

Butler University [Digital Commons @ Butler University](https://digitalcommons.butler.edu/)

[Scholarship and Professional Work - LAS](https://digitalcommons.butler.edu/facsch_papers) College of Liberal Arts & Sciences

1984

A SCC MO Calculation on the Tetracyanoethylene-benzene **Complex**

Joe Kirsch

Shannon Lieb Butler University, slieb@butler.edu

Mark Cisneros

Follow this and additional works at: [https://digitalcommons.butler.edu/facsch_papers](https://digitalcommons.butler.edu/facsch_papers?utm_source=digitalcommons.butler.edu%2Ffacsch_papers%2F778&utm_medium=PDF&utm_campaign=PDFCoverPages)

Part of the Chemistry Commons

Recommended Citation

Kirsch, Joe; Lieb, Shannon; and Cisneros, Mark, "A SCC MO Calculation on the Tetracyanoethylenebenzene Complex" Proceedings of the Indiana Academy of Science / (1984): 181-185. Available at [https://digitalcommons.butler.edu/facsch_papers/778](https://digitalcommons.butler.edu/facsch_papers/778?utm_source=digitalcommons.butler.edu%2Ffacsch_papers%2F778&utm_medium=PDF&utm_campaign=PDFCoverPages)

This Article is brought to you for free and open access by the College of Liberal Arts & Sciences at Digital Commons @ Butler University. It has been accepted for inclusion in Scholarship and Professional Work - LAS by an authorized administrator of Digital Commons @ Butler University. For more information, please contact [digitalscholarship@butler.edu.](mailto:digitalscholarship@butler.edu)

A SCC MO Calculation on the Tetracyanoethylene-benzene Complex

Joe Kirsch, Shannon Lieb and Mark Cisneros Department of Chemistry Butler University Indianapolis, Indiana 46208

Introduction

The tetracyanoethylene-benzene charge transfer complex is formed through the interaction of the pi electron density of the benzene and the pi antibonding orbitals of the tetracyanoethylene. Two sandwich type structures of the complex have been proposed in the literature (2) and are shown in Figure ¹ . Molecular orbital calculations

FIGURE 1. Proposed Structures of Benzene-Tetracyanoethylene Complex.

can be used to determine the nature and energy of the absorption which results from complex formation. If the calculations are carried out as a function of the benzenetetracyanoethylene intermolecular distance, the intermolecular distance that yields the best agreement with the observed charge transfer energy can be obtained.

Calculations

The MO calculations were carried out on ^a VAX 11/780 computer. The necessary input data for the SCC MO program are given in Table 1. The atomic coordinates are calculated from standard bond lengths and bond angles. These molecular parameters are also listed in Table 1.

The SCC MO calculations require the evaluation of overlap integrals, coulomb integrals, and resonance integrals. These integrals are the elements of the secular deter minant. Calculation of the overlap integrals, S_{ij} , using Slater type atomic orbitals, the effective nuclear charge, and the atomic coordinates have been described in the literature (1). Valence orbital ionization energies, VOIE, are used to approximate the coulomb

integrals, H_{ii} (1). The Wolfsberg-Helmholtz approximation is used to determine the resonance integrals, H_{ii} (1). The secular determinant is then solved from these values of S_{ij} , H_{ii} , H_{ij} to obtain two sets of MO mixing coefficients (Eigenvectors) and MO energies (eigenvalues).

The eigenvectors resulting from the calculation are used in a Mulliken population analysis to calculate new atomic charges for each atom in the complex. These new atomic charges, output atomic charges, are compared to the input atomic charges. If the input and the output charges are different, the difference times 0.1 is used as a new input charge; and the calculation is recycled until the input and output atomic charges converge and self consistent charges on the atoms are obtained.

Results and Discussion

The benzene-tetracyanoethylene complex has 74 valence electrons and 70 valence atomic orbitals in its basis set. This basis set and collection of valence electrons will yield 70 molecular orbitals with the first 37 molecular orbitals being doubly populated with electrons. The 38th molecular orbital is the lowest unoccupied molecular orbital. The lowest energy electronic transition, the charge transfer band, will then occur bet ween the 37th MO and the 38th MO.

Examination of the eigenvectors for molecular orbitals 37 and 38 show that all of the atomic orbital coefficients are near zero except those for the pi 2p type atomic orbitals. This indicates that molecular orbitals 37 and 38 are primarily pi type molecular orbitals. The values of the eigenvectors of the pi 2p atomic orbitals for molecular orbitals 37 and 38 are listed in Table 2. The atom numbering system is given in Figure 2 and ³ for each proposed structure. Further examination of the eigenvectors show that MO ³⁷ is bonding for benzene (C2-C3 and C5-C6), bonding for tetracyanoethylene (CN groups), and bonding for the complex for both structures. Molecular orbital 38, however, is antibonding for the tetracyanoethylene part of the complex for both structures. Finally, it can be noted that the eigenvectors indicate more electron density, larger values for the eigenvectors, on the tetracyanoethylene for MO ³⁸ than for MO ³⁷ in both structures. In summary, the analysis of the eigenvectors for MO 37, highest occupied, and MO 38, lowest empty, supports the notion of an electronic transition, charge transfer, from ^a benzene pi bonding MO to ^a tetracyanoethylene pi antibonding MO as ^a description of the charge transfer band.

Atom Number*		Structure A		Structure B		
		MO37	MO38	MO37	MO38	
Benzene	$C-1$	$\boldsymbol{0}$	$\overline{0}$	$\mathbf{0}$	θ	
Benzene	$C-2$	$-.36$.38	$-.32$	$-.25$	
Benzene	$C-3$	$-.36$	$-.38$	$-.32$	$-.25$	
Benzene	$C-4$	θ	$\mathbf{0}$	Ω	θ	
Benzene	$C-5$.36	$-.38$.32	.25	
Benzene	$C-6$.36	.38	.32	.25	
TCNE	$C-7$	$\mathbf{0}$	θ	.07	.47	
TCNE	$C-8$	θ	$\mathbf{0}$	$-.07$	$-.47$	
TCNE	$C-9$.13	$-.21$.14	.22	
TCNE	$C-10$	$-.13$.21	.14	.22	
TCNE	$C-11$	$-.13$	$-.21$	$-.14$	$-.22$	
TCNE	$C-12$.13	.21	$-.14$	$-.22$	
TCNE	$N-13$	$-.29$	$-.35$.32	$-.30$	
TCNE	$N-14$.29	.35	.32	$-.30$	
TCNE	$N-15$	$-.29$.35	$-.32$.30	
TCNE	$N-16$.29	$-.35$	$-.32$.30	

Table 2. Pi AO Eigenvectors for the Benzene—Tetracyanoethylene Complex

*Refers to the numbering system in figures 2 and ³

Table ³ shows the energy difference of MO ³⁸ and MO 37, energy of the charge transfer band, as a function of the benzene-tetracyanoethylene intermolecular distance. Figures 4 and ⁵ are plots of this data for each proposed structure. The plots for both structures show minimum near the observed charge transfer absorption energy for an intermolecular distance of ² angstroms, structure A—³⁸⁶ nm, 2.04 A; structure B—³⁸⁹ nm, 2.09 A.

B

Eigenvector Analysis in Table 2. Eigenvector Analysis in Table 2.

FIGURE 2. Atom Numbering System for FIGURE 3. Atom Numbering System for

	Structure A
r (angstroms)