

Thermochemical Value Tables Enthalpy of Formation and Bond Dissociation Energy for Multi-Fluorinated Ethanol's and Its Radicals at Standard Conditions

Hebah Abdel-Wahab, Joseph Bozzelli

Department of Chemistry and Environmental Science, New Jersey Institute of Technology, Newark, United States

Email address:

Dr.heathera@gmail.com (H. Abdel-Wahab), bozzellij@njit.edu (J. Bozzelli)

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Abstract: Fluorinated alcohols are used as solvents for proteins, organic compounds, and peptides, they are also known to be used in the organic synthesis industry due to its strong hydrogen bonding character. Halogenated hydrocarbons are mostly synthetically produced, and they don't exist naturally in the environment. Bond dissociation energies values can be used to explain reactivity and stability of chemical compounds. The objective of this research work is to calculate standard enthalpy of formation, and bond dissociation energy values for 18 different fluorinated ethanol; tri- fluorinated ethanol, tetra- fluorinated ethanol, and penta- fluorinated ethanol using Gaussian M-062x/6-31+g (d,p) method. Structures and thermochemical properties of Tri-, Tetra-, and Penta-Fluorinated Ethanol's and its Radicals were determined by the Gaussian M-062x/6-31+g (d,p) calculation: Enthalpies of formation for 18 fluorinated ethanol and some radicals were calculated with a popular ab initio and density functional theory methods: the Gaussian M-062x/6-31+g (d,p) via several series of isodesmic reactions. Bond dissociation energies for these fluorinated ethanol's and its radicals were also calculated.

Keywords: Enthalpy of Formation, Bond Dissociation Energy, Fluorinated Ethanol's, Gaussian M-062x/6-31+g (d,p), Computation, Thermochemical, Tri-Fluorinated Ethanol, Tetra-Fluorinated Ethanol, Penta-Fluorinated Ethanol

1. Introduction

Fluorinated alcohols are known to be used in the organic synthesis industry. They have strong hydrogen bonding donor character, and they are strong nucleophiles which allow organic reactions to occur without the using of a catalyst. Fluorinated alcohols have been used as solvents in epoxidation reactions, annulation reactions, nucleophilic substitution reactions, electrophilic reactions, and functionalization of multiple bonds [1].

Fluorinated alcohols are excellent solvents of proteins, peptides, and other organic compounds due to its physicochemical properties. Fluoro-alcohols, like other alcohols, can alter lipid bilayer properties and stability, and protein function. [2]

Most Halogenated hydrocarbons don't exist naturally in the environment and are synthetically produced. The major source of halogenated hydrocarbons to the atmosphere is agriculture. Crop spraying introduces halogenated

hydrocarbons to the environment entering through adsorption and deposition onto airborne particles or directly entering the aquatic system. Some halogenated hydrocarbons, PCBs, furans, and dioxins are by products, industrial waste that would unintentionally enter the atmosphere. [3]

The onset temperatures for energetic materials in the calorimetric measurements has been roughly predicted using molecular orbital calculations of bond dissociation energies. Stability and reactivity of chemical compounds can be explained using bond dissociation energies values. Standard enthalpies of formation estimated using semi-empirical MO calculations, MOPAC-PM7 package has been used previously to derive bond dissociation energy values for chemical compounds. [4]

Bond dissociation energies (BDEs) values for some inorganic compounds, lanthanide selenides and sulfides were measured in 2021 using Resonant two-photon ionization spectroscopy. The predissociation thresholds were found to be the BDE values for these molecules. The 0 K gaseous heat

of formation, $\Delta_f H^\circ_K$ for each molecule was also reported using this method. [5]

The amount of energy used to break a mole of covalently bonded gas molecule to pair of radicals is the bond dissociation energy. The units used for the bond dissociation energy is commonly kJ/mol. Covalent bonds can be broken heterolytically or homolytically. The heterolytic breaking of a covalent bond would result in the pair of electrons going to only one atom, either A or B.

$C-D \rightarrow C^+ + D^-$ or $C-D \rightarrow C^- + D^+$. The homolytic breaking of a covalent bond on the other hand would result in one electron staying with each atom, $\cdot A-B \rightarrow A\cdot + B\cdot$. Bond dissociation energy can be calculated for molecules as the difference in enthalpy of formation of products and reactants. Bond dissociation energy is a state function, as it doesn't depend on the mechanism or pathway on how bonds form or break. Energy of chemical reactions can be assessed using values for the bond dissociation energy. There are some systematic trends for the bond dissociation values; bonds dissociation energy varies with hybridization. For example, sp^3 hybridized carbons in hydrocarbons have smaller bond dissociation values compared to sp^2 hybridized carbons. The longer and weaker sp^3 hybridized bond are easier to break compared to the shorter and stronger sp^2 and sp hybridized bond (double and triple bonds). Among sp^3 hybridized bonds, bond dissociation values depend on its position, where it's on a primary, secondary or tertiary carbon. Methane has the strongest C-H bond with highest bond dissociation values, following C-H bonds on primary carbons, following C-H bonds on secondary carbons, following C-H bonds on tertiary carbons. [6]

Energetics of chemical processes can be assessed using bond dissociation energy values. Hess's Law has been used in the past and is currently being used to estimate reaction enthalpies by combining bond dissociation energies of bonds formed and bond dissociation energies of bonds broken. [23]

The energy change when forming a mole of compound from its component elements is called the enthalpy of formation, $\Delta_f H^\circ$. If heat is released when the elements combine to form the compound, enthalpy of formation would have a negative sign. If heat is absorbed when the elements combine to form the compound, enthalpy of formation would have a positive sign. Values of Enthalpy of formation is dependent upon temperature, pressure and physical states of reactants and products in the chemical reaction. Standard enthalpy of formation ($\Delta_f H^\circ$) is the enthalpy of formation at standard conditions; 1 atm pressure, 25°C, and 1 M aqueous solution concentration. Any element in its most stable form has a standard enthalpy of formation has a value of zero. Tabulated enthalpy of formation values can be used to calculate standard enthalpy of any reaction whose standard enthalpy of formation values are well known, [7]

$$\Delta H^\circ_{rxn} = \sum m \Delta_f H^\circ_{(products)} - \sum n \Delta_f H^\circ_{(reactants)}$$

The heat involved in chemical or physical change at constant temperature and pressure is enthalpy of reaction (H), and it's a thermodynamic quantity, $q = \Delta H$. [8]

