Chem 106	
TA	Name

# **Worksheet: Crystal Violet and Quantum Theory**

\*\*Where short answers are required, please use complete sentences.\*\*

This experiment requires 4 Molecular Energy, and 2 Molecular Orbital calculations. List the job numbers your answers are based on:

	Molecular Energy	Molecular Orbitals
Crystal violet cation CV		
Crystal violet alcohol CVOH		
OH4-water cluster		
CV-OH4-water product		
CVOH4-water transition state		

## A. Partial charges in the CV<sup>+</sup> ion



In WebMO, open your completed **Molecular Energy** calculation for **CV**. Check that the atoms are labeled with the **index number**. (If not, click the Preferences tool on the left toolbar, and under Labels, select Index.)



1. (a) Scroll down to the Partial Charges table. **Fill in** the following table using the tabulated values or click the blue magnifying glass icon within the column heading of the table: To switch back to atom indexes, click the Reset Viewer button at the bottom of the display.

CV		
	Charge	
Atom	(±0.001)	
N 2		
N 24		
N 43		
C 7		
C 19		
C 38		
Central C 15		
Total charge	1.000	
Total minus C15		

•	) What <b>percent</b> of t r <b>bon</b> (C 15)?	he total charge is on th	e <b>central</b>
	hat <b>percent</b> of the gether?	total charge is on <b>all</b> th	e other atoms
th		ft, look at the charges opears that the 3 rings a	
	quite different	slightly different	identical

(c). Calculations show that **CV** is resonance stabilized with the (+) charge spread out equally among the three aromatic rings. This is analogous to the way negative charge is shared in the formate ion. See page 2 in the handout.

The following formulas show several resonance forms of CV. Within each circled region change **four** single bonds into double bonds and add **one** "+" symbol. There are several ways of doing this, but **only one** is consistent with the charges in the table in (a).

(d) Go back to the <u>informational webpage</u> and follow the link to **CV cation surfaces.** This page shows several surfaces:

- the total electron density distribution of the ion. The total electron density surface shows the volume around the nuclei where the electron density is equal to or greater than 0.025 electrons per unit volume. Recall that electron density is highest close to the nucleus, and gradually decreases going out from the nucleus. Thus, an electron density surface can be drawn at any arbitrary value. The value used here is consistent with our notion of the shape of the molecule.
- The **electrostatic potential** surface combines the total electron density surface with a color scale that reports a **local excess of electrons** (red) or **lack of electrons** (blue) at points around the surface. **Red** spots occur where electrons protrude, such as lone pairs or  $\pi$  bonds. **Green** areas occur where there is zero formal charge and no excess or relative lack of electrons.
- The **electrophilic reactivity** surface combines the total electron density surface with a color scale showing the preferred location for **addition of electrons** (**blue**), which, for example, might originate from OH<sup>-</sup> ion<sup>-</sup>. This happens to coincide with the center of the lowest unoccupied molecular orbital (LUMO).

(1) Is the overall appearance of the **electron density** surface consistent with the **resonance description** of CV? What do you see, and what is the significance of this?

(2) What is the color of the **electrostatic potential** surface **above the N atoms**? What causes this color to appear on the surface there?

#### B. The OH<sup>-</sup> reactant and the CV-OH<sup>-</sup> transition state

#### 1. Reaction energy and activation energy

Open the completed molecular energy jobs of CV, OH-4water complex, the CVOH-4water complex, and the CVOH-4water transition state. For each one, on the View Job page, record the energy in the table below. Look for "RB3LYP Energy" under Calculated Quantities, Overview.

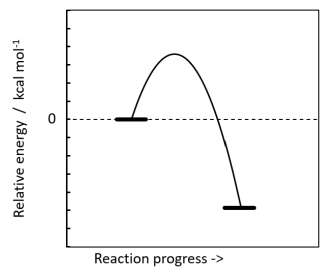
For the  $\Delta E$  rows, convert Hartrees to kcal/mol using the factor 627.5095 kcal/mol per Hartree.

		RB3LYP (Hartrees)* (to the 6 <sup>th</sup> decimal place)	
Doostonts	OH-4-water		
Reactants	CV		
Product	CVOH-4-water		
Transition state	CVOH-4-water TS		keel/mel/record out to the
			kcal/mol (record out to the 3 <sup>rd</sup> decimal place)
	$\Delta E_{reaction}$		
	$\Delta E_{activation}$		

<sup>\*&</sup>quot;R" stands for Restricted, meaning all the electrons occur in pairs in the molecule. "B3LYP" is the type of density functional theory we are using to calculate molecular energy.

- Label the graph at the right with "reactants," "TS" and "product".
- Label several tick marks on the y-axis that are consistent with the ΔE values in the table.
- Draw a vertical **up** arrow and label it " $\Delta E_{activation}$ ".
- Draw a vertical **down** arrow and label it " $\Delta E_{reaction}$ ".

Is the crystal violet +	OH- reaction	
endothermic	or <b>exothermic</b>	<u></u> ?
(Check one)		



#### 2. Carbon-oxygen and hydrogen bond changes

(a) In addition to formation of a new C-O bond, several H-bonds undergo significant changes during the CV + OH<sup>-</sup> reaction. In WebMO, open the molecular energy jobs for the OH-4-water reactant, the CVOH-4-water transition state, and the CVOH-4-water product. In each one, use the **Adjust tool** to click the C-O and H-O pairs indicated in the table below. The next page has a table with corresponding atom numbers. The distance is displayed in the **lower left-hand corner**. Enter distances in the table.



