4.34462800 1.18194500

C -1.46416700 4.11968700 -0.08781100

H -1.72921700 1.98262600 -0.02135600

C -2.05271000 5.37490200 0.11651100

H -3.84425200 6.42272600 0.73669100

H -0.43408700 4.05370600 -0.45251500

H -1.48788900 6.28958100 -0.08493700

Br -6.83112500 1.61530300 0.78921500

C -4.86396000 3.06426000 -2.38815600

C -3.64646000 3.80785300 -2.72703000

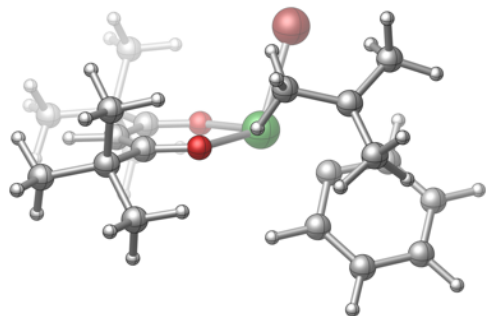
H -2.77298600 3.16917100 -2.90126600

H -3.89618800 4.33714800 -3.67169600

H	-3.43193200	4.59688000	-1.98973500
C	-6.02730300	3.75295600	-1.85560300
H	-6.27492900	3.23522900	-0.88809900
H	-5.88983200	4.82841100	-1.70092200
H	-6.91126200	3.52976500	-2.47876300
C	-4.92032100	1.62609300	-2.60312500
H	-4.39980600	1.18120800	-1.71248700
H	-5.93601900	1.21352600	-2.61584200
H	-4.31876400	1.30951200	-3.46806300
C	-3.99765900	-2.04299800	-2.41666900
H	-4.50980500	-1.09613700	-2.63543100
H	-3.62975600	-2.46321500	-3.36682800
H	-4.73811400	-2.74187400	-1.99634900
C	-1.79247300	-0.86164300	-2.05512900
H	-0.94894400	-0.69247700	-1.36629000
H	-1.39075900	-1.28343000	-2.98995400
H	-2.24412200	0.11364500	-2.27809000
C	-2.13552100	-3.18274700	-1.17725000
H	-1.33979400	-3.09453500	-0.42124800
H	-2.84676500	-3.95210300	-0.84154600
H	-1.67782500	-3.54796000	-2.11010200
C	-4.03481600	-3.86790200	3.15263100
H	-4.54509200	-4.33078300	2.29399500
H	-2.94829100	-3.93038100	2.98467900
H	-4.26982000	-4.47309900	4.04257900
C	-3.78742500	-1.85256500	4.62981700
H	-4.10196500	-0.81629000	4.81677700
H	-4.03378000	-2.46071900	5.51490000
H	-2.69233900	-1.86283400	4.50307800
C	-6.02375400	-2.39487900	3.59346500
H	-6.29684300	-3.00026000	4.47277700
H	-6.37558700	-1.36501800	3.74989800
H	-6.55016300	-2.80707100	2.71709800

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -5053.62941616

**<sup>3</sup>C-TS<sup>TMHD</sup>**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.490206 (Hartree/Particle)

Thermal correction to Energy= 0.521083

Thermal correction to Enthalpy= 0.522027

Thermal correction to Gibbs Free Energy= 0.426260

Sum of electronic and zero-point Energies= -5051.532776

Sum of electronic and thermal Energies= -5051.501899

Sum of electronic and thermal Enthalpies= -5051.500955

Sum of electronic and thermal Free Energies= -5051.596722

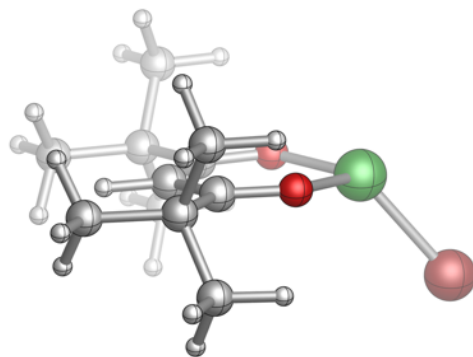
C	-4.14753400	-1.48132700	2.30746700
O	-4.44355000	-0.25668700	2.44671400
C	-3.54744000	-2.03086700	1.15083200
H	-3.31137500	-3.08747600	1.15977900
C	-3.22455700	-1.29905500	-0.00872700
O	-3.47446100	-0.06153200	-0.17935400
C	-2.54038500	-1.98320100	-1.21504200
C	-4.51150100	-2.36708500	3.51972900
Ni	-4.34960200	1.19437100	1.09370400
C	-3.58220100	2.93087200	0.48779900
C	-4.14597100	4.19912300	0.75030900
C	-2.24245000	2.92270400	0.02939800
C	-3.42897200	5.38828400	0.57152100
H	-5.18348000	4.25104300	1.09742400
C	-1.50639600	4.09787000	-0.14584200
H	-1.77131500	1.96058700	-0.20018500
C	-2.10333200	5.34043800	0.11802800
H	-3.90014900	6.35387200	0.77915200
H	-0.47168900	4.05531300	-0.49971000
H	-1.53881400	6.26508900	-0.03090500
Br	-6.81827500	1.41346700	0.81869300
C	-4.80157500	3.10131700	-2.19963000
C	-3.66950100	3.87909300	-2.71678800
H	-2.79502500	3.26976500	-2.96958300
H	-4.04306200	4.39922900	-3.62213900
H	-3.39483700	4.68633700	-2.01619500

C	-6.03327600	3.78608800	-1.79967500
H	-6.44540600	3.30572100	-0.89038200
H	-5.92497300	4.87084800	-1.68731900
H	-6.77949600	3.55714800	-2.58873600
C	-4.81055300	1.64007500	-2.33682300
H	-3.89070000	1.19522500	-1.92766300
H	-5.68174500	1.16981800	-1.86640100
H	-4.80286000	1.43280500	-3.42681200
C	-1.36682000	-1.09243100	-1.67365800
H	-0.60786900	-1.00101100	-0.87959800
H	-0.88241900	-1.53201300	-2.56002000
H	-1.71247300	-0.08167900	-1.93039000
C	-3.59341200	-2.07608500	-2.34200800
H	-4.44565900	-2.70332800	-2.03499400
H	-3.97871700	-1.08155200	-2.60630200
H	-3.14588500	-2.52411300	-3.24397500
C	-2.00793200	-3.39155900	-0.90638300
H	-2.81605800	-4.09348400	-0.65093200
H	-1.49371400	-3.79320600	-1.79391100
H	-1.28532900	-3.37895800	-0.07559900
C	-6.04880200	-2.30525100	3.66341200
H	-6.37116500	-2.86187300	4.55816400
H	-6.38771800	-1.26317100	3.75271400
H	-6.54428300	-2.75089000	2.78539500
C	-4.07225900	-3.83210800	3.38081800
H	-4.55138400	-4.32532800	2.52106400
H	-2.98057600	-3.92334400	3.26785400
H	-4.36123100	-4.39066500	4.28530800
C	-3.84905200	-1.75312600	4.77088600
H	-4.14529300	-2.31349400	5.67213300
H	-2.74995400	-1.78908800	4.69275900
H	-4.15064900	-0.70358900	4.89435000

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5053.62798843

$^3\text{P}_\text{A}^{TMHD}$



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.282662 (Hartree/Particle)

Thermal correction to Energy= 0.301298

Thermal correction to Enthalpy= 0.302242

Thermal correction to Gibbs Free Energy= 0.233217

Sum of electronic and zero-point Energies= -4662.566267

Sum of electronic and thermal Energies= -4662.547631

Sum of electronic and thermal Enthalpies= -4662.546686

Sum of electronic and thermal Free Energies= -4662.615712

C -4.16700400 -0.40222700 2.65529700

O -5.13971600 0.35433000 2.31766400

C -3.23345300 -0.95046200 1.75314300

H -2.45486600 -1.58098000 2.16264600

C -3.21113900 -0.71121000 0.36412700

O -4.06759300 0.00109200 -0.26275100

C -2.11244400 -1.30436300 -0.53872000

C -4.08893700 -0.67638000 4.16906700

Ni -5.62926000 0.77793600 0.52055600

Br -7.73773800 0.11643200 -0.24995500

C -2.90452500 -1.56399200 4.57828100

H -1.93833400 -1.10485800 4.31781100

H -2.91897000 -1.71112000 5.66947900

H -2.95499600 -2.55864500 4.10923100

C -3.97676100 0.68838700 4.88340500

H -4.81907900 1.34057700 4.61305300

H -3.98111300 0.54324000 5.97520400

H -3.03991500 1.20067100 4.61027400

C -5.41118300 -1.36744300 4.57247800

H -6.27548000 -0.74839600 4.29302600

H -5.51205200 -2.34592400 4.07557900

H -5.43309500 -1.53298600 5.66122100

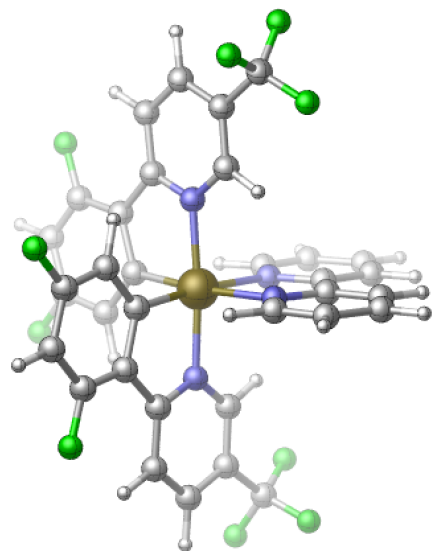
C -1.08285500 -2.15697000 0.21714600

H -0.54178200 -1.57110500 0.97611600

H -1.55059600 -3.02123300 0.71340400

H -0.33824900 -2.54663100 -0.49437100  
C -1.39251800 -0.12376400 -1.22725300  
H -0.63530700 -0.50315200 -1.93149900  
H -2.10740000 0.49898900 -1.78337600  
H -0.88157600 0.51258700 -0.48657700  
C -2.81648900 -2.17637200 -1.60240600  
H -3.33119000 -3.03092200 -1.13404800  
H -3.56097400 -1.58898700 -2.15814200  
H -2.07612800 -2.57259800 -2.31519100  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4664.03136026