Molecules, ions, or atoms containing at least an unpaired

electron in the valance shell are called free radicals. Free radicals are chemically reactive, unstable, and mostly short lived. Heat, electrolysis, electrical discharge, and ionizing radiation can generate free radicals. Free radicals are intermediates in many chemical reactions. Free radicals are important in atmospheric chemistry, combustion, plasma chemistry, polymerization, biochemistry, and they are important in many chemical reactions. [9]

In 2016 Hang Wang studied the thermodynamic properties of fluorinated methanol using CBS-QB3, M06-2X, WB97X, W1U, M06, B3LYP, CBS-APNO and G4 Calculations. Small standard deviation suggests good error cancellation of work reactions and accuracy. Small values for standard deviations was obtained in calculations using M06-2x/6-31+g (d,p) Gaussian method; it is an accurate method to calculate Enthalpy of fluorinated alcohols, it shows the second smallest standard deviation after CBS-QB3 method of calculation. Enthalpy of fluorinated methanol were studies in the past. [10]

Halogenated compounds have low reactivity, are highly stable, and are used in industry. They are of concern to the environment due to their persistence in the environment, and its widespread use. Their thermochemical properties must be studied in order to understand the reduction and oxidation reactions involving these molecules. [11]

2. Computational Method

The Global-hybrid meta-GGA density functional approximation, GGA, DFT Gaussian M-062x/6-31+g (d,p) method of calculation has been used to initially analyze frequencies, optimized structures, and thermo energies of the molecules studied. In the GGA, generalized gradient approximation, the density functional depends on the down and up spin densities and the reduced gradient. In the meta GGA, the functional also depends on the up and down spin kinetic energy densities. A hybrid GGA is a combination of GGA with Hartree-Fock exchange. The hybrid meta GGA is a combination of meta GGA with Hartree-Fock exchange. [12]

A series of Isodesmic Reactions and composite calculations were employed to calculate enthalpy of formation of tri-, terta-, and penta- fluorinated ethanol's. Calculations were performed using Gaussian 16 program. All reported calculations of enthalpy of formation are for standard conditions of 1 atm pressure and 298K. The Gaussian M-062x/6-31+g (d,p) level of calculation has been used for this study as this method of calculation were successfully employed in the past when applied to fluoro hydrocarbons 6 with small reported standard deviations values. [13]

The standard deviation is calculated using the following formula. [14]

$$B = \sqrt{1/N \sum (x_i - \mu)^2}$$

Where, X_i is the mean; the average of the numbers, μ is the actual numbers to be calculated the standard deviation of, and $1/N \sum (x_i - \mu)^2$ is the variance.

3. Isodesmic and Isogyric Reaction

The Gaussian M-062x/6-31+g (d,p) method of calculation has been used to calculate the enthalpy of formation of tri-, tera-, and penta-fluorinated ethanol's and its radicals. Work reactions and reference species have been used to calculate enthalpy of formation of fluorinated ethanol's using this method.

To cancel any systematic error in the molecular orbital calculations using this method, the number of each type of bond must be conserved in each isodesmic reactions. By the careful choice of the isodesmic reactions, all enthalpies of formation calculations are allowed to accuracies close to experimental values.

The ΔH_{f298K° values of all reference species but the fluoroethanols are known, the ΔH_{f298K° of the target species fluoro-ethanols, is obtained from this data and the calculated ΔH_{rxn} , 298°. ΔH_{f298K° calculated using two different reference molecules are within \pm (0 to 0.60 Kcal mol⁻¹).

4. Reference Species

Table 1 lists the Standard enthalpy of formation for the reference species used in isodesmic reactions with their uncertainties. Table 2 provides all calculated standard enthalpy of formation values, $\Delta_f H^\circ_{(298)}$ for of tri-, tera-, and penta-fluorinated ethanol's and its radicals.

Table 1. Reference Species in the Isodesmic Reactions Standard Enthalpy of Formation Values (kcal mol⁻¹) [15].

Species	$\Delta_f H^\circ_{(298)}$	Species	$\Delta_f H^\circ_{(298)}$
CH ₃ F	-56.54 ± 0.07 ^a	CH ₃ OOH	-30.96 ± 0.67 ^b
CH ₃ CH ₂ F	-56.62 ± 0.48 ^b	CH ₃ CH ₂ OOH	-38.94 ± 0.81 ^b
CH ₃ CH ₂ CH ₂ F	-65.42 ± 1.11 ^a	CH ₃ CH ₂ CH ₂ OOH	-44.03 ± 0.67 ^b
CH ₂ F ₂	-70.24 ± 1.30 ^a	CH ₃ OO [•]	2.37 ± 1.24 ^b
CH ₃ CHF ₂	-108.07 ± 1.46 ^a	CH ₃ CH ₂ OO [•]	-6.19 ± 0.92 ^b
CH ₃ CH ₂ CHF ₂	-107.67 ± 0.48 ^h	CH ₃ CH ₂ CH ₂ OO [•]	-11.35 ± 1.24 ^b
CHF ₃	-120.87 ± 1.62 ^a	CH ₄	-17.81 ± 0.01 ^c
CH ₃ CH ₂ CHF ₂	-125.82 ± 1.65 ^a	CH ₃ CH ₃	-20.05 ± 0.04 ^c
CH ₃ CF ₃	-166.71 ± 1.97 ^h	CH ₃ CH ₂ CH ₃	-25.01 ± 0.06 ⁱ
CH ₃ CH ₂ CF ₃	-166.09 ± 0.48 ^h	CH ₃ CH ₂ CH ₂ CH ₃	-30.07 ± 0.08 ^j
CH ₃ [•]	-180.51 ± 2.05 ^a	CH ₃ O [•]	5.15 ± 0.08 ^c
CH ₃ CH ₂ [•]	-185.48 ± 2.15 ^a	CH ₃ CH ₂ O [•]	-3.01 ^d
CH ₃ CH ₂ CH ₂ [•]	34.98 ± 0.02 ^c	OH	8.96 ± 0.01 ^c
H	28.65 ± 0.07 ^c	CH ₃ OH	-47.97 ± 0.04 ^c
O	24.21 ± 0.24 ^{gj}	CH ₃ CH ₂ OH	-56.07 ± 0.05 ⁱ
CF ₄ ¹³	24.18 ⁱ	HOO [•]	2.94 ^{ej}
CF ₃ CH ₂ CHF ₂ ¹³	52.10 ^c	HOOH	-32.39 ± 0.04 ^{fi}
	59.57 ^c		-32.37 ⁱ
	-223.15	CHF ₂ CHF ₂ ¹³	-212.13
	-286.18	CF ₃ CHF ₂ ¹³	-267.79

^aWang [10], ^bWang [15], ^cRuscic [16], ^dBurke [17], ^eChase [18], ^fLuo [19], ^gBodi [20], ^hPedley, ⁱATcT Tables [21], ^jCsontos [22].