		C-O or OH distance (Å)	
H-bond	OH4-water	CV-OH-TS-4-water	CVOH-4-water
1			
2			
3			
4			
C-O			
	3	3 1————————————————————————————————————	3 - 4 - 2 CO
_		(The ring-groups remove	d to clarify the graphics.)

H-bond		Indexes of the H-bonded aton	ns
#	OH4-water	CV-OH-TS-4-water	CVOH-4-water
1	4, 13	62, 71	62, 71
2	4, 6	62, 64	62, 64
3	4, 11	62, 69	62, 69 (not an H-bond, but record the distance)
4	1, 10	59, 68	59, 68

- (b) Describe how these bonds **change** going from **reactants to T.S.** and from **T.S.to products**.
- #1.
- #2.
- #3.
- #4.
- C--O
- (c) Qualitatively, what is the overall energy impact of these changes during reaction? In other words, why does the energy **go up** to the transition state, then **down** to the product?

### C. The violet color of CV

In WebMO open your CV and CVOH Molecular Orbital calculation jobs by clicking the job name in the Job Manager. Scroll down to the Molecular Orbitals table near the bottom of the View Job page. For each, record the energies of the HOMO and LUMO in the table (next page).

Here is a section of the CV MO table with some explanatory annotations. Note that the MOs are listed in order of most stable (lowest energy) at the top, to least stable (highest energy) at the bottom.

Orbital	<b>+</b>	Symmetry	Occupancy	♦ Spin ▲	Energy (au) Hartrees	•	
MO#			electrons/MO		Hartrees		occupied
97		-	2	-	-0.25776	õ	<b>↑</b>
98		-	2	-	-0.23237	P	
99		-	2	-	-0.19428	P	
100		-	2	-	-0.19422	P	номо
101		-	0	-	-0.09613	P	LUMO
102		-	0	-	-0.01531	<i>≨</i> ⊃	
103		-	0	-	-0.00465	P	
							unoccupie

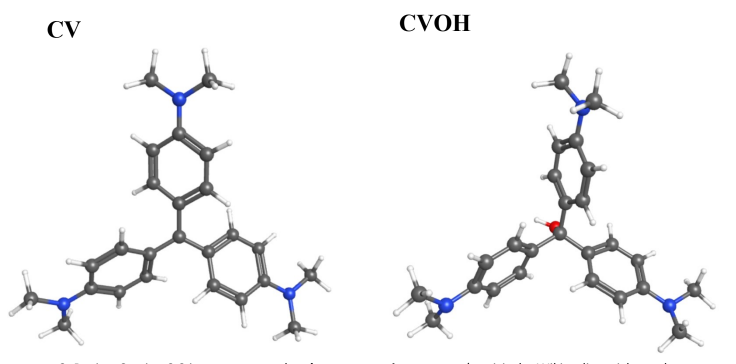
1. Fill in the boxes based on your two Molecular Orbital calculations. Calculate the predicted the predicted wavelength  $\lambda$  for each species. Assume that the main peak in the UV-VIS spectrum which occurs at the  $\lambda_{\text{max}}$  is due to absorption of a photon whose energy equals the energy gap ( $\Delta E$ ) between LUMO and HOMO.

	cv	сvон	
HOMO energy (Hartrees)			
LUMO energy (Hartrees)			
ΔЕ <sub>LUMO – номо</sub> (Hartrees)			
ΔЕ <sub>LUMO – HOMO</sub> (kcal/mol)*			
Calculated $\lambda$ (nm) (the $\lambda_{max}$ )			
*1 Hartree = 627.5095 kcal/mol			
Use $\Delta E = hv = hc/\lambda$ $h = Planck's constant = 6.626 \times 10^{-34}$ $c = speed of light = 3.000 \times 10^{10} c$ Avogadro's number = 6.022 x 10 <sup>23</sup> n 1 kcal = 4.184 x 10 <sup>3</sup> Joule	m/sec		

Show your work here. Include units and the correct number of significant figures.

2. Compared to the LUMO in CVOH, the **LUMO of CV is spread over more atoms,** which is the main reason for the lower energy of CV's LUMO. Make rough drawings of the two LUMOs on the structures below. Hint: To get the best view of these MOs, set the "MO Isosurface value" to 0.030 by clicking Preferences button on the left-hand toolbar.

Preferences	
Display	Isosurfaces Colors Mechanics
MO isosurface value	0.030



3. Review Section 3.2 in your text on the **electromagnetic spectrum** (or visit the Wikipedia article on the same topic). Circle the spectral range where the calculated  $\lambda_{max}$  wavelengths occur.

CVOH	ultraviolet	visible	infrared
CV	ultraviolet	visible	infrared

4. When an **electron-rich ion**, such as **OH**<sup>-</sup>, reacts with with a cation like CV, it pours its excess valence electrons into the cation's LUMO because the LUMO provides the lowest energy orbital to hold the electrons. The CV atom where the **largest lobe of the LUMO** is located is therefore the one most likely to bond to OH-.

Which CV atom is this - judging from your drawing in (2) or looking at the LUMO graphic in WebMO?

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