**Figure S17**  
**[Ir(II)]**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.456088 (Hartree/Particle)

Thermal correction to Energy= 0.496055

Thermal correction to Enthalpy= 0.496999

Thermal correction to Gibbs Free Energy= 0.378449

Sum of electronic and zero-point Energies= -2626.184144

Sum of electronic and thermal Energies= -2626.144177

Sum of electronic and thermal Enthalpies= -2626.143232

Sum of electronic and thermal Free Energies= -2626.261783

Ir -0.38361500 0.77402400 0.05316900

C 2.34735200 1.55708000 1.07420000

C 2.41755700 1.29983400 -1.28265800

C 3.73792100 1.88540800 1.05493200

C 1.54618300 1.50257900 2.25738700

C 3.76397300 1.61776800 -1.34905400

H 1.85216300 1.05616000 -2.18579600

C 4.43596300 1.91595000 -0.13134000

H 4.24922400 2.11309300 1.99056100

C 2.02866400 1.78209300 3.57294900

H 4.27848300 1.63319400 -2.31040600

H 5.49928800 2.16794500 -0.13425200

C -0.59876000 1.08108500 3.17806200

C 1.18832500 1.70340600 4.66064600

H 3.07308700 2.06117100 3.71419800

C -0.17324600 1.33815200 4.47076900

H -1.63515300 0.80142300 2.97206200

H 1.56779800 1.92063400 5.66211800

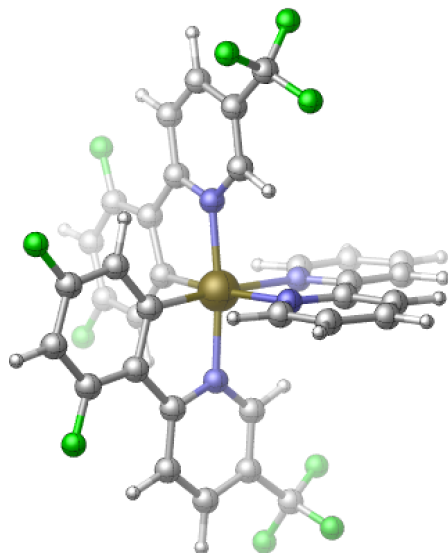
H -0.87109400 1.26044100 5.30511000

C -1.20298700 -1.96383800 0.73819500

C	1.11681300	-1.82859300	0.34511200
C	-1.07689600	-3.34293800	0.99669600
C	-2.43196900	-1.17336500	0.77340100
C	1.28877300	-3.18376500	0.59613000
H	1.95705500	-1.18870400	0.07846900
C	0.16627400	-3.95482400	0.92500200
H	-1.95872600	-3.92382500	1.24996100
C	-2.28624200	0.21283100	0.45110000
C	-3.71127200	-1.66010700	1.10138600
H	0.26512200	-5.02467200	1.11845200
C	-3.42294200	1.03566800	0.46870100
C	-4.83725100	-0.84913300	1.12405700
C	-4.66164900	0.49780900	0.80247100
H	-3.36520800	2.09860200	0.22890800
H	-5.81457800	-1.25438600	1.38393400
C	-0.76136600	0.59493300	-1.92704200
C	-1.08185900	1.82964800	-2.57472800
C	-0.70798800	-0.57879600	-2.69448400
C	-1.32492000	1.82253700	-3.96108600
C	-1.12033100	3.00700200	-1.70896600
C	-0.95672700	-0.52336500	-4.06206400
H	-0.47155300	-1.54613700	-2.24825800
C	-1.26781200	0.66511300	-4.72473000
C	-1.42632300	4.32796300	-2.09075400
H	-1.45934100	0.69105800	-5.79690900
C	-0.81789200	3.72327200	0.51586400
C	-1.42840100	5.34401800	-1.14574900
H	-1.66027100	4.54557000	-3.12862000
C	-1.11862300	5.03850900	0.18583900
H	-1.66255400	6.36897900	-1.43944600
N	1.71837900	1.25851900	-0.13231300
N	0.20508800	1.16057400	2.10042000
N	-0.08694600	-1.23930600	0.40954300
N	-0.82238600	2.74098200	-0.39775800
C	2.66020500	-3.80107100	0.55323200
F	3.14500300	-4.01512800	1.79241600
F	3.54522900	-3.02117500	-0.09014500
F	2.63996800	-4.99550400	-0.06605700
F	-3.88932800	-2.95793500	1.41527800
F	-5.73932900	1.29702900	0.81882300
F	-1.62428200	2.96511100	-4.60875900
F	-0.89844900	-1.65281300	-4.78414100
H	-0.56136900	3.43599300	1.53475600
C	-1.15339100	6.10205200	1.24945300
F	-0.61759400	7.25480300	0.80869300
F	-0.48167500	5.73342700	2.35350300

F -2.41670600 6.38074700 1.62781400  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2629.5151152  
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2627.9322792

**[Ir(III)]<sup>+</sup>**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.459873 (Hartree/Particle)

Thermal correction to Energy= 0.499499

Thermal correction to Enthalpy= 0.500443

Thermal correction to Gibbs Free Energy= 0.383450

Sum of electronic and zero-point Energies= -2626.063035

Sum of electronic and thermal Energies= -2626.023409

Sum of electronic and thermal Enthalpies= -2626.022465

Sum of electronic and thermal Free Energies= -2626.139458

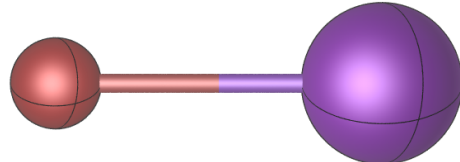
Ir	-0.36720300	0.77952700	0.06402700
C	2.38652400	1.58502300	1.06698800
C	2.42805500	1.35755700	-1.25950400
C	3.73997300	1.94563400	1.07249400
C	1.56470700	1.49006000	2.29981200
C	3.77401700	1.71107200	-1.32161800
H	1.86155100	1.11139000	-2.15945200
C	4.43993800	2.00930000	-0.13187700
H	4.24890400	2.17751000	2.00668500
C	2.07363700	1.73195000	3.58207600
H	4.28084200	1.74808300	-2.28671300
H	5.49488900	2.28997700	-0.13730600
C	-0.54779900	1.03770800	3.19369300
C	1.23267100	1.61942200	4.68865100
H	3.11804600	2.00575800	3.72187300
C	-0.10296800	1.26420200	4.49413000
H	-1.58236600	0.75935700	2.98495900
H	1.62051500	1.80683600	5.69171500
H	-0.79616300	1.16232700	5.33004100
C	-1.21972100	-1.95400300	0.73314100

C	1.10163800	-1.85646100	0.34063800
C	-1.11821500	-3.33733900	0.97548300
C	-2.43541000	-1.14432500	0.77199500
C	1.25013700	-3.21667800	0.57522400
H	1.95744300	-1.23755900	0.07567900
C	0.11416200	-3.97017800	0.89631400
H	-2.01018300	-3.90644400	1.21964200
C	-2.27236700	0.23887500	0.45318600
C	-3.72190900	-1.61132400	1.10260600
H	0.19325500	-5.04396300	1.07620300
C	-3.39069500	1.08218300	0.47576200
C	-4.83382600	-0.78149000	1.12989900
C	-4.63836000	0.56239400	0.81197100
H	-3.32047300	2.14392000	0.23766600
H	-5.81688700	-1.17171500	1.39127000
C	-0.74495800	0.58128800	-1.90904100
C	-1.07074200	1.80607900	-2.56857000
C	-0.67417000	-0.60085500	-2.65716800
C	-1.30553300	1.78075600	-3.95663300
C	-1.12303600	2.99278100	-1.71722400
C	-0.91559400	-0.56296500	-4.02840200
H	-0.43349300	-1.56155800	-2.20088300
C	-1.23351300	0.61441100	-4.70518000
C	-1.45072100	4.30320700	-2.11465900
H	-1.41910400	0.62615600	-5.77867700
C	-0.83773400	3.74010900	0.49917100
C	-1.47314600	5.32942900	-1.18093400
H	-1.68875200	4.50473300	-3.15477800
C	-1.16335400	5.04493300	0.15494100
H	-1.72632500	6.34619300	-1.48708800
N	1.75547400	1.29387500	-0.09938000
N	0.26119100	1.15194300	2.12847300
N	-0.09061500	-1.24482100	0.41597400
N	-0.81828700	2.74747400	-0.40373300
C	2.61267600	-3.85661200	0.52181200
F	3.10234800	-4.07120500	1.75753400
F	3.50139300	-3.08854900	-0.13119600
F	2.56742400	-5.04948500	-0.09414100
F	-3.91702700	-2.90403800	1.41404900
F	-5.69845700	1.37943900	0.83251100
F	-1.60903600	2.91184300	-4.61605200
F	-0.84204900	-1.69891000	-4.73266500
H	-0.58538500	3.47760800	1.52538800
C	-1.22539400	6.11997400	1.20806100
F	-0.68430200	7.26779300	0.76615200
F	-0.57444400	5.76211400	2.32835200

F -2.49680900 6.39308100 1.55489100  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2629.3946502  
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2627.8122578

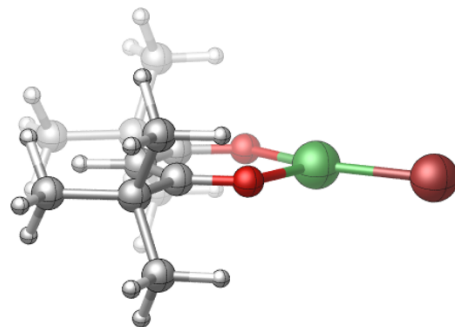
**K<sup>+</sup>**

UB3LYP-D3/def2-SVP-CPCM(THF)  
Zero-point correction= 0.000000 (Hartree/Particle)  
Thermal correction to Energy= 0.001416  
Thermal correction to Enthalpy= 0.002360  
Thermal correction to Gibbs Free Energy= -0.015176  
Sum of electronic and zero-point Energies= -599.792231  
Sum of electronic and thermal Energies= -599.790815  
Sum of electronic and thermal Enthalpies= -599.789871  
Sum of electronic and thermal Free Energies= -599.807407  
K 0.67977500 1.81953700 -2.01383600  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -599.8697705  
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -599.8424387

**KBr**

UB3LYP-D3/def2-SVP-CPCM(THF)  
Zero-point correction= 0.000291 (Hartree/Particle)  
Thermal correction to Energy= 0.003334  
Thermal correction to Enthalpy= 0.004279  
Thermal correction to Gibbs Free Energy= -0.024774  
Sum of electronic and zero-point Energies= -3173.856696  
Sum of electronic and thermal Energies= -3173.853653  
Sum of electronic and thermal Enthalpies= -3173.852709  
Sum of electronic and thermal Free Energies= -3173.881762  
K -0.27573900 1.80647700 -1.00469500  
Br -3.38333800 1.80647700 -1.00469500  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -3174.256792  
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -3174.1001508

**<sup>1</sup>P<sub>A</sub>-TMHD**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.282899 (Hartree/Particle)

Thermal correction to Energy= 0.301442

Thermal correction to Enthalpy= 0.302386

Thermal correction to Gibbs Free Energy= 0.234658

Sum of electronic and zero-point Energies= -4662.552097

Sum of electronic and thermal Energies= -4662.533554

Sum of electronic and thermal Enthalpies= -4662.532610

Sum of electronic and thermal Free Energies= -4662.600337

C	-2.67689100	1.50064700	-0.05133500
O	-1.42106200	1.53065300	-0.28586800
C	-3.43835400	0.32407000	0.07753400
H	-4.49775900	0.42432000	0.27535300
C	-2.90671800	-0.97398200	-0.03591800
O	-1.67346600	-1.22093200	-0.26675900
C	-3.30475800	2.90087100	0.08111300
C	-3.78209700	-2.23131300	0.10708500
Ni	-0.31332000	0.03984100	-0.51201000
Br	1.82020300	0.91198800	-0.90887100
C	-3.677444000	-3.01696800	-1.21927400
H	-2.63039000	-3.26130100	-1.44801600
H	-4.24888200	-3.95565800	-1.14550000
H	-4.08809500	-2.43169500	-2.05782800
C	-5.25691000	-1.91964900	0.40030800
H	-5.81737900	-2.86280300	0.49247700
H	-5.37928900	-1.36675700	1.34431100
H	-5.72151600	-1.33529900	-0.40862600
C	-3.19667000	-3.07570700	1.26065900
H	-3.25819300	-2.53293700	2.21771400
H	-3.76359900	-4.01435100	1.36297400
H	-2.14248000	-3.32164900	1.07013400
C	-3.04457300	3.64761900	-1.24638900
H	-3.42761500	4.67791500	-1.17842900
H	-1.96868200	3.68623800	-1.46782300
H	-3.55352200	3.14797900	-2.08647800

C -4.81319500 2.87253300 0.36555800  
H -5.37475200 2.38061600 -0.44351400  
H -5.04277600 2.35931100 1.31202700  
H -5.18707500 3.90483500 0.44886800  
C -2.57677900 3.62410700 1.23623200  
H -2.95320300 4.65492400 1.32904500  
H -2.74929600 3.10915900 2.19514100  
H -1.49362900 3.66017400 1.05302600

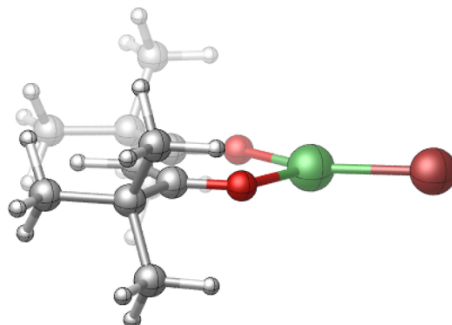
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4664.0202018

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4663.333533

**$^3P_A$ -TMHD**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.282628 (Hartree/Particle)

Thermal correction to Energy= 0.301243

Thermal correction to Enthalpy= 0.302187

Thermal correction to Gibbs Free Energy= 0.233454

Sum of electronic and zero-point Energies= -4662.565357

Sum of electronic and thermal Energies= -4662.546741

Sum of electronic and thermal Enthalpies= -4662.545797

Sum of electronic and thermal Free Energies= -4662.614530

C	-2.71245200	1.52103100	-0.04857400
O	-1.45837800	1.56881700	-0.27941600
C	-3.45883500	0.33159800	0.07913400
H	-4.52137600	0.42417300	0.26347600
C	-2.92327900	-0.96773600	-0.01162300
O	-1.68892300	-1.22719900	-0.21770400
C	-3.37150100	2.90790100	0.08917000
C	-3.80975600	-2.22005900	0.12992000
Ni	-0.28827500	0.05093900	-0.50691000
Br	1.92911300	0.60440700	-1.08862900
C	-3.68687100	-3.01752700	-1.18789800
H	-2.63661800	-3.26429600	-1.39857800
H	-4.26046700	-3.95521300	-1.11606200
H	-4.08376300	-2.43856900	-2.03759700
C	-5.28877600	-1.90374100	0.39590100
H	-5.85325700	-2.84488000	0.48603200
H	-5.42618500	-1.34293600	1.33320700
H	-5.73872800	-1.32467400	-0.42510700
C	-3.24759500	-3.05863700	1.29887700
H	-3.32562900	-2.50982300	2.25143700
H	-3.81724600	-3.99619600	1.39731400
H	-2.19040400	-3.30667500	1.12888900
C	-3.09963700	3.68145600	-1.21992800
H	-3.49950700	4.70484300	-1.14282400
H	-2.02075400	3.73993500	-1.42127400
H	-3.58547300	3.18839400	-2.07755400