5. Results and Discussion

5.1. Standard Enthalpy Values

Isodesmic work reactions from M-062x/6-31+g (d,p) method of calculation has been utilized to perform calculation of standard enthalpy of formations for tri-, tera-, and penta-fluorinated ethanol's and some radicals. Reference species and their standard enthalpy of formation along with their uncertainties has been used in all isodesmic work reactions.

The calculated sum of thermal enthalpies using the Gaussian M06-2x/6-31+g (d,p) Level of Theory for all target fluorinated ethanol's, their radicals, and for reference species has also been used. The standard enthalpy of formation in kcal mol⁻¹ for all reference species are listed, table 1. Standard deviation values 14 are listed for all calculated Gaussian M06-2x/6-31+g (d,p) standard enthalpy of formations values. The calculated Gaussian M06-2x/6-31+g (d,p) standard enthalpy of formation for tri-, tera-, and penta- fluorinated ethanol's and some radicals are listed table 2.

Table 2. Isodesmic reactions used in calculating standard enthalpy of formation, ΔH°_{Rxn} for tri-, tera-, and penta-fluorinated ethanol's and its radicals using the Gaussian M06-2x/6-31+g (d,p) Level of Theory.

Isodesmic Reactions Target Specie	$\Delta H^\circ_{Rxn(298)}$ Hartrees	$\Delta H^\circ_{Rxn(298)}$ Kcal/mole ¹	$\Delta_f H^\circ_{(298)}$ kcal mol ⁻¹	Error kcal mol ⁻¹
CF ₃ CH ₂ OH + CH ₄ = CH ₃ CH ₂ OH + CHF ₃				
-452.696144 -40.447961 -154.926666 -338.204669	0.01277	8.013303	-213.1	±2.03
-17.81 -56.21 -166.71				
CF ₃ CH ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CH ₃ CF ₃				
-452.696144 -79.717768 -154.926666 -377.492814	-0.00557	-3.49398	-213.2	±2.14
-20.05 -56.21 -180.51				

Isodesmic Reactions Target Specie	$\Delta H^\circ_{\text{Rxn (298)}}$ Hartrees	$\Delta H^\circ_{\text{Rxn (298)}}$ Kcal/mole ¹	$\Delta_f H^\circ_{(298)}$ kcal mol ⁻¹	Error kcal mol ⁻¹
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-213.15 ±2.09	
Standard Deviation over rxns 0.05 kcal mol ⁻¹				
CF ₃ CH ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CF ₃				
-452.048269 -79.717768 -154.268107 -377.492814	0.005116	3.210341	-166.7	± 2.09
-20.05 -3.01 -180.51				
CF ₃ CH ₂ OH + CH ₄ = CH ₃ CH ₂ O• + CHF ₃				
-452.048269 -40.447961 -154.268107 -338.204669	0.023454	14.71762	-166.6	±1.98
-17.81 -3.01 -166.71				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-166.65±2.04	
Standard Deviation over rxns 0.05 kcal mol ⁻¹				
CF ₃ CH ₂ O• + CH ₃ CH ₂ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CH ₂ CF ₃				
-452.023032 -118.99092 -154.268107 -416.766483	-0.02064	-12.9537	-150.5	±2.21
-25.02 -3.01 -185.48				
CF ₃ CH ₂ O• + CH ₄ = CH ₃ CH ₂ O• + CHF ₃				
-452.023032 -40.447961 -154.268107 -338.204669	-0.00178	-1.11885	-150.8	±1.98
-17.81 -3.01 -166.71				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-150.70±2.09	
Standard Deviation over rxns 0.16 kcal mol ⁻¹				
CHF ₂ CHFOH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CH ₃ CF ₃				
-452.686514 -79.717768 -154.926666 -377.492814	-0.0152	-9.5369	-207.1	±2.14
-20.05 -56.21 -180.51				
CHF ₂ CHFOH + CH ₄ = CH ₃ CH ₂ OH + CHF ₃				
-452.686514 -40.447961 -154.926666 -338.204669	0.00314	1.970381	-207.1	±2.03
-17.81 -56.21 -166.71				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-207.10±2.09	
Standard Deviation over rxns 0.00 kcal mol ⁻¹				
C•F ₂ CHFOH + CH ₄ = CH ₃ CH ₂ O• + CHF ₃				
-452.032761 -40.447961 -154.268107 -338.204669	0.007946	4.986194	-156.9	±1.98
-17.81 -3.01 -166.71				
C•F ₂ CHFOH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CF ₃				
-452.032761 -79.717768 -154.268107 -377.492814	-0.01039	-6.52108	-156.9	±2.09
-20.05 -3.01 -180.51				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-156.90 ±2.04	
Standard Deviation over rxns 0.00 kcal mol ⁻¹				
CHF ₂ C•FOH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CF ₃				
-452.028711 -79.717768 -154.268107 -377.492814	-0.01444	-9.0625	-154.4	±2.09
-20.05 -5.01 -180.51				
CHF ₂ C•FOH + CH ₃ CH ₂ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CH ₂ CF ₃				
-452.028711 -118.990915 -154.268107 -416.766483	-0.01496	-9.39006	-154.1	±2.21
-25.02 -5.01 -185.48				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-154.30±2.15	
Standard Deviation over rxns 0.16 kcal mol ⁻¹				
CHF ₂ CHFO• + CH ₄ = CH ₃ CH ₂ O• + CHF ₃				
-452.019492 -40.447961 -154.268107 -338.204669	-0.00532	-3.34024	-148.6	±1.98
-17.81 -3.01 -166.71				
CHF ₂ CHFO• + CH ₃ CH ₂ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CH ₂ CF ₃				
-452.019492 -118.99092 -154.268107 -416.766483	-0.02418	-15.1751	-148.3	±2.21
-25.02 -3.01 -185.48				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-148.50±2.09	
Standard Deviation over rxns 0.16 kcal mol ⁻¹				
CH ₂ FCF ₂ OH + CH ₄ = CH ₃ CH ₂ OH + CHF ₃				
-452.696009 -40.447961 -154.926666 -338.204669	0.012635	7.928589	-213.0	±2.03
-17.81 -56.21 -166.71				
CH ₂ FCF ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CH ₃ CF ₃				
-452.696009 -79.717768 -154.926666 -377.492814	-0.0057	-3.57869	-213.1	±2.14
-20.05 -56.21 -180.51				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-213.05±2.09	
Standard Deviation over rxns 0.05 kcal mol ⁻¹				
C•HFCF ₂ OH + CH ₄ = CH ₃ CH ₂ O• + CHF ₃				
-452.036986 -40.447961 -154.268107 -338.204669	0.012171	7.637424	-159.5	±1.98
-17.81 -3.01 -166.71				
C•HFCF ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CF ₃				
-452.036986 -79.717768 -154.268107 -377.492814	-0.00617	-3.86985	-159.6	±2.09
-20.05 -3.01 -180.51				
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-159.55±2.04	
Standard Deviation over rxns 0.08 kcal mol ⁻¹				