C -4.88485300 2.85280500 0.34352000  
H -5.42231700 2.35841700 -0.48042200  
H -5.12521900 2.32757400 1.28069900  
H -5.27822900 3.87785000 0.42845200  
C -2.68038400 3.62616200 1.26988500  
H -3.07617000 4.64901400 1.37174600  
H -2.86289300 3.09311000 2.21711100  
H -1.59448700 3.68392200 1.10943900

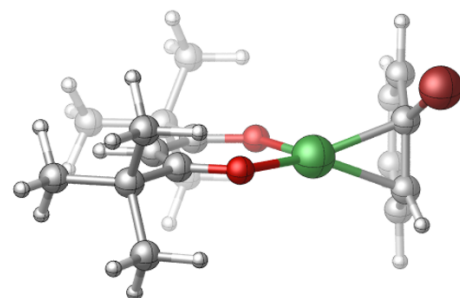
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4664.0306008

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4663.3442077

## <sup>2</sup>A-TMHD



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.371812 (Hartree/Particle)

Thermal correction to Energy= 0.396012

Thermal correction to Enthalpy= 0.396956

Thermal correction to Gibbs Free Energy= 0.314402

Sum of electronic and zero-point Energies= -4893.918513

Sum of electronic and thermal Energies= -4893.894313

Sum of electronic and thermal Enthalpies= -4893.893369

Sum of electronic and thermal Free Energies= -4893.975922

C -2.35056300 1.49763400 -0.08085100

O -1.10960500 1.45110100 -0.33763200

C -3.19367000 0.36601100 0.03381700

H -4.23921800 0.54154200 0.25423300

C -2.77418500 -0.97096700 -0.12013600

O -1.57666200 -1.32919700 -0.37629300

C -2.90157200 2.93116400 0.09863400

C -3.76813200 -2.14650600 0.00901700

Ni -0.07491500 -0.16653800 -0.64206800

C 1.55418400 2.72254700 -1.71182500

C 1.45782900 1.54447300 -2.43036700

C 1.62478300 0.28492900 -1.78974300

C 1.91005300 0.27106800 -0.39159800

C 2.04367800 1.49041300 0.32913700

C 1.84316700 2.69133100 -0.32679000

H 1.41396600 3.68253200 -2.21343400

H 1.26792300 1.56399800 -3.50600800

H 2.28974700 1.46223700 1.39153000

H 1.91409000 3.62603100 0.23439000

H 1.76342200 -0.61570400 -2.39502800

Br 2.65175100 -1.34156700 0.40594200

C -2.09139300 3.59678000 1.23197500

H -1.01591800 3.56642800 1.00633500

H -2.39813000 4.64815900 1.35425500

H -2.25721700 3.07951100 2.19139100

C -4.39753200 2.99040200 0.43959900

H -5.01698000 2.54991700 -0.35695600

H -4.62378400 2.46802800 1.38216500  
H -4.70817700 4.04070800 0.55863100  
C -2.64839900 3.68495300 -1.22544000  
H -3.21915200 3.23119400 -2.05236800  
H -2.96191300 4.73731100 -1.13208300  
H -1.58124000 3.65678700 -1.48825900  
C -3.24958800 -3.07040100 1.13205500  
H -3.88771200 -3.96486700 1.21565800  
H -2.21897900 -3.39134100 0.92429600  
H -3.25977900 -2.55155500 2.10463900  
C -5.20695800 -1.71727400 0.33082900  
H -5.26878700 -1.17815200 1.28885400  
H -5.62897000 -1.07314100 -0.45610100  
H -5.84886200 -2.60918200 0.40975400  
C -3.75272900 -2.90842500 -1.33445600  
H -2.73172900 -3.22922200 -1.58600900  
H -4.39793100 -3.79982300 -1.27598900  
H -4.12491600 -2.27066600 -2.15299400

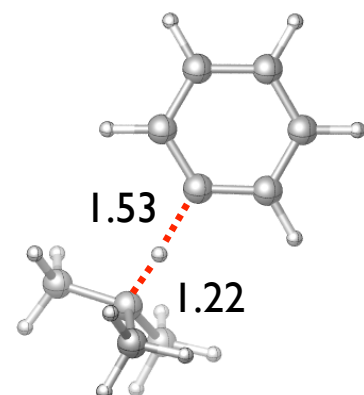
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4895.7207494

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4894.8531818

**Figure S18**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.213786 (Hartree/Particle)

Thermal correction to Energy= 0.225178

Thermal correction to Enthalpy= 0.226122

Thermal correction to Gibbs Free Energy= 0.173602

Sum of electronic and zero-point Energies= -389.540592

Sum of electronic and thermal Energies= -389.529200

Sum of electronic and thermal Enthalpies= -389.528256

Sum of electronic and thermal Free Energies= -389.580776

C -1.32452900 -5.70999100 -0.64601400

C -0.12003600 -5.86266600 -1.56424400

H 0.68500100 -6.43887600 -1.07969700

H -0.40152200 -6.39634600 -2.49114200

H 0.29018200 -4.88221700 -1.85654500

C -2.41255400 -4.81606700 -1.22439400

H -2.85986500 -5.27816400 -2.12420100

H -3.22412400 -4.64874100 -0.49758000

H -2.01124200 -3.83307100 -1.52055500

C -1.85409300 -7.03864200 -0.12381800

H -1.05791200 -7.62793500 0.35995400

H -2.66327700 -6.88962700 0.60966700

H -2.26260400 -7.64800700 -0.95155300

H -0.91115000 -5.12004400 0.34065500

C -0.40944900 -4.39769200 1.59793900

C -0.09322600 -3.04791900 1.51310700

C -0.29669600 -5.12910100 2.77318600

C 0.36202000 -2.39719300 2.67184400

H -0.19342200 -2.49804600 0.57204800

C 0.15955400 -4.46946500 3.92654200

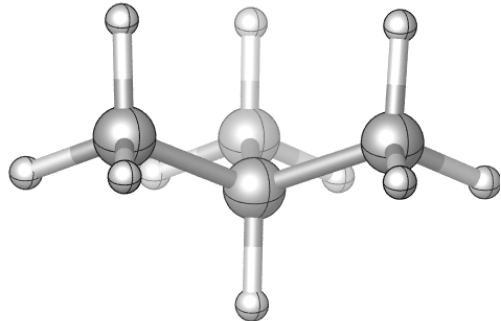
H -0.55527800 -6.19197700 2.80758700

C 0.48683500 -3.10845500 3.87230300

H 0.61887800 -1.33435500 2.63568700

H 0.25853600 -5.01968900 4.86691500

H 0.84136300 -2.59837600 4.77177200  
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -390.1859701  
UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -389.8284779



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.130747 (Hartree/Particle)

Thermal correction to Energy= 0.136352

Thermal correction to Enthalpy= 0.137297

Thermal correction to Gibbs Free Energy= 0.103241

Sum of electronic and zero-point Energies= -158.220710

Sum of electronic and thermal Energies= -158.215105

Sum of electronic and thermal Enthalpies= -158.214161

Sum of electronic and thermal Free Energies= -158.248216

C -1.35880900 -5.74058300 -0.69920900

C -0.82369700 -4.98911000 -1.92428400

H 0.27857000 -4.96727300 -1.93406700

H -1.15671900 -5.47685400 -2.85723600

H -1.18260900 -3.94683300 -1.94599900

C -2.89193400 -5.72268600 -0.66606700

H -3.30687200 -6.23708900 -1.55062800

H -3.28129200 -6.23329100 0.23015200

H -3.28288100 -4.69183400 -0.66506000

C -0.82205700 -7.17657400 -0.65350800

H 0.28028500 -7.19472100 -0.64411100

H -1.17672700 -7.71082400 0.24334200

H -1.15825600 -7.74716700 -1.53709700

H -0.99537000 -5.21550400 0.20414900

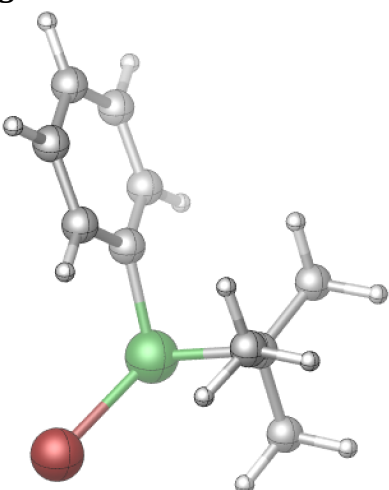
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -158.5324782

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -158.3793557

**Figure S19**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.211223 (Hartree/Particle)

Thermal correction to Energy= 0.225662

Thermal correction to Enthalpy= 0.226607

Thermal correction to Gibbs Free Energy= 0.167821

Sum of electronic and zero-point Energies= -4471.112669

Sum of electronic and thermal Energies= -4471.098230

Sum of electronic and thermal Enthalpies= -4471.097286

Sum of electronic and thermal Free Energies= -4471.156071

C 2.85880600 0.74317800 0.23337800

C 3.89346800 -0.36358800 0.23547100

C 2.89142500 1.59192300 -1.03876300

H 3.79736600 -1.00840200 -0.65008500

H 3.80865900 -0.98854000 1.13475300

H 2.83855000 0.98998300 -1.95855600

H 2.09760800 2.35745600 -1.06205300

Br 1.03762700 -1.52014500 -2.04620800

Ni 1.11076700 -0.01515500 -0.20188300

C 0.91749900 -0.80812000 1.49745800

C 0.23763100 -0.12580100 2.51524500

C 1.28575200 -2.14699300 1.68565300

C -0.07912200 -0.78179600 3.71312800

H -0.04145600 0.92453500 2.38833100

C 0.97451500 -2.79953400 2.88509200

H 1.80903300 -2.68372300 0.89031400

H -0.61299000 -0.24450900 4.50216000

H 1.26392600 -3.84500600 3.02485900

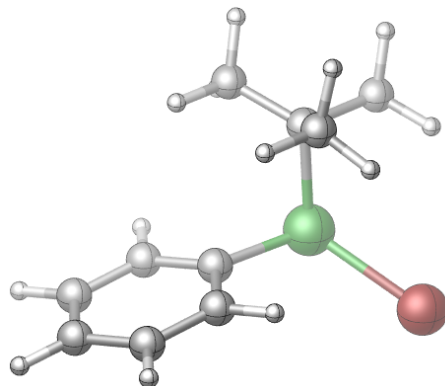
C 0.29169800 -2.11808000 3.90012700

H 0.05055000 -2.62804200 4.83649700

H 4.90621400 0.08337100 0.22516700

H 3.85381700 2.13871200 -1.06186100

C 2.86935900 1.59626600 1.48856300  
H 2.00503800 2.27775800 1.52880400  
H 2.86822500 0.98299600 2.39880000  
H 3.78220400 2.22164500 1.49347700  
UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4472.2797924  
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4471.7023257



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.211070 (Hartree/Particle)

Thermal correction to Energy= 0.225625

Thermal correction to Enthalpy= 0.226569

Thermal correction to Gibbs Free Energy= 0.166740

Sum of electronic and zero-point Energies= -4471.112594

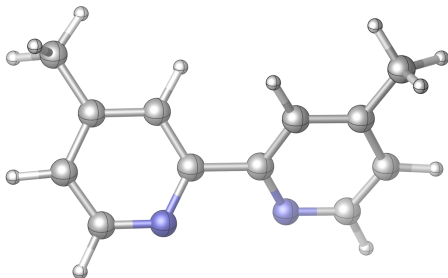
Sum of electronic and thermal Energies= -4471.098040