Isodesmic Reactions Target Specie	$\Delta H^\circ_{\text{Rxn (298)}}$ Hartrees	$\Delta H^\circ_{\text{Rxn (298)}}$ Kcal/mole ¹	$\Delta_f H^\circ_{(298)}$ kcal mol ⁻¹	Error kcal mol ⁻¹
CH ₂ FCF ₂ O• + CH ₄ = CH ₃ CH ₂ O• + CHF ₃ -452.019556 -40.447961 -154.268107 -338.204669 -17.81 -3.01 -166.71	-0.00526	-3.30008	-148.6	±1.98
CH ₂ FCF ₂ O• + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CH ₃ CF ₃ -452.019556 -79.717768 -154.268107 -377.492814 -20.05 -3.01 -180.51	-0.0236	-14.8074	-148.7	±2.09
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.05 kcal mol ⁻¹			-148.65±2.04	
CF ₃ CFHOH + CH ₄ = CH ₃ CH ₂ OH + CF ₄ -551.950082 -40.447961 -154.926666 -437.46666 -17.81 -56.21 -223.15	0.004717	2.959965	-264.5	±0.10
CF ₃ CFHOH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CHF ₂ CHF ₂ -551.950082 -79.717768 -154.926666 -476.716001 -20.05 -56.21 -212.13	0.025183	15.80258	-264.1	±0.09
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-264.30±0.10	
CF ₃ C•FOH + CH ₄ = CH ₃ CH ₂ O• + CF ₄ -551.293882 -40.447961 -154.268107 -437.46666 -17.81 -3.01 -223.15	0.007076	4.440261	-212.8	±0.10
CF ₃ C•FOH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CHF ₂ CHF ₂ -551.293882 -79.717768 -154.268107 -476.716001 -20.05 -3.01 -212.13	0.027542	17.28288	-212.4	±0.40
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-212.60±0.30	
CF ₃ CFHO• + CH ₄ = CH ₃ CH ₂ O• + CF ₄ -551.281264 -40.447961 -154.268107 -437.46666 -17.81 -3.01 -223.15	-0.00554	-3.47766	-204.9	±0.10
CF ₃ CFHO• + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CHF ₂ CHF ₂ -551.281264 -79.717768 -154.268107 -476.716001 -20.05 -3.01 -212.13	0.014924	9.364959	-204.5	±0.40
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-204.70±0.30	
CF ₂ HCF ₂ OH + CH ₄ = CH ₃ CH ₂ OH + CF ₄ -551.947553 -40.447961 -154.926666 -437.46666 -17.81 -56.21 -223.15	0.002188	1.372992	-262.9	±0.10
CF ₂ HCF ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CHF ₂ CHF ₂ -551.947553 -79.717768 -154.926666 -476.716001 -20.05 -56.21 -212.13	0.022654	14.21561	-262.5	±0.10
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-262.70±0.10	
C•F ₂ CF ₂ OH + CH ₄ = CH ₃ CH ₂ O• + CF ₄ -551.286788 -40.447961 -154.268107 -437.46666 -17.81 -3.01 -223.15	-1.8E-05	-0.0113	-208.3	±0.10
C•F ₂ CF ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CHF ₂ CHF ₂ -551.286788 -79.717768 -154.268107 -476.716001 -20.05 -3.01 -212.13	0.020448	12.83132	-207.9	±0.40
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-208.10±0.30	
CF ₂ HCF ₂ O• + CH ₄ = CH ₃ CH ₂ O• + CF ₄ -551.26873 -40.447961 -154.268107 -437.46666 -17.81 -3.01 -223.15	-0.01808	-11.3429	-197.0	±0.10
CF ₂ HCF ₂ O• + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CHF ₂ CHF ₂ -551.26873 -79.717768 -154.268107 -476.716001 -20.05 -3.01 -212.13	0.00239	1.499749	-196.6	±0.40
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹ Standard Deviation over rxns 0.20 kcal mol ⁻¹			-196.80±0.30	
CF ₃ CF ₂ OH + CH ₃ CH ₂ CH ₃ = CH ₃ CH ₂ OH + CF ₃ CH ₂ CHF ₂ -651.207522 -118.99092 -154.926666 -615.260935 -25.02 -56.21 -286.18	0.010836	6.799698	-324.2	±0.10
CF ₃ CF ₂ OH + CH ₃ CH ₃ = CH ₃ CH ₂ OH + CF ₃ CHF ₂ -651.207522 -79.717768 -154.926666 -575.976217 -20.05 -56.21 -267.79	0.022407	14.06062	-318.0	±1.70
CF ₃ CF ₂ OH + CH ₄ = CF ₃ OH + CH ₃ CHF ₂ -651.207522 -40.447961 -413.437575 -278.225303 -17.81 -218.11 -120.87	-0.00739	-4.64044	-316.5	±1.70
Reported $\Delta_f H^\circ$ (298) kcal mol ⁻¹			-319.60±1.20	

Isodesmic Reactions Target Specie	$\Delta H^\circ_{\text{Rxn}} (298)$ Hartrees	$\Delta H^\circ_{\text{Rxn}} (298)$ Kcal/mole ¹	$\Delta_f H^\circ (298)$ kcal mol ⁻¹	Error kcal mol ⁻¹
Standard Deviation over rxns 3.30 kcal mol ⁻¹				
CF ₃ CF ₂ O• + CH ₄ = CHF ₂ O• + CH ₃ CF ₃				
-650.527081 -40.447961 -313.498636 -377.492814	-0.01641	-10.2962	-250.2	±2.06
-17.81 -97.82 -180.51				
CF ₃ CF ₂ O• + CH ₃ CH ₂ CH ₃ = CH ₃ CH ₂ O• + CF ₃ CH ₂ CHF ₂				
-650.527081 -118.990915 -154.268107 -615.260935	-0.01105	-6.93148	-257.2	±0.10
-25.02 -3.01 -286.18				
CF ₃ CF ₂ O• + CH ₃ CH ₃ = CH ₃ CH ₂ O• + CF ₃ CHF ₂				
-650.527081 -79.717768 -154.268107 -575.976217	0.000525	0.329443	-251.1	±1.70
-20.05 -3.01 -267.79				
Reported $\Delta_f H^\circ (298)$ kcal mol ⁻¹			-252.80±1.30	
Standard Deviation over rxns 3.11 kcal mol ⁻¹				

Hartrees, kcal mole⁻¹

*SD Standard Deviation kcal mol⁻¹

Errors reported avg of sum of uncertainties in rxn's reference species.

5.2. Bond Energies

The difference in bond dissociation energies of products and reactants for the hemolysis is called the bond dissociation energy. The bond dissociation energy values don't depend on the pathway by which it occurs, it doesn't depend on how bonds are

formed or on how bonds break, therefore bond dissociation energies are state functions. Energetics of chemical processes can be assessed using bond dissociation energy values.

The calculated bond dissociation energy values in kcal/mol for Methyl C-H, Ethyl C-H, and hydroxyl O-H bonds are listed table 3.

Table 3. Bond Dissociation Energy (BDE's) of Tri-, Trtra-, and Penta- Fluorinated Ethanol's Using Gaussian M-062x/6-31+g (d,p) method of calculation.

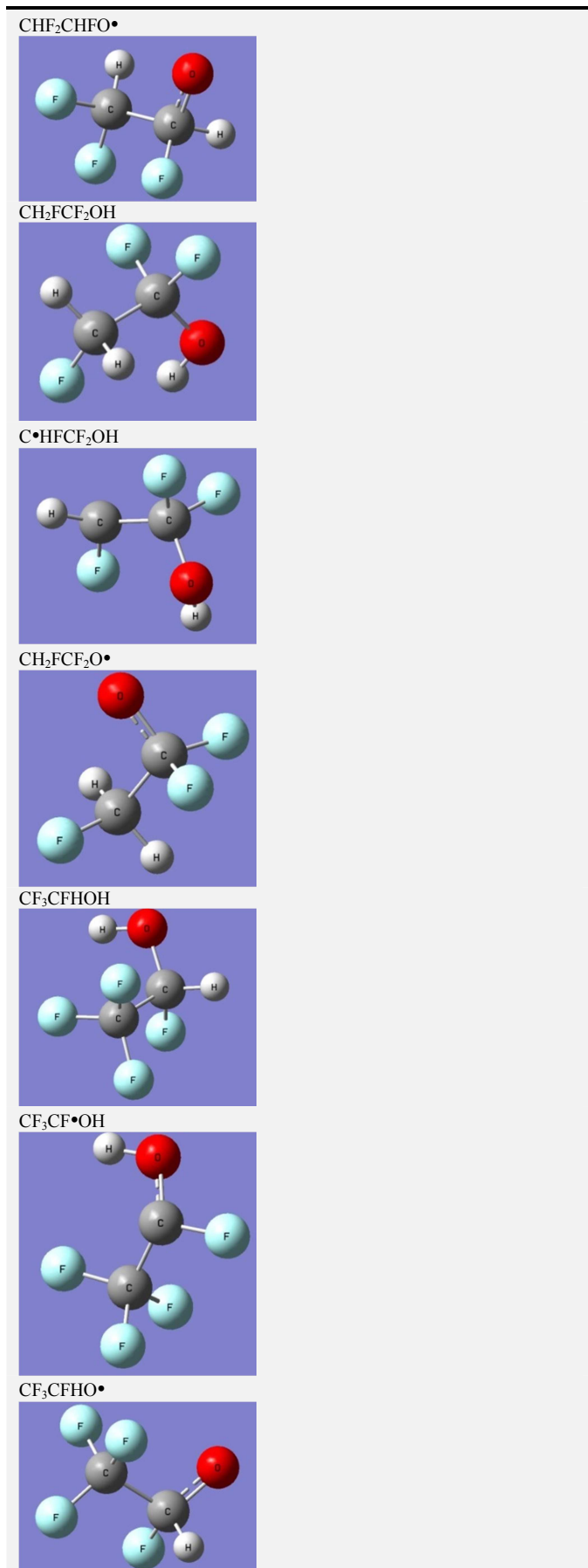
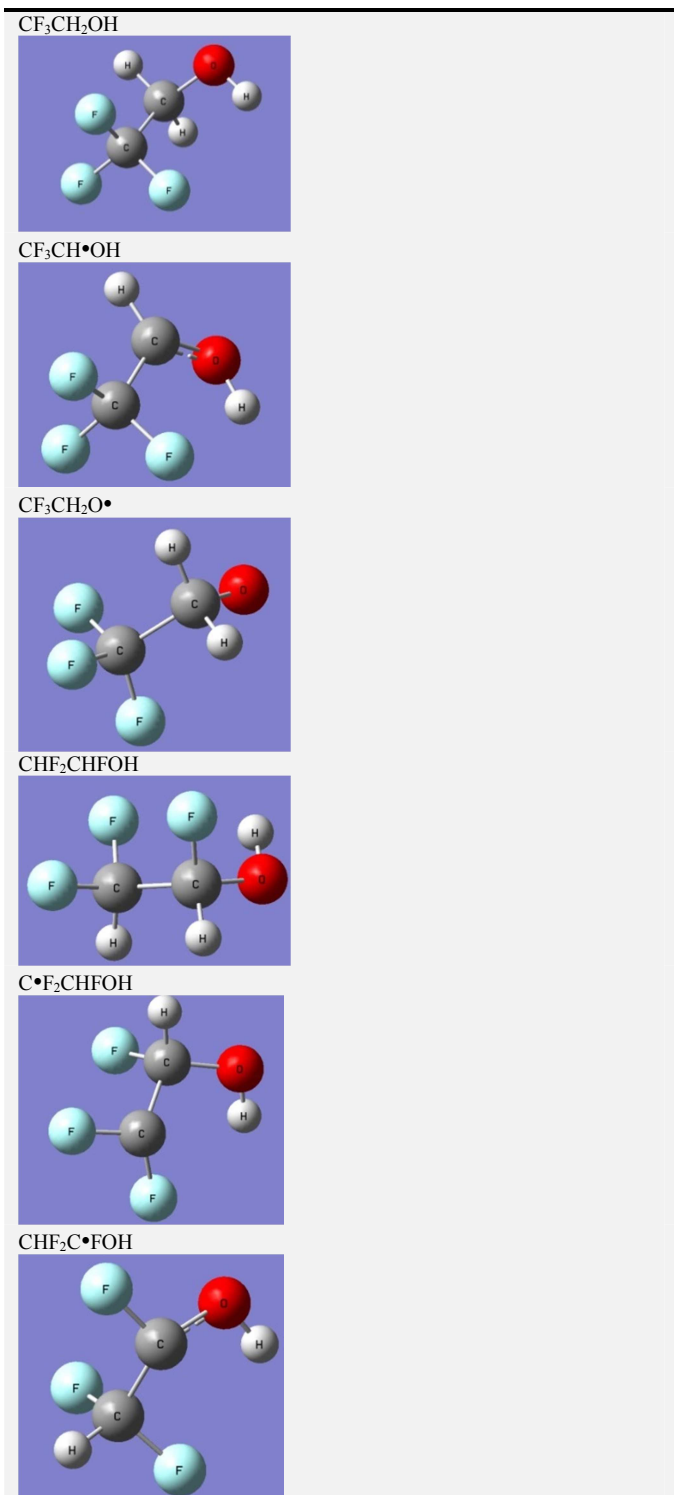
Reactions	Bond Dissociation Energy ^a (Kcal mol ⁻¹)	* Error Kcal mol ⁻¹
CF ₃ CH-HOH		
CF ₃ CH-HOH = H• + CF ₃ CH• OH	98.60± 2.07	± 2.07
-213.15 ±2.09 52.10 -166.65±2.04		
CF ₃ CH ₂ O-H		
CF ₃ CH ₂ O-H = H• + CF ₃ CH ₂ O•	114.55±2.09	±2.09
-213.15 ±2.09 52.10 -150.70±2.09		
C-HF ₂ CHFOH		
C-HF ₂ CHFOH = H• + C• F ₂ CHFOH	102.30±2.07	±2.07
-207.10±2.09 52.10 -156.90 ±2.04		
CHF ₂ C-HFOH		
CHF ₂ C-HFOH = H• + CHF ₂ C• FOH	104.90±2.10	±2.10
-207.10±2.09 52.10 -154.30±2.15		
CHF ₂ CHFO-H		
CHF ₂ CHFO-H = H• + CHF ₂ CHFO•	110.70±2.09	±2.09
-207.10±2.09 52.10 -148.50±2.09		
HC-HFCF ₂ OH		
HC-HFCF ₂ OH = H• + C• HFCF ₂ OH	105.60±2.07	±2.07
-213.05±2.09 52.10 -159.55±2.04		
CH ₂ FCF ₂ O-H		
CH ₂ FCF ₂ O-H = H• + CH ₂ FCF ₂ O•	116.50±2.07	±2.07
-213.05±2.09 52.10 -148.65±2.04		
CF ₃ CF-HOH		
CF ₃ CF-HOH = H• + CF ₃ CF• OH	103.80±0.20	±0.20
-264.30±0.10 52.10 -212.60±0.30		
CF ₃ CFHO-H		
CF ₃ CFHO-H = H• + CF ₃ CFHO•	111.70±0.20	±0.20
-264.30±0.10 52.10 -204.70±0.30		
CF ₂ -HCF ₂ OH		
CF ₂ -HCF ₂ OH = H• + CF ₂ • CF ₂ OH	106.70±0.20	±0.20
-262.70±0.10 52.10 -208.10±0.30		
CF ₂ HCF ₂ O-H		
CF ₂ HCF ₂ O-H = H• + CF ₂ HCF ₂ O•	118.00±0.20	±0.20
-262.70±0.10 52.10 -196.80±0.30		
CF ₃ CF ₂ O-H		
CF ₃ CF ₂ O-H = H• + CF ₃ CF ₂ O•	118.90±1.30	±1.30
-319.60±1.20 52.10 -252.80±1.30		