Sum of electronic and thermal Enthalpies= -4471.097096

Sum of electronic and thermal Free Energies= -4471.156924

Ni	-0.69015100	-0.10048000	0.06690000
C	-1.96369900	1.24956000	-0.26283500
C	-3.75785000	3.24547600	-1.08695500
C	-3.28369100	0.91615600	-0.59356900
C	-1.54374900	2.58408400	-0.34634400
C	-2.43860100	3.58003200	-0.76273900
C	-4.17889400	1.91239800	-1.00231600
H	-3.61298600	-0.12420000	-0.53461900
H	-0.52004100	2.86087800	-0.07631200
H	-2.10321300	4.61904300	-0.82766500
H	-5.20867000	1.64572900	-1.25680800
Br	-1.01456200	-2.11044000	-1.14130800
C	-1.03216700	-0.11144300	1.98866700
H	-4.45861200	4.02270400	-1.40299600
C	0.07143200	-1.13149100	2.27540100
H	0.11860800	-1.28593800	3.37058600
H	-0.12193100	-2.10897700	1.81056400
H	1.07300700	-0.78180000	1.97154100
C	-2.41490100	-0.64855700	2.29100800
H	-3.19423400	0.09351200	2.07311600
H	-2.62479500	-1.55971000	1.71081700
H	-2.47953600	-0.90882600	3.36492000
C	-0.73489600	1.24114900	2.61122300
H	0.21051300	1.66488300	2.23478800
H	-1.53801100	1.96418300	2.41945300
H	-0.62753400	1.12405900	3.70638000

UB3LYP-D3/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4472.279756  
UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -4471.7027401



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.212040 (Hartree/Particle)

Thermal correction to Energy= 0.224472

Thermal correction to Enthalpy= 0.225417

Thermal correction to Gibbs Free Energy= 0.170903

Sum of electronic and zero-point Energies= -573.414733

Sum of electronic and thermal Energies= -573.402300

Sum of electronic and thermal Enthalpies= -573.401356

Sum of electronic and thermal Free Energies= -573.455869

C -2.31575100 -1.15742200 -0.05825600

C -0.91697700 -1.27194600 0.02278700

C -0.22447200 -0.70695500 1.10164900

C -0.99218900 -0.04466300 2.07014800

C -2.37809600 0.02559400 1.90563400

C -3.07072300 -1.75985500 -1.20231200

C -2.63843200 -2.95842800 -1.79813600

C -3.35500200 -3.51359600 -2.86541400

C -4.49796800 -2.81983100 -3.28994600

C -4.85559900 -1.63801000 -2.63699400

H -0.36149000 -1.78401300 -0.76571600

H -0.51932900 0.41415800 2.94221800

H -2.98834200 0.54166500 2.65702300

H -1.75618600 -3.47459000 -1.41410900

H -5.10737900 -3.19713900 -4.11514400

H -5.74810300 -1.08816000 -2.96001800

N -3.03302700 -0.50956800 0.87394100

N -4.17073800 -1.11301100 -1.61872400

C -2.91853600 -4.78894400 -3.53673600

H -2.16322000 -5.32437400 -2.94346500

H -3.77370400 -5.46175800 -3.70579800

H -2.47910300 -4.57147800 -4.52509600

C 1.27668300 -0.77649400 1.19822700

H 1.60731200 -0.85611700 2.24488300

H 1.68075800 -1.63018200 0.63440200

H 1.72933200 0.14008200 0.78228300

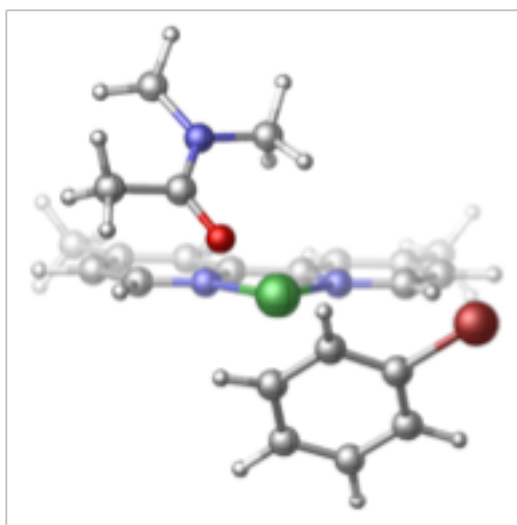
UB3LYP-D3/def2-TZVPP-CPCM(THF) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -574.2558479

UM06/def2-TZVPP-CPCM(THF)//UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -573.7901558

**<sup>1</sup>F-DMA-O**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.435353 (Hartree/Particle)

Thermal correction to Energy= 0.465364

Thermal correction to Enthalpy= 0.466308

Thermal correction to Gibbs Free Energy= 0.371774

Sum of electronic and zero-point Energies= -5174.442026

Sum of electronic and thermal Energies= -5174.412014

Sum of electronic and thermal Enthalpies= -5174.411070

Sum of electronic and thermal Free Energies= -5174.505604

C -1.99573800 1.70563600 1.27592200

O -1.57857800 0.71672600 0.64048900

N -2.89472400 2.56501500 0.75010400

C -1.47410900 1.95364900 2.67628200

H -1.11216300 2.98325600 2.81198000

H -2.26334600 1.76787100 3.42184500

H -0.65298600 1.24992700 2.85587800

C -3.33995100 2.40820300 -0.63035700

H -4.25568400 1.79961900 -0.69048700

H -3.54156900 3.40275500 -1.05441600

H -2.55717000 1.92011100 -1.22281600

C -3.58663900 3.59887300 1.50728500

H -4.67135200 3.39904700 1.50070200

H -3.25493700 3.63050100 2.54901400

H -3.41506200 4.58905900 1.05431700

Ni -2.83422400 -0.70330200 -0.47307400

C -2.12470800 -3.48520000 -0.24809200

C -1.46424900 -2.23640700 -0.06587200

C -0.92329800 -1.56582200 -1.20744300

C -1.06246300 -2.17899900 -2.47563400

C -1.73841200 -3.37384900 -2.65124100

C	-2.27633700	-4.02521900	-1.51351100
H	-2.51370100	-4.01365500	0.62537200
H	-1.14288900	-1.93603900	0.93594900
H	-1.85101900	-3.80954600	-3.64483900
H	-2.80536800	-4.97206300	-1.64706800
H	-0.24789300	-0.71828200	-1.08343100
Br	-0.30699000	-1.28484100	-3.99442100
C	-5.58488100	-0.71657100	-1.26670300
C	-6.73781400	-0.77640800	-2.10429200
C	-6.62620500	-0.92475200	-3.47360800
C	-5.31084700	-1.01892300	-4.02328900
C	-4.22414500	-0.96917600	-3.16702200
C	-5.62452000	-0.53277000	0.15671400
C	-6.81183900	-0.35572400	0.92827500
C	-6.75854700	-0.17742000	2.29830500
C	-5.47215700	-0.17843900	2.92135100
C	-4.35230300	-0.35121800	2.12428400
H	-7.72971900	-0.70607800	-1.65381300
H	-5.15633400	-1.12736700	-5.09888200
H	-3.20818600	-1.04370200	-3.56061600
H	-7.78206500	-0.35573300	0.42787800
H	-5.36804800	-0.05364900	4.00110500
H	-3.34962000	-0.36151400	2.56330400
N	-4.32004800	-0.82323300	-1.83343800
N	-4.39581800	-0.51921400	0.79480000
C	-7.99766800	0.01327800	3.13136200
H	-8.91250800	-0.01112300	2.52081600
H	-8.07819400	-0.77190300	3.90309800
H	-7.96779700	0.97738100	3.66821900
C	-7.82822500	-0.98815700	-4.37703000
H	-7.85017400	-1.93823500	-4.93878200
H	-8.77061600	-0.89980000	-3.81620200
H	-7.79867700	-0.18096600	-5.12951400

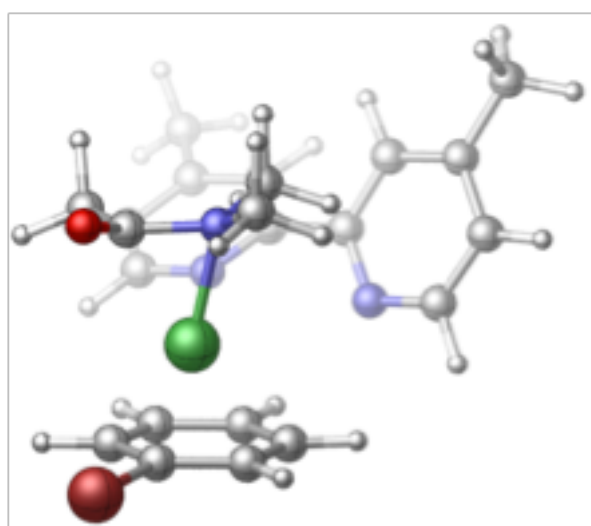
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5176.5999679

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5175.5019016

**<sup>1</sup>F-DMA'**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.434398 (Hartree/Particle)

Thermal correction to Energy= 0.465055

Thermal correction to Enthalpy= 0.465999

Thermal correction to Gibbs Free Energy= 0.367305

Sum of electronic and zero-point Energies= -5174.382824

Sum of electronic and thermal Energies= -5174.352168

Sum of electronic and thermal Enthalpies= -5174.351223

Sum of electronic and thermal Free Energies= -5174.449917

C -1.97016600 -0.98194000 -1.08653400

O -2.20401200 0.23211900 -0.94905100

N -0.60758500 -1.45789200 -1.14294800

C -3.05520500 -1.90925000 -1.62444700

H -3.12003200 -1.79774700 -2.72214000

H -2.91213600 -2.96911300 -1.38454000

H -4.00633000 -1.56981000 -1.19390600

C -0.25349500 -2.60168900 -1.99171600

H -0.17849100 -2.30293900 -3.05367600

H 0.71643900 -3.00280500 -1.66692700

H -0.99124600 -3.40058900 -1.88880200

C 0.42535300 -0.41680500 -1.13844900

H 0.58143400 -0.01349500 -2.15619400

H 0.11223800 0.39839900 -0.47949900

H 1.36587800 -0.85076800 -0.77147100

Ni -1.23286100 -1.97005100 0.65837000

C -1.08992900 -3.90407700 1.69927000

C -2.39999500 -3.36935600 1.70329300

C -2.63609200 -2.04654400 2.16848500

C -1.53656200 -1.26833800 2.61729100

C -0.22206000 -1.78959600 2.61961200

C	-0.01065000	-3.10841500	2.14799900
H	-0.90472400	-4.88789900	1.26762300
H	-3.23163500	-3.96770100	1.32849900
H	0.61175300	-1.18009900	2.96852300
H	1.00757500	-3.49857800	2.10094800
H	-3.64243600	-1.62574500	2.16820100
Br	-1.85461700	0.49234200	3.32735600
C	-0.09040200	-5.98907000	-1.79741600
C	0.72717800	-5.82898000	-2.92735000
C	2.08116800	-5.50137600	-2.77491200
C	2.55601200	-5.36665300	-1.46251900
C	1.67347400	-5.56036500	-0.39654700
C	-1.55072400	-6.27313800	-1.95183000
C	-2.01040200	-7.17779800	-2.92166800
C	-3.38412000	-7.41845300	-3.06355400
C	-4.24220200	-6.71770300	-2.20384900
C	-3.69645700	-5.83388600	-1.26960100
H	0.29925600	-5.92628100	-3.92752400
H	3.60125500	-5.11375200	-1.26868300
H	2.03505800	-5.46701800	0.63429100
H	-1.29511900	-7.71102700	-3.55197400
H	-5.32455600	-6.85832800	-2.25656300
H	-4.35677400	-5.27266200	-0.59793000
N	0.38261000	-5.86107300	-0.54841000
N	-2.38753300	-5.61150600	-1.13621600
C	-3.91142400	-8.36728900	-4.10664000
H	-3.18760700	-9.16524600	-4.32932600
H	-4.85947500	-8.82725000	-3.79075500
H	-4.10602100	-7.82680900	-5.04905000
C	2.96833300	-5.25888600	-3.96637600
H	4.01884800	-5.49575400	-3.74191900
H	2.64880300	-5.85264100	-4.83564100
H	2.92585600	-4.19588000	-4.26012500

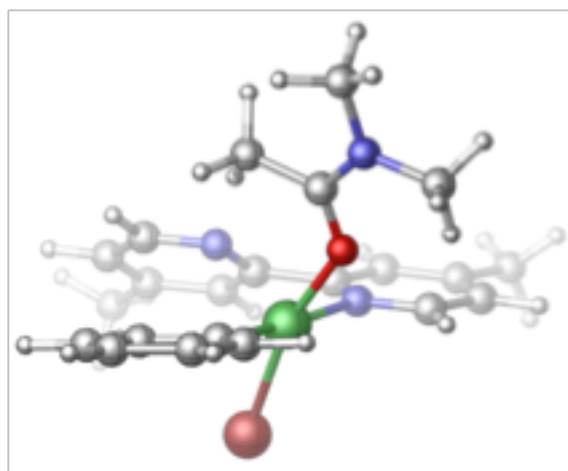
UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5176.5457794

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5175.463106

**<sup>1</sup>G-DMA-O**



UB3LYP-D3/def2-SVP-CPCM(THF)

Zero-point correction= 0.437417 (Hartree/Particle)

Thermal correction to Energy= 0.467940

Thermal correction to Enthalpy= 0.468884

Thermal correction to Gibbs Free Energy= 0.371580

Sum of electronic and zero-point Energies= -5174.486401

Sum of electronic and thermal Energies= -5174.455878

Sum of electronic and thermal Enthalpies= -5174.454934

Sum of electronic and thermal Free Energies= -5174.552238

C -2.51811700 0.31763100 0.07638600

O -1.75704900 -0.42918400 -0.58355500

N -2.03794100 0.99075400 1.13756200

C -3.96681200 0.45866400 -0.30611300

H -4.55773600 -0.27525200 0.26498400

H -4.36819500 1.46308700 -0.12181600

H -4.04949800 0.20330700 -1.37052500

C -2.87060900 1.84720400 1.97767100

H -2.39333500 1.94038300 2.96236100

H -2.97962400 2.85839700 1.55033500

H -3.86665800 1.41301600 2.13146500

C -0.62541000 0.90751900 1.49514000

H -0.19668200 1.92010700 1.56276800

H -0.50684800 0.41033900 2.47185100

H -0.08655700 0.33533700 0.73370900

Ni -2.17675500 -1.82835800 -1.86714500

C -3.43413900 -3.49580300 1.78010300

C -2.67291600 -2.89108000 0.76966300

C -3.15204600 -2.77163900 -0.54921900

C -4.43301900 -3.29451900 -0.81448900

C -5.20936200 -3.88521500 0.19066900

C -4.71214100 -3.99090300 1.49636400

H -3.02949300 -3.57586900 2.79444400

H	-1.68660800	-2.48703600	1.01852400
H	-4.82632300	-3.24467200	-1.83340900
H	-6.20606200	-4.27154400	-0.04668700
H	-5.31338900	-4.45711600	2.28225500
Br	-2.08778200	-3.71370700	-3.31942000
C	-3.16586000	-0.65969400	-4.64107200
C	-3.62524100	-0.99946500	-5.91967800
C	-4.96027400	-1.38708400	-6.10195100
C	-5.77973200	-1.40094300	-4.96412000
C	-5.23882800	-1.03497200	-3.72942700
C	-1.74267500	-0.29293700	-4.39998500
C	-0.99936100	0.43769100	-5.33616700
C	0.33428100	0.77675600	-5.07833500
C	0.86918600	0.36373200	-3.84796800
C	0.07257000	-0.35560600	-2.96397400
H	-2.93801600	-0.99718100	-6.76814700
H	-6.82987100	-1.69394000	-5.03522200
H	-5.86283400	-1.03861300	-2.82874600
H	-1.47541400	0.76390200	-6.26241400
H	1.90000700	0.60148200	-3.57663900
H	0.45598600	-0.68479900	-1.99576300
N	-3.96721800	-0.67229400	-3.56429500
N	-1.19861900	-0.68518600	-3.23111300
C	1.16437100	1.54228400	-6.07178600
H	0.54561600	1.98976200	-6.86246500
H	1.73798500	2.34099500	-5.57658600
H	1.89556400	0.87062000	-6.55264800
C	-5.47169300	-1.81243900	-7.45188100
H	-6.55881200	-1.66825800	-7.53590100
H	-4.97664600	-1.25818600	-8.26334200
H	-5.26605300	-2.88448100	-7.61430200

UB3LYP-D3/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

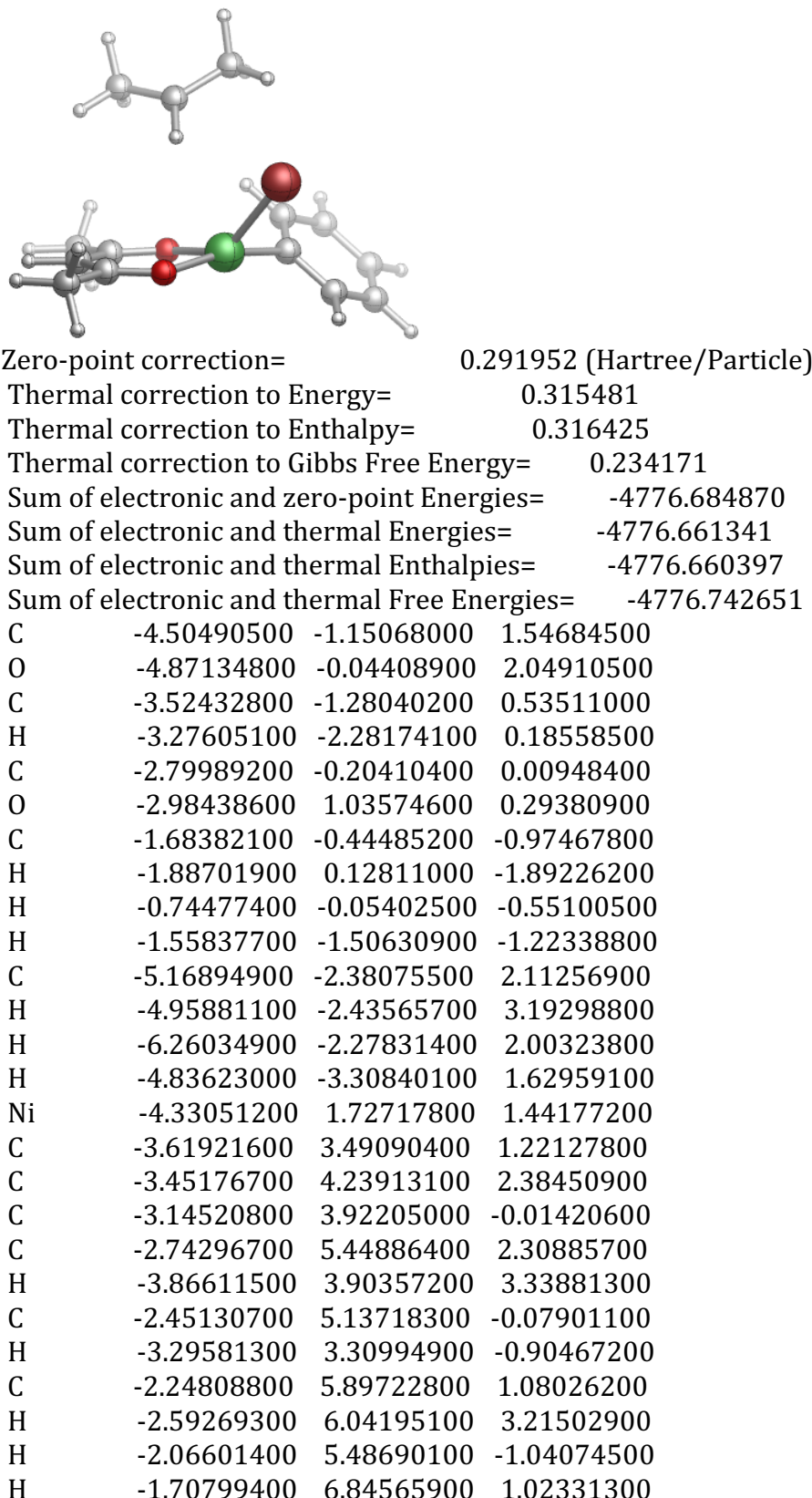
HF= -5176.6518013

UM06/def2-TZVPP-CPCM(THF)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5175.5492669

**Figure S21 and S22**

${}^3\text{C}$ -complex $i\text{-Pr}$



Br -6.44771500 2.65360700 1.23309800  
C -5.11552700 1.62544300 -2.28403600  
C -4.42971800 0.77253600 -3.29549900  
H -4.25594200 -0.25077100 -2.92809100  
H -5.00850200 0.70566000 -4.23717200  
H -3.44069100 1.18927200 -3.58548700  
C -5.60698200 2.98852600 -2.63087500  
H -6.09475100 3.47343300 -1.77207700  
H -4.78208400 3.65352000 -2.96502400  
H -6.32623700 2.96524500 -3.47394800  
H -5.13465700 1.30791000 -1.23963800

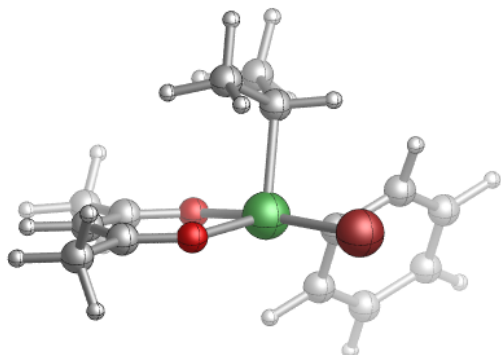
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF=-4817.61341725

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4816.80636293

**<sup>1</sup>C*i*-Pr**



Zero-point correction= 0.297520 (Hartree/Particle)  
Thermal correction to Energy= 0.318715  
Thermal correction to Enthalpy= 0.319659  
Thermal correction to Gibbs Free Energy= 0.246597  
Sum of electronic and zero-point Energies= -4776.718715  
Sum of electronic and thermal Energies= -4776.697520  
Sum of electronic and thermal Enthalpies= -4776.696576  
Sum of electronic and thermal Free Energies= -4776.769638  
C -4.26273100 -0.32975800 0.88325900  
O -5.48402200 0.00752800 0.68169600  
C -3.38034400 -0.82945700 -0.07996300  
H -2.36837900 -1.08522300 0.23092600  
C -3.76492900 -1.08155300 -1.41643700  
O -4.90428300 -0.79971000 -1.89114600  
C -2.79089900 -1.72159500 -2.37446300  
H -3.24196500 -2.63947000 -2.78393400  
H -2.62784600 -1.03858900 -3.22370400  
H -1.82661000 -1.96303700 -1.90881300  
C -3.81451800 -0.15170900 2.31204900  
H -2.78277800 -0.48724800 2.47756500  
H -3.90213400 0.91250200 2.58507400  
H -4.49525900 -0.70914500 2.97462600  
Ni -6.41932000 0.06169200 -0.95042200  
C -8.05687900 0.29108000 0.01760800  
C -9.04317800 1.26600900 -0.15218700  
C -8.23551000 -0.72293200 0.96570000  
C -10.19712900 1.24092600 0.64018600  
H -8.93270500 2.04653300 -0.90690000  
C -9.40114700 -0.75480000 1.74569400  
H -7.47139600 -1.48860900 1.11625400  
C -10.38063200 0.22978700 1.59065800  
H -10.95983800 2.01277900 0.50390800  
H -9.53226000 -1.55152300 2.48338700  
H -11.28413500 0.20987500 2.20541700  
Br -7.66733300 -0.10032400 -2.93012200

C -6.09728600 2.06783100 -1.09823400  
C -4.86421300 2.12587800 -1.95693100  
H -3.98270700 1.71952500 -1.43941700  
H -4.99714800 1.61084700 -2.91533100  
H -4.66010000 3.19576300 -2.15927600  
H -7.00456200 2.34137600 -1.64664100  
C -5.98780400 2.75420900 0.23531300  
H -6.84282300 2.54899700 0.88981400  
H -5.06152300 2.49389800 0.76277600  
H -5.97146100 3.84284700 0.03369600

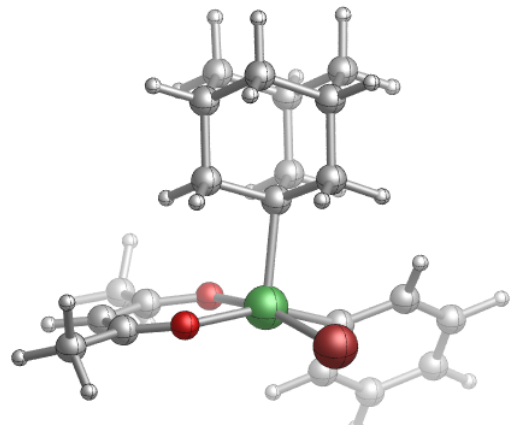
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4778.31828595

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4777.53911766

**<sup>1</sup>C'Ad**



Zero-point correction= 0.437415 (Hartree/Particle)