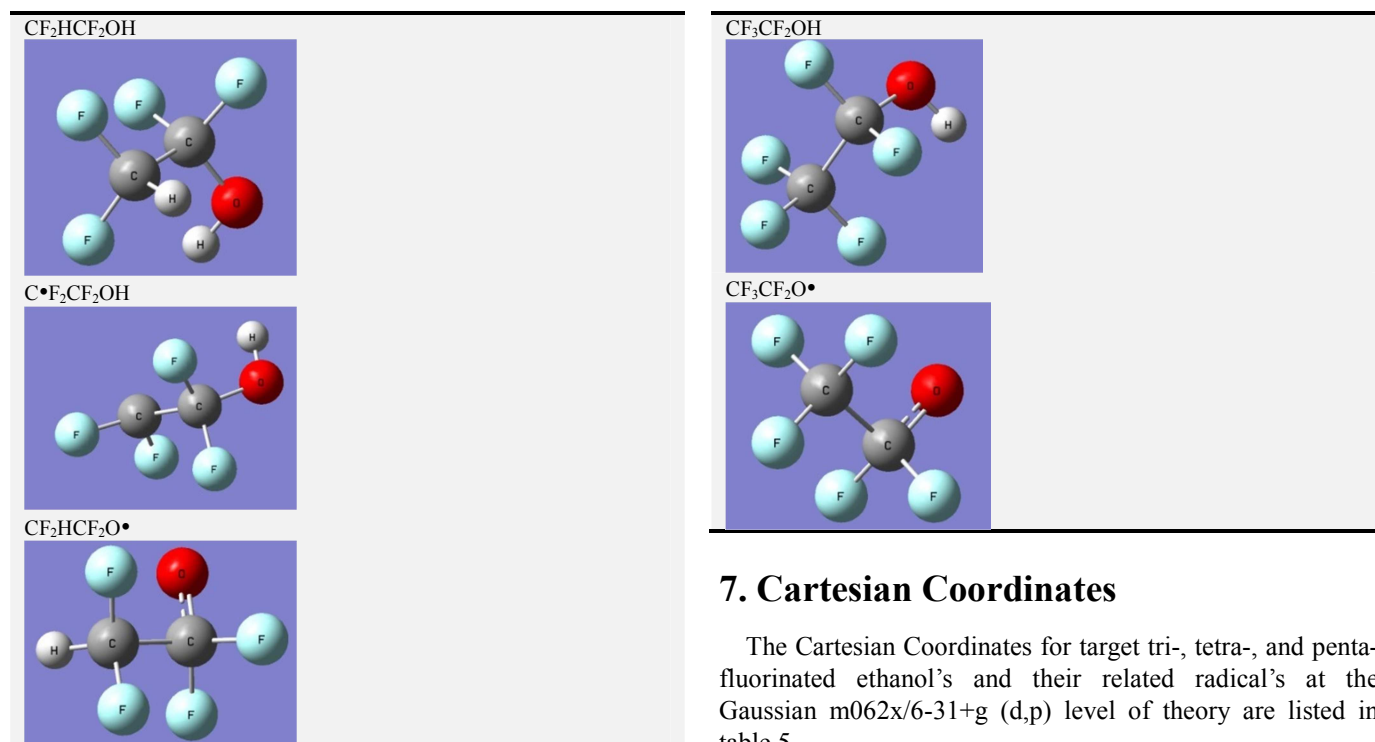
*Errors reported avg of sum of uncertainties in rxn' s reference specie.