Thermal correction to Energy= 0.461985

Thermal correction to Enthalpy= 0.462930

Thermal correction to Gibbs Free Energy= 0.382855

Sum of electronic and zero-point Energies= -5047.993260

Sum of electronic and thermal Energies= -5047.968689

Sum of electronic and thermal Enthalpies= -5047.967745

Sum of electronic and thermal Free Energies= -5048.047820

C -4.43468200 -0.23717600 0.96563600

O -5.63063900 0.17719300 0.76541200

C -3.53913800 -0.66722700 -0.01969400

H -2.54843700 -0.99537800 0.29203500

C -3.86018300 -0.69469700 -1.39633600

O -4.97313600 -0.33049400 -1.87551300

C -2.82915900 -1.15406300 -2.39894800

H -3.28876100 -1.88280400 -3.08401900

H -2.52215300 -0.28777900 -3.00863400

H -1.93913200 -1.59320500 -1.92938900

C -4.01701000 -0.21032800 2.41514700

H -4.75884300 -0.75938900 3.01585500

H -3.01925100 -0.63829200 2.57712800

H -4.02588200 0.83402600 2.76813400

Ni -6.58537800 0.27486400 -0.87403600

C -8.23065100 0.18830000 0.11302300

C -9.38838100 0.95737400 -0.03924200

C -8.23130200 -0.87031500 1.03189300

C -10.51933900 0.70313800 0.74559700

H -9.42878500 1.75477900 -0.78071100

C -9.37136000 -1.13661200 1.80513200

H -7.34523200 -1.49331100 1.16546000

C -10.51541800 -0.34591600 1.67233500

H -11.41160100 1.32293000 0.61937100

H -9.35412900 -1.96526100 2.51897100

H	-11.40032900	-0.54739500	2.28160600
Br	-7.82631400	-0.08224600	-2.83680800
C	-4.86888900	2.49814000	-1.18010100
H	-4.66416300	2.08205600	-2.17429500
H	-4.24168300	1.97428200	-0.44909800
C	-6.34184500	2.39288000	-0.84527500
C	-4.50641700	4.01939000	-1.16399200
H	-3.43319900	4.10577900	-1.40007500
C	-5.34808000	4.75003500	-2.22143000
H	-5.13013300	4.34938300	-3.22530000
H	-5.08808900	5.82190700	-2.23530800
C	-7.18702000	3.04934700	-1.91445700
H	-8.25940200	2.91950200	-1.72799600
H	-6.96178200	2.63617100	-2.90423200
C	-6.83901300	4.57486400	-1.89206200
H	-7.46108900	5.06625700	-2.65779700
C	-7.14221400	5.15760600	-0.50287900
H	-6.91855300	6.23784100	-0.49205000
H	-8.21399500	5.04675800	-0.26777500
C	-6.65936500	2.91466300	0.54025900
H	-6.08277400	2.37795100	1.30202100
H	-7.72461700	2.79909500	0.77294500
C	-4.79585900	4.59391900	0.23195600
H	-4.18630300	4.07566800	0.99119800
H	-4.51625400	5.66051200	0.26485800
C	-6.29075800	4.43118800	0.55111100
H	-6.51108900	4.82321400	1.55745700

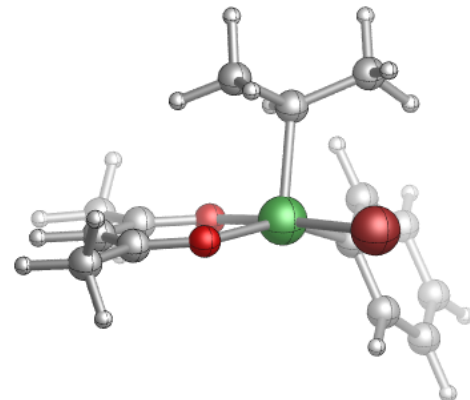
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5050.01092587

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5049.00795149

**<sup>1</sup>C''-complex<sup>i</sup>-Pr**



Zero-point correction= 0.297586 (Hartree/Particle)

Thermal correction to Energy= 0.318784

Thermal correction to Enthalpy= 0.319728

Thermal correction to Gibbs Free Energy= 0.246639

Sum of electronic and zero-point Energies= -4776.714077

Sum of electronic and thermal Energies= -4776.692879

Sum of electronic and thermal Enthalpies= -4776.691935

Sum of electronic and thermal Free Energies= -4776.765024

C	-4.71394900	-0.83682200	1.41221400
O	-4.96334000	0.31835300	1.91336500
C	-3.79101400	-1.11077400	0.39737800
H	-3.69214900	-2.14071900	0.05733900
C	-3.01247000	-0.11220400	-0.23128300
O	-3.01644500	1.10855800	0.10530200
C	-2.09747300	-0.47872000	-1.37378800
H	-2.11986300	-1.54930700	-1.61525800
H	-2.38705800	0.10316800	-2.26352600
H	-1.06870200	-0.17986100	-1.11674100
C	-5.53236100	-1.94915000	2.01885800
H	-5.31999400	-2.00026600	3.09889300
H	-6.60215600	-1.70916400	1.91294500
H	-5.32724400	-2.92455600	1.55948600
Ni	-4.09501000	1.95354900	1.54168100
C	-3.04178000	1.74801500	3.28615000
C	-2.78031600	3.00241500	4.07332000
H	-2.44997000	2.68019700	5.08024400
H	-1.98767900	3.61381600	3.62531200
H	-3.67732100	3.61969300	4.20166700
C	-1.81892600	1.00651300	2.82222100
H	-1.22553000	0.76413800	3.72576300
H	-2.05787700	0.05950600	2.32267400
H	-1.19275500	1.62443600	2.16442000
C	-5.60201600	2.80182900	2.36851200

C	-6.08675100	2.53451400	3.65025400
C	-6.37717500	3.55231700	1.47551300
C	-7.33172500	3.03849900	4.05039900
H	-5.51981400	1.92294500	4.35362900
C	-7.62931900	4.04192000	1.87471000
H	-6.01458000	3.77186700	0.46955100
C	-8.10706200	3.79351300	3.16459400
H	-7.69723100	2.82679200	5.05920700
H	-8.22491700	4.62800800	1.16911600
H	-9.07898300	4.18346400	3.47749200
Br	-3.10094400	4.00210200	0.96810200
H	-3.77534000	1.09081800	3.76577800

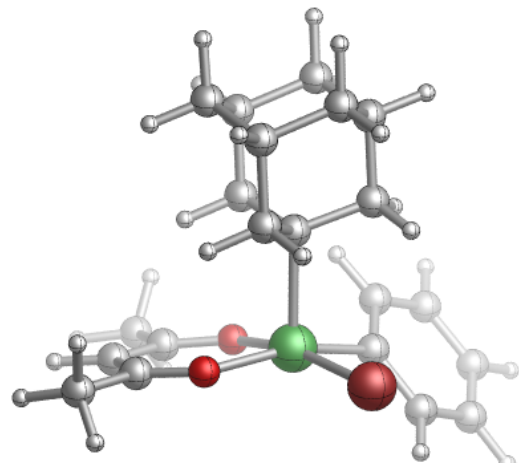
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4778.31370771

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4777.53485701

**<sup>1</sup>C''-complex<sup>Ad</sup>**



Zero-point correction= 0.437465 (Hartree/Particle)

Thermal correction to Energy= 0.462034

Thermal correction to Enthalpy= 0.462978

Thermal correction to Gibbs Free Energy= 0.382950

Sum of electronic and zero-point Energies= -5047.990295

Sum of electronic and thermal Energies= -5047.965725

Sum of electronic and thermal Enthalpies= -5047.964781

Sum of electronic and thermal Free Energies= -5048.044810

C -5.00425600 -0.26788700 0.89142900

O -6.20763300 -0.09658300 0.49061000

C -3.92294100 -0.63981800 0.08489000

H -2.94211000 -0.74516000 0.54650700

C -4.05834100 -0.91786900 -1.29336000

O -5.14597900 -0.82709000 -1.93726700

C -2.85361600 -1.33961400 -2.09877100

H -3.09933600 -2.24865600 -2.66908500

H -2.62663900 -0.55028500 -2.83475200

H -1.96501300 -1.51818300 -1.47930700

C -4.80793300 -0.01038900 2.36445900

H -3.78175400 -0.21092500 2.69861400

H -5.06313200 1.04066800 2.57685000

H -5.51238900 -0.63330200 2.93768500

Ni -6.90807900 -0.25022800 -1.26341200

C -8.63211600 -0.13074400 -0.38893200

C -8.86934400 0.84796400 0.58480500

C -9.56877900 -1.16303200 -0.53966600

C -10.03786500 0.83357700 1.35447300

H -8.13073900 1.62753500 0.77055200

C -10.73242100 -1.19106000 0.24524300

H -9.40861800 -1.95228000 -1.27397800

C -10.97907400 -0.18855300 1.18566700

H -10.20362900 1.61892600 2.09767500

H	-11.44968000	-2.00568000	0.10819400
H	-11.89158500	-0.20537700	1.78748400
Br	-7.87407100	-0.91544200	-3.32141600
C	-6.95241000	4.63241400	-1.45962800
H	-6.77787700	5.68900000	-1.72357900
H	-7.57063200	4.62556600	-0.54643000
C	-5.60910400	3.93191200	-1.19863500
H	-5.07774200	4.41776800	-0.36418300
C	-7.68932700	3.91934400	-2.60538300
H	-8.66405700	4.39922800	-2.79056300
C	-6.83291200	3.93696800	-3.87980400
H	-6.65234000	4.97726200	-4.19944100
H	-7.36516400	3.43248800	-4.70306600
C	-4.74589100	3.95079400	-2.46905000
H	-3.77825900	3.45721400	-2.27934300
H	-4.52747500	4.99215200	-2.75978100
C	-5.49624600	3.23318400	-3.60106100
H	-4.87864000	3.20487900	-4.51356400
C	-5.77110800	1.75099000	-3.18032800
H	-6.28957300	1.22636700	-3.98975300
H	-4.81947800	1.24232800	-2.99613300
C	-5.87600300	2.44612500	-0.78263100
H	-6.45320600	2.42802400	0.14897700
H	-4.91587500	1.94283500	-0.60564500
C	-7.96891600	2.43660100	-2.18948600
H	-8.60512100	2.43346300	-1.29925300
H	-8.49574500	1.90864900	-2.99410700
C	-6.62208300	1.80550400	-1.92723900

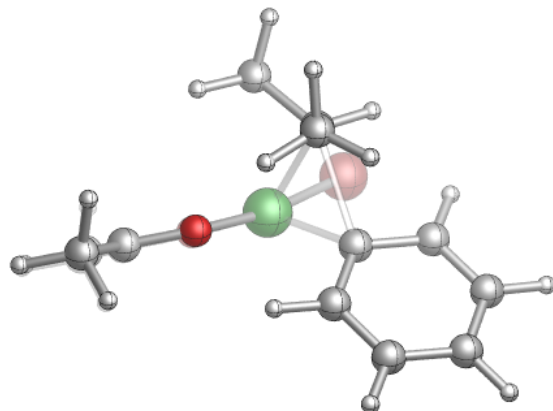
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5050.01092587

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5049.00795149

**<sup>1</sup>C'-TS<sup>i</sup>-Pr**



Zero-point correction= 0.297700 (Hartree/Particle)  
Thermal correction to Energy= 0.318234  
Thermal correction to Enthalpy= 0.319178  
Thermal correction to Gibbs Free Energy= 0.247594  
Sum of electronic and zero-point Energies= -4776.703604  
Sum of electronic and thermal Energies= -4776.683070  
Sum of electronic and thermal Enthalpies= -4776.682126  
Sum of electronic and thermal Free Energies= -4776.753710

C	-3.98117100	-0.38342000	0.70411400
O	-5.08969800	0.21721900	0.49156100
C	-3.32399300	-1.20475000	-0.21926400
H	-2.38337900	-1.67194700	0.06838000
C	-3.85483700	-1.46525500	-1.49621700
O	-4.93772100	-0.97201700	-1.94555200
C	-3.13566300	-2.39152600	-2.44479200
H	-3.80094700	-3.23338200	-2.69534700
H	-2.93190400	-1.85372300	-3.38431600
H	-2.19593600	-2.77930000	-2.03101300
C	-3.39798000	-0.14585800	2.07400300
H	-2.44765700	-0.67334000	2.22592400
H	-3.24443600	0.93573600	2.21689000
H	-4.12400600	-0.47172600	2.83614500
Ni	-6.13948800	0.26022100	-1.10166300
C	-7.81306000	0.66666600	-0.10279400
C	-9.05265800	1.00209600	-0.67045700
C	-7.79056700	0.05730400	1.16296900
C	-10.24405400	0.69568300	-0.00790000
H	-9.09530700	1.49703500	-1.63995100
C	-8.98807100	-0.25809500	1.81696400
H	-6.84295300	-0.16692600	1.64708500
C	-10.21906500	0.05747300	1.23709300
H	-11.19717400	0.96200700	-0.47238400
H	-8.94779400	-0.74867400	2.79324000
H	-11.15168100	-0.18170300	1.75423300

Br -7.17326300 0.42517600 -3.27868800  
C -6.51392300 2.34400900 -0.60210600  
C -5.16174000 2.60170600 -1.26675000  
H -4.31501000 2.28670100 -0.64282500  
H -5.06437200 2.17265500 -2.27356000  
H -5.09625500 3.69978600 -1.38743300  
H -7.31815200 2.70006500 -1.25050200  
C -6.56569800 2.90752400 0.80142100  
H -7.58235300 2.93153300 1.21220100  
H -5.91979500 2.32549000 1.47258200  
H -6.17602500 3.94075700 0.77039600

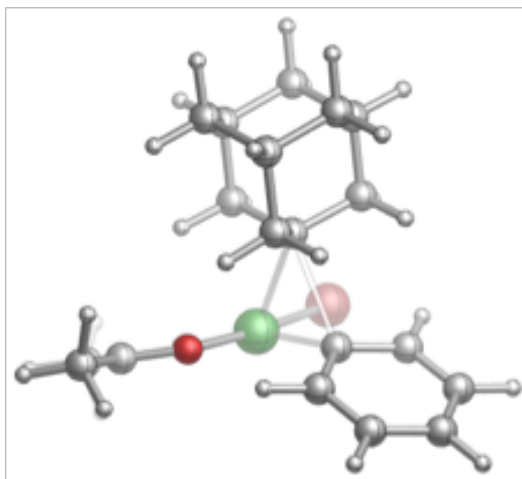
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4778.30356914

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4777.52976312

**<sup>1</sup>C'-TS<sup>Ad</sup>**



Zero-point correction= 0.437403 (Hartree/Particle)

Thermal correction to Energy= 0.461446

Thermal correction to Enthalpy= 0.462390

Thermal correction to Gibbs Free Energy= 0.382978

Sum of electronic and zero-point Energies= -5047.979137

Sum of electronic and thermal Energies= -5047.955094

Sum of electronic and thermal Enthalpies= -5047.954150

Sum of electronic and thermal Free Energies= -5048.033562

C -4.15139200 -0.32333600 0.86761600

O -5.33202900 0.08503200 0.60028300

C -3.28524600 -0.92518300 -0.05312300

H -2.29807700 -1.24196500 0.28039100

C -3.67414100 -1.18129800 -1.38283800

O -4.79198000 -0.84913700 -1.88498500

C -2.74210200 -1.91202600 -2.31851000

H -3.20762500 -2.86806000 -2.60894000

H -2.62121000 -1.32162500 -3.24013500

H -1.75870800 -2.10984600 -1.87251700

C -3.72030600 -0.10058300 2.29585600

H -4.45272300 -0.56802400 2.97288600

H -2.71979700 -0.50074500 2.50485600

H -3.73066700 0.98136300 2.50662900

Ni -6.24602900 0.14729400 -1.08859200

C -7.98762200 0.57629700 -0.17833400

C -9.20253900 0.66459100 -0.87981900

C -8.03340200 0.19172400 1.17451300

C -10.41614200 0.34639100 -0.26192300

H -9.20741400 0.97120500 -1.92361800

C -9.24668200 -0.12980900 1.79215800

H -7.11224400 0.13968000 1.75013500

C -10.44603100 -0.05293300 1.07805200

H -11.34361400 0.41580200 -0.83693300

H	-9.24851800	-0.43760300	2.84154300
H	-11.39547300	-0.29623200	1.56217100
Br	-7.11843200	0.19947500	-3.36062400
C	-5.28375200	2.52892500	-1.24878000
H	-5.30605900	2.19079400	-2.29364600
H	-4.52331400	1.95547200	-0.70399300
C	-6.66621600	2.43410400	-0.59177100
C	-4.85052900	4.03020500	-1.22245600
H	-3.85529700	4.08862300	-1.69206100
C	-5.86741000	4.86152700	-2.01666000
H	-5.89981800	4.52240400	-3.06533100
H	-5.56363200	5.92190700	-2.02670500
C	-7.69195500	3.21864500	-1.38395500
H	-8.68517300	3.14425400	-0.92525600
H	-7.75299900	2.84885700	-2.41591000
C	-7.25041000	4.71496200	-1.36622500
H	-8.00136400	5.28767600	-1.93433800
C	-7.19782800	5.20698000	0.09103600
H	-6.91456200	6.27252200	0.11987200
H	-8.19384400	5.12403000	0.55743900
C	-6.61585500	2.87362600	0.85785500
H	-5.90737800	2.25638300	1.42358300
H	-7.60348100	2.79088600	1.32881700
C	-4.78425100	4.50871200	0.23563300
H	-4.04311600	3.91510700	0.79703700
H	-4.45395100	5.56043200	0.27309400
C	-6.17330500	4.36820400	0.87545500
H	-6.14156600	4.69343500	1.92815500

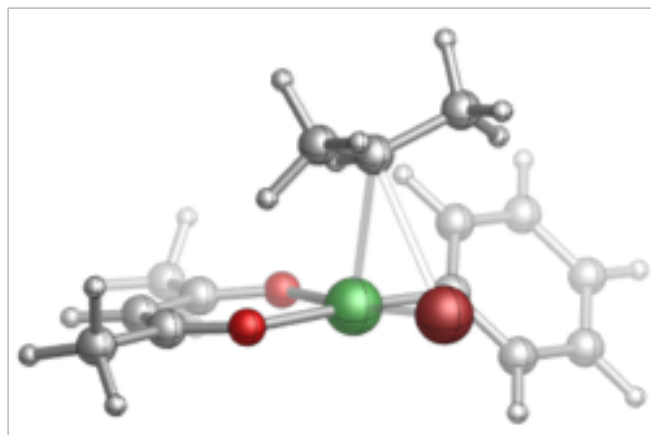
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5049.99792520

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -5048.99950709

**<sup>1</sup>C''-TS<sup>i</sup>-Pr**



Zero-point correction= 0.295724 (Hartree/Particle)

Thermal correction to Energy= 0.316798

Thermal correction to Enthalpy= 0.317742

Thermal correction to Gibbs Free Energy= 0.244376

Sum of electronic and zero-point Energies= -4776.696349

Sum of electronic and thermal Energies= -4776.675275

Sum of electronic and thermal Enthalpies= -4776.674331

Sum of electronic and thermal Free Energies= -4776.747697

C	-5.04608300	-1.01406600	1.21106900
O	-5.32751400	0.16285200	1.62034400
C	-3.95817400	-1.36070000	0.39392100
H	-3.83217900	-2.40749900	0.11902000
C	-3.01109300	-0.42613100	-0.07225700
O	-3.04483700	0.82052700	0.17074100
C	-1.83955400	-0.89548300	-0.90294300
H	-1.92024200	-1.94887900	-1.20229400
H	-1.75036900	-0.26041400	-1.79788800
H	-0.91319000	-0.76089700	-0.31938400
C	-6.00259300	-2.08229600	1.68641000
H	-6.03083400	-2.07627200	2.78789300
H	-7.01889000	-1.83443700	1.33978700
H	-5.73264600	-3.08619400	1.33347400
Ni	-4.41067800	1.79213500	1.23167700
C	-2.57075800	2.20308600	3.02216800
C	-2.82417200	3.25030200	4.03568000
H	-2.53735600	2.79351300	5.00681500
H	-2.21473800	4.14976000	3.88032600
H	-3.88908900	3.50542500	4.10354200
C	-1.20542500	1.90133300	2.53608000
H	-0.75543000	1.23275600	3.29992000
H	-1.23654200	1.34670300	1.58995900
H	-0.58140300	2.79910800	2.43890900
C	-5.79914400	2.69344700	2.16757700

C	-6.17978900	2.25492700	3.44617500
C	-6.52459700	3.74546800	1.58528700
C	-7.23572400	2.86391200	4.13724900
H	-5.65589900	1.41241200	3.90724800
C	-7.59492600	4.34545600	2.26323800
H	-6.24598500	4.11342900	0.59465800
C	-7.94995400	3.91227500	3.54554300
H	-7.50686200	2.51117100	5.13705900
H	-8.14794900	5.16173000	1.78842100
H	-8.77793000	4.38672600	4.07943700
Br	-3.19307500	3.77569500	0.91731000
H	-3.24161700	1.33852300	3.07332400

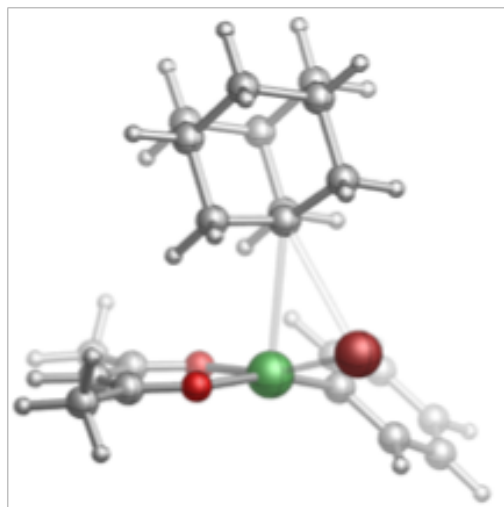
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4778.29784834

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4777.52344646

**<sup>1</sup>C''-TS<sup>Ad</sup>**



Zero-point correction= 0.435959 (Hartree/Particle)

Thermal correction to Energy= 0.460362

Thermal correction to Enthalpy= 0.461306

Thermal correction to Gibbs Free Energy= 0.380306

Sum of electronic and zero-point Energies= -5047.974046

Sum of electronic and thermal Energies= -5047.949643

Sum of electronic and thermal Enthalpies= -5047.948699

Sum of electronic and thermal Free Energies= -5048.029699

C -4.70640500 -0.24338900 0.58006500

O -5.97458200 -0.12831900 0.47699400

C -3.85228100 -0.76873500 -0.40277800

H -2.78784300 -0.83737700 -0.18002100

C -4.31201900 -1.22967700 -1.65667100

O -5.51710500 -1.16812900 -2.04300200

C -3.33209400 -1.81335900 -2.64797000

H -3.68092300 -2.81241100 -2.95423400

H -3.32458100 -1.18588500 -3.55440000

H -2.31077800 -1.88798900 -2.25150300

C -4.13796000 0.25448500 1.88799800

H -3.05636400 0.08748700 1.97593300

H -4.34547400 1.33343400 1.97952500

H -4.65733200 -0.24303300 2.72225700

Ni -7.06881100 -0.54722400 -1.01368500

C -8.60538600 -0.38655400 0.10336900

C -8.68547700 0.51094000 1.18279500

C -9.66517000 -1.30313800 -0.04643800

C -9.78274500 0.52056400 2.05456800

H -7.86177200 1.20203100 1.36953200

C -10.75773200 -1.31420800 0.83143800

H -9.64558000 -2.01968100 -0.87083800

C -10.82770600 -0.39436100 1.88335900

H	-9.81496200	1.24065900	2.87856800
H	-11.56155900	-2.04311400	0.68677900
H	-11.68402100	-0.39280000	2.56374300
Br	-8.31327900	-0.51928400	-3.01682200
C	-7.41274300	4.85063700	-2.19040200
H	-7.04767900	5.88730600	-2.27380500
H	-8.47811100	4.90440700	-1.91455800
C	-6.61749100	4.10709200	-1.10810600
H	-6.75092400	4.58013100	-0.12309100
C	-7.24506500	4.12846500	-3.53562100
H	-7.83647000	4.62051700	-4.32301200
C	-5.76139800	4.05320100	-3.93024100
H	-5.35591700	5.07032300	-4.05623700
H	-5.64836100	3.53864300	-4.89788200
C	-5.12920000	4.03202000	-1.48564400
H	-4.56189000	3.50136600	-0.70414600
H	-4.71189100	5.04971300	-1.55445600
C	-4.97413900	3.31588400	-2.83695500
H	-3.91267100	3.21821500	-3.11167500
C	-5.53837000	1.83511100	-2.68920300
H	-5.43283400	1.29008600	-3.63465600
H	-4.98455500	1.32238000	-1.89644300
C	-7.17747600	2.62440500	-0.97500200
H	-8.23936600	2.64724600	-0.70277200
H	-6.60664700	2.09380900	-0.20610700
C	-7.80033700	2.65138900	-3.39786300
H	-8.86019700	2.67321400	-3.11572600
H	-7.68131800	2.12275100	-4.35041500
C	-6.94726400	2.08232300	-2.32846000