6. Optimized Structures

The GaussView software has been used along with gaussian output files to provide a picture of the Gaussian M06-2x/6-31+g (d,p) optimized structure for each molecule in this study table 4.

Table 4. Optimized geometries for target tri-, tetra-, and penta- fluorinated ethanol's and their related radicals calculated by Gaussian M06-2x/6-31+g (d,p) level of theory.





7. Cartesian Coordinates

The Cartesian Coordinates for target tri-, tetra-, and penta-fluorinated ethanol's and their related radical's at the Gaussian m062x/6-31+g (d,p) level of theory are listed in table 5.

Table 5. Cartesian Coordinates in Angstroms for target fluorinated ethanol and their related radical's geometries at the m062x/6-31+g (d,p) level of theory.

Molecule	Cartesian Coordinates				
	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
CF ₃ CH ₂ OH	1	6	0.000015792	0.000045146	0.000015456
	2	6	0.000056272	-0.000131331	-0.000001091
	3	9	-0.000036015	-0.000005130	-0.000021513
	4	9	0.000035235	0.000054970	-0.000018991
	5	9	-0.000001701	0.000051428	0.000034529
	6	1	-0.000016200	0.000007968	0.000014306
	7	1	-0.000023506	-0.000003074	-0.000011946
	8	8	-0.000022602	-0.000015046	0.000001930
	9	1	-0.000007275	-0.000004931	-0.000012678
CF ₃ C•HOH	1	6	-0.000055227	-0.000030630	-0.000017226
	2	6	0.000117503	0.000159657	-0.000153816
	3	9	-0.000061675	0.000035455	0.000071731
	4	9	-0.000053827	-0.000176074	0.000047967
	5	9	0.000080802	-0.000106445	0.000024638
	6	1	0.000001337	0.000046611	-0.000002493
	7	8	0.000004111	0.000030529	-0.000039199
	8	1	-0.000033025	0.000040898	0.000068398

Molecule	Cartesian Coordinates				
CF ₃ CH ₂ O•	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	-0.000037529	0.000088167	0.000122350
	2	6	-0.000033636	-0.000121335	-0.000217962
	3	9	0.000020892	0.000000631	0.000073365
	4	9	0.000016383	-0.000001502	0.000002646
	5	9	-0.000021622	0.000041297	0.000066627
	6	1	-0.000020139	-0.000044741	-0.000019536
CHF ₂ CHFOH	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	0.000034684	-0.000097837	0.000016859
	2	6	0.000077734	-0.000008144	0.000008869
	3	9	-0.000027798	0.000012351	0.000007573
	4	9	-0.000002203	0.000054383	-0.000025902
	5	1	-0.000018584	0.000015264	0.000009448
	6	9	0.000005269	0.000001678	-0.000006105
C•F ₂ CHFOH	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	-0.000071214	0.000095917	0.000002631
	2	6	-0.000097297	-0.000427988	0.000070858
	3	9	0.000109064	0.000106013	0.000068648
	4	9	0.000002068	0.000252877	-0.000098474
	5	1	-0.000001211	0.000016670	-0.000018084
	6	9	0.000033735	-0.000023289	-0.000027474
CHF ₂ C•FOH	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	-0.000038782	0.000051546	0.000051509
	2	6	-0.000109395	-0.000076573	-0.000071886
	3	9	0.000040508	0.000010628	0.000077586
	4	9	0.000020881	0.000006702	-0.000016053
	5	1	0.000007533	-0.000016674	-0.000041204
	6	9	0.000090694	-0.000038267	-0.000029261
CHF ₂ CHFOH	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	-0.000038782	0.000051546	0.000051509
	2	6	-0.000109395	-0.000076573	-0.000071886
	3	9	0.000040508	0.000010628	0.000077586
	4	9	0.000020881	0.000006702	-0.000016053
	5	1	0.000007533	-0.000016674	-0.000041204
	6	9	0.000090694	-0.000038267	-0.000029261

Molecule	Cartesian Coordinates				
	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
CHF ₂ CHFO•	1	6	0.000055628	-0.000106091	0.000003381
	2	6	0.000053463	-0.000000718	-0.000007006
	3	9	-0.000043989	-0.000020986	0.000005782
	4	9	-0.000016210	-0.000035868	0.000004447
	5	1	-0.000011020	-0.000013533	0.000007886
	6	1	-0.000010458	0.000024649	-0.000005524
	7	9	-0.000051835	0.000115358	-0.000005383
	8	8	0.000022121	0.000009470	-0.000053144
	9	1	0.000002301	0.000027718	0.000049560
CH ₂ FCF ₂ OH	1	6	0.000135651	0.000057453	0.000051033
	2	6	-0.000243304	0.000078716	-0.000013330
	3	9	-0.000116433	0.000005813	0.000097761
	4	1	-0.000077576	-0.000009293	0.000043932
	5	1	-0.000038364	-0.000018670	0.000006345
	6	9	0.000035849	0.000027631	-0.000096988
	7	9	0.000045365	0.000036806	-0.000102497
	8	8	0.000181449	-0.000179561	-0.000003213
	9	1	0.000077364	0.000001106	0.000016957
C•HFCF ₂ OH	1	6	-0.000009430	0.000025594	0.000005992
	2	6	0.000017955	-0.000004162	0.000000569
	3	9	0.000001322	0.000052928	-0.000001501
	4	1	-0.000003779	0.000024041	0.000008036
	5	9	-0.000012190	-0.000046066	0.000011421
	6	9	-0.000025702	-0.000032617	0.000028822
	7	8	0.000010500	-0.000018093	-0.000017353
	8	1	0.000021325	-0.000001626	-0.000035986
CH ₂ FCF ₂ O•	1	6	0.000157230	-0.000033765	0.000067368
	2	6	-0.000084968	-0.000038414	-0.000127598
	3	9	-0.000060007	0.000028432	0.000052860
	4	1	-0.000002632	-0.000019789	-0.000013222
	5	1	-0.000027022	0.000016735	-0.000013331
	6	9	-0.000031881	-0.000031453	0.000013157
	7	9	0.000080040	0.000071255	0.000005918
	8	8	-0.000030759	0.000006999	0.000014849