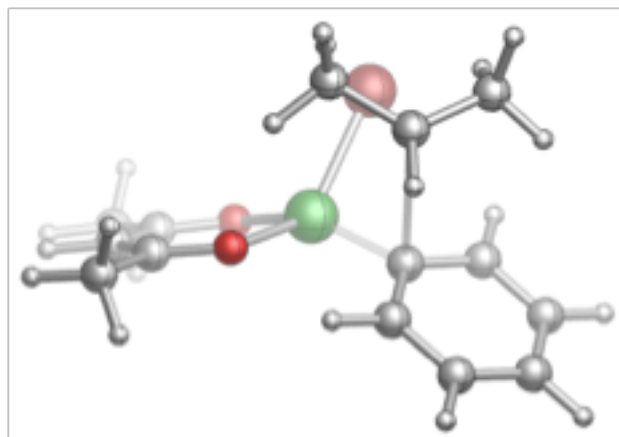
UB3LYP-D3/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -5050.00227554

UM06/def2-TZVPP-CPCM(DMA) // UB3LYP-D3/def2-SVP-CPCM(THF)

HF = -5049.00355154

<sup>3</sup>C-TS<sup>i</sup>-Pr



Zero-point correction= 0.291168 (Hartree/Particle)  
Thermal correction to Energy= 0.314140  
Thermal correction to Enthalpy= 0.315084  
Thermal correction to Gibbs Free Energy= 0.234321  
Sum of electronic and zero-point Energies= -4776.683307  
Sum of electronic and thermal Energies= -4776.660335  
Sum of electronic and thermal Enthalpies= -4776.659391  
Sum of electronic and thermal Free Energies= -4776.740155

C	-2.36118600	-0.64377600	0.45185900
O	-3.12407300	0.38453200	0.41725900
C	-2.78580900	-1.98098800	0.41828900
H	-2.02349700	-2.75834000	0.45386000
C	-4.14152700	-2.37888300	0.40968500
O	-5.12445300	-1.57532400	0.35697800
C	-4.49654500	-3.84327300	0.48017400
H	-5.11652200	-4.01802700	1.37437500
H	-5.11524600	-4.10273900	-0.39380100
H	-3.61609400	-4.49770600	0.51421700
C	-0.89165200	-0.31740600	0.54991400
H	-0.71761600	0.28210100	1.45786000
H	-0.25571800	-1.21146100	0.57453900
H	-0.61047800	0.31243000	-0.30920600
Ni	-5.02932200	0.35599300	0.15613900
C	-5.28126300	2.24073700	0.47824200
C	-6.58594000	2.74199900	0.42739100
C	-4.21984700	3.01190600	0.95673400
C	-6.84311500	4.01928800	0.94287500
H	-7.39661700	2.14897500	0.00031200
C	-4.49017800	4.29328700	1.45962300
H	-3.20813100	2.61056000	0.95600400
C	-5.79496800	4.79583100	1.45187400
H	-7.86489500	4.40854200	0.93342600
H	-3.66792500	4.90225700	1.84538300

H	-5.99591200	5.79966300	1.83490400
Br	-5.82173700	0.50856100	-2.05487700
C	-2.99932200	3.69583100	-1.82226500
C	-2.43639500	2.38994100	-2.26258900
H	-2.47089500	2.27999300	-3.36550800
H	-1.39395400	2.25214000	-1.93660100
H	-3.03033300	1.54850300	-1.85893200
C	-4.31868200	4.17966700	-2.31192400
H	-4.23469800	4.66402400	-3.30939100
H	-5.02961500	3.34405400	-2.42737500
H	-4.76335300	4.92062300	-1.62850200
H	-2.40445600	4.34732800	-1.17510300

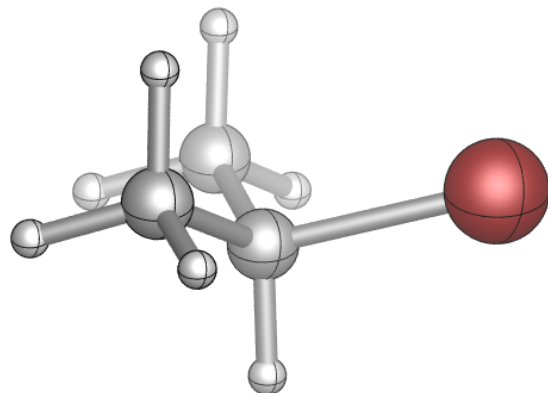
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4778.28000251

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -4777.49651484

**iPr-Br**



Zero-point correction= 0.093570 (Hartree/Particle)

Thermal correction to Energy= 0.098960

Thermal correction to Enthalpy= 0.099905

Thermal correction to Gibbs Free Energy= 0.064321

Sum of electronic and zero-point Energies= -2692.245354

Sum of electronic and thermal Energies= -2692.239964

Sum of electronic and thermal Enthalpies= -2692.239020

Sum of electronic and thermal Free Energies= -2692.274603

C -1.76973700 2.94632600 2.80829800

C -1.50495700 2.87037900 1.31519000

H -0.41627400 2.79919900 1.14944300

H -1.97839200 1.97918900 0.87478400

H -1.87574800 3.76236000 0.79039500

Br -3.74989200 3.16938400 3.11257100

C -1.28411200 1.73632900 3.58641000

H -0.19151300 1.64615500 3.46014100

H -1.50019600 1.82967400 4.66027400

H -1.75050200 0.81207500 3.21146000

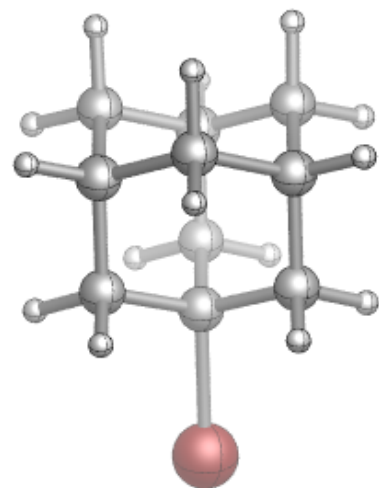
H -1.35916500 3.87336400 3.23130400

UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2692.78972539

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -2692.55656411

**Ad-Br**

Zero-point correction= 0.233256 (Hartree/Particle)

Thermal correction to Energy= 0.241651

Thermal correction to Enthalpy= 0.242595

Thermal correction to Gibbs Free Energy= 0.199568

Sum of electronic and zero-point Energies= -2963.523396

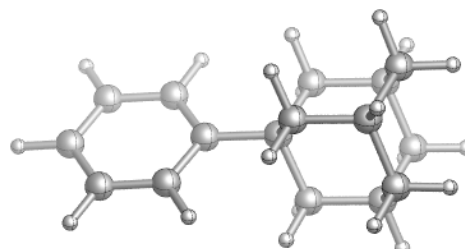
Sum of electronic and thermal Energies= -2963.515001

Sum of electronic and thermal Enthalpies= -2963.514057

Sum of electronic and thermal Free Energies= -2963.557084

C	-1.88329500	1.37104900	-0.00315300
H	-0.78191400	1.37449800	0.00737900
H	-2.22085500	2.41940600	0.00779100
C	-2.41569700	0.63798100	1.23278300
C	-2.40864800	0.64841100	-1.26606800
H	-2.02694600	1.17358800	-2.15702900
C	-1.91092000	-0.81021500	-1.26434300
H	-0.80809900	-0.83507000	-1.28427500
H	-2.26341100	-1.32960300	-2.17139100
C	-1.91038800	-0.80883400	1.25596600
H	-2.26836300	-1.31683900	2.16532200
H	-0.80930600	-0.82190600	1.27737400
C	-2.43463400	-1.53167700	-0.00756200
H	-2.07298600	-2.57298400	0.00601100
C	-3.97509900	-1.51171600	-0.00439200
H	-4.36116700	-2.04310000	-0.89055900
H	-4.35988500	-2.04142200	0.88367200
C	-3.94751600	0.67076200	1.25572500
H	-4.30007200	1.71402200	1.27552400
H	-4.32001500	0.17411300	2.16540400
C	-3.94917600	0.67029900	-1.26386000
H	-4.31347900	1.71155400	-1.28172800
H	-4.33425000	0.17570100	-2.17139400

C -4.47192100 -0.05272800 -0.00743600  
H -5.57405200 -0.03143000 0.00639700  
Br -1.72961200 1.58334000 2.88316800  
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2692.78972539  
UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -2692.55656411

**Ad-Ph**

Zero-point correction= 0.324546 (Hartree/Particle)

Thermal correction to Energy= 0.335948

Thermal correction to Enthalpy= 0.336892

Thermal correction to Gibbs Free Energy= 0.287610

Sum of electronic and zero-point Energies= -621.049416

Sum of electronic and thermal Energies= -621.038014

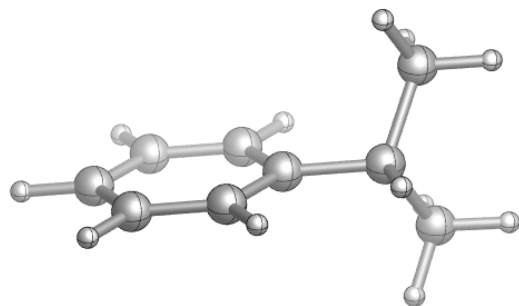
Sum of electronic and thermal Enthalpies= -621.037070

Sum of electronic and thermal Free Energies= -621.086353

C	-9.05162300	0.85471200	0.04815300
C	-8.76142400	0.74758300	1.42334000
C	-10.16553900	0.14499500	-0.43192500
C	-9.54413700	-0.03042700	2.27881000
H	-7.90510000	1.28308600	1.83989900
C	-10.95474300	-0.63687700	0.42163000
H	-10.43668600	0.19213500	-1.48658400
C	-10.64968800	-0.73005800	1.78097200
H	-9.28914200	-0.09075600	3.34040300
H	-11.81463900	-1.17621500	0.01482700
H	-11.26506400	-1.34040000	2.44703400
C	-7.75260900	4.04495400	-2.68996900
H	-7.11367000	4.69035700	-3.31727000
H	-8.78114200	4.43953100	-2.76026000
C	-7.27177300	4.08183800	-1.22799900
H	-7.30785900	5.11786900	-0.85006300
C	-7.70650700	2.59392500	-3.20303400
H	-8.06397000	2.55806500	-4.24614000
C	-6.26146400	2.06767800	-3.13250800
H	-5.60116700	2.68458200	-3.76635600
H	-6.21435600	1.03603700	-3.52239900
C	-5.82724500	3.55160200	-1.14811100
H	-5.46724600	3.58930600	-0.10516500
H	-5.15543600	4.19174800	-1.74563400
C	-5.77961700	2.10281000	-1.67094500
H	-4.74640800	1.72033500	-1.61021100
C	-6.69528200	1.21605600	-0.80637400
H	-6.66147600	0.17105500	-1.15967100
H	-6.32977800	1.21373200	0.23318200

C -8.18905000 3.19860600 -0.36157700  
H -9.22626000 3.57420400 -0.39577800  
H -7.86515600 3.25088600 0.69044200  
C -8.62079800 1.71159200 -2.33219600  
H -9.65923600 2.07586400 -2.40624800  
H -8.61137500 0.67811800 -2.71792900  
C -8.16600400 1.72198700 -0.85193500  
UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -622.03871909  
UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)  
HF= -621.50649605

**iPr-Ph**



Zero-point correction= 0.184466 (Hartree/Particle)

Thermal correction to Energy= 0.193027

Thermal correction to Enthalpy= 0.193971

Thermal correction to Gibbs Free Energy= 0.150892

Sum of electronic and zero-point Energies= -349.775767

Sum of electronic and thermal Energies= -349.767205

Sum of electronic and thermal Enthalpies= -349.766261

Sum of electronic and thermal Free Energies= -349.809340

C	-7.85943500	1.84940600	0.01558800
C	-9.01702800	1.45077700	-0.67127300
C	-7.00753200	0.84660600	0.51272700
C	-9.31823500	0.09662500	-0.85914900
H	-9.69435700	2.21412100	-1.06585100
C	-7.30281600	-0.50701700	0.32813500
H	-6.09905000	1.12607500	1.05317600
C	-8.46112100	-0.88852700	-0.35980900
H	-10.22624400	-0.18796500	-1.39771100
H	-6.62558100	-1.26896300	0.72361700
H	-8.69289700	-1.94693800	-0.50445000
C	-7.53947100	3.32441200	0.21624600
C	-6.22164800	3.71973500	-0.47099300
H	-5.36471200	3.18920800	-0.02409600
H	-6.24387100	3.47828000	-1.54546900
H	-6.03644800	4.80119500	-0.36592400
H	-8.34966000	3.89909500	-0.26480700
C	-7.52492300	3.70544400	1.70627000
H	-8.48052200	3.45256900	2.19213900
H	-6.72234500	3.17529600	2.24515400
H	-7.35204200	4.78702500	1.82998700

UB3LYP-D3/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -350.34501770

UM06/def2-TZVPP-CPCM(DMA)// UB3LYP-D3/def2-SVP-CPCM(THF)

HF= -350.03343472

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