Molecule	Cartesian Coordinates				
	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
CF ₃ CFHOH	1	6	-0.000139743	-0.000019177	-0.000028107
	2	6	-0.000044583	-0.000023228	-0.000002049
	3	9	0.000017824	0.000030702	0.000057967
	4	9	-0.000069952	-0.000066058	-0.000012198
	5	9	0.000058051	-0.000059520	-0.000018868
	6	9	0.000166545	0.000025560	-0.000007913
	7	1	0.000003967	0.000022634	0.000010499
	8	8	-0.000043206	0.000028605	-0.000066782
	9	1	0.000051097	0.000060482	0.000067450
CF ₃ C•FOH	1	6	0.000033580	0.000004460	0.000005322
	2	6	0.000020098	0.000029526	-0.000031153
	3	9	0.000021581	-0.000000752	0.000016405
	4	9	0.000016549	0.000006426	0.000007990
	5	9	-0.000041946	0.000031215	0.000004491
	6	9	-0.000000439	-0.000039185	-0.000010685
	7	8	-0.000042830	-0.000023720	0.000011783
	8	1	-0.000006593	-0.000007971	-0.000004153
CF ₃ CFHO•	1	6	0.000536610	0.000642487	-0.000238587
	2	6	-0.000336421	-0.000010459	-0.000050530
	3	9	0.000179754	0.000022776	0.000069636
	4	9	0.000013212	-0.000037718	0.000038270
	5	9	0.000033732	0.000056583	-0.000043087
	6	9	-0.000277185	-0.000172926	0.000053380
	7	1	-0.000020915	-0.000096792	0.000011479
	8	8	-0.000128787	-0.000403952	0.000159440
CF ₂ HCF ₂ OH	1	6	0.000034684	-0.000097837	0.000016859
	2	6	0.000077734	-0.000008144	0.000008869
	3	9	-0.000027798	0.000012351	0.000007573
	4	9	-0.000002203	0.000054383	-0.000025902
	5	1	-0.000018584	0.000015264	0.000009448
	6	9	0.000005269	0.000001678	-0.000006105
	7	9	-0.000005043	-0.000008859	0.000006453
	8	8	-0.000067666	0.000036885	-0.000008025
	9	1	0.000003607	-0.000005722	-0.000009170

Molecule	Cartesian Coordinates				
	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
C•F ₂ CF ₂ OH	1	6	0.000024554	0.000023392	0.000004415
	2	6	-0.000034477	0.000024861	-0.000019704
	3	9	0.000055726	0.000012426	0.000032150
	4	9	0.000019915	0.000028025	-0.000017047
	5	9	0.000004005	-0.000011188	-0.000003720
	6	9	0.000015833	-0.000043273	0.000042429
	7	8	-0.000068386	-0.000006654	0.000010769
	8	1	-0.000017169	-0.000027587	-0.000049293
CF ₂ HCF ₂ O•	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	0.000006128	-0.000032081	0.000064624
	2	6	0.000009899	-0.000032233	-0.000045587
	3	9	-0.000037818	-0.000059393	0.000013603
	4	9	0.000041223	0.000037095	-0.000050906
	5	1	-0.000021281	0.000006425	-0.000018170
	6	9	-0.000013457	0.000031071	0.000014828
CF ₃ CF ₂ OH	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	-0.000029279	0.000029059	-0.000014751
	2	6	0.000037295	0.000003602	0.000009039
	3	9	-0.000004027	-0.000031190	-0.000022595
	4	9	-0.000016447	-0.000026160	-0.000006065
	5	9	0.000008918	-0.000023108	0.000015387
	6	9	-0.000006450	0.000029260	-0.000000832
CF ₃ CF ₂ O•	Center Number	Atomic Number	Forces (Hartrees/Bohr)		
			X	Y	Z
	1	6	0.000090683	0.000175823	0.000176247
	2	6	-0.000004772	-0.000005489	-0.000074419
	3	9	-0.000035079	0.000031055	-0.000015124
	4	9	-0.000026035	-0.000069252	0.000016300
	5	9	0.000038428	-0.000012873	-0.000000972
	6	9	-0.000080090	-0.000021765	-0.000033887
	7	9	0.000031147	-0.000065731	-0.000034377
	8	8	-0.000014283	-0.000031768	-0.000033768

8. Conclusions

The ab initio and Global-hybrid meta-GGA density function method, Gaussian M06-2x/6-31+g (d,p) has been used to calculate thermodynamic properties of 18 tri-, tetra-, and penta- fluorinated ethanol's and their related radicals. Standard enthalpy of formation at the Gaussian M06-2x/6-31+g (d,p) calculation level has been calculated using multiple work reactions, work reactions were employed for cancellation of calculation errors.

The Thermochemical properties: the O—H, secondary methyl C—H, and ethyl C-H Bond Dissociation Energies (BDEs), and Standard Enthalpy of formation (298K) values has been calculated for tri-, tetra-, and penta- fluorinated Ethanol's and their related Radicals: CH₃CH₂-xFxOH, CH₃-xFxCH₂-xFxOH, and CH₂-xFxCH₂OH.

The C-H bond dissociation energies range from 98.6 to 104.9 Kcal mol⁻¹ on the secondary ethyl carbons, and from 103.3 to 106.7 Kcal mol⁻¹ on the primary methyl carbons. The O-H bond energies range from 110.7 to 118.9 Kcal mol⁻¹.

Calculated values for the O-H bond energies for penta-fluorinated ethanol are higher than those of O-H bond energies for tri- fluorinated ethanol. The calculated the O-H bond energies increased by intruding more fluorine atoms to either methyl or ethyl carbons.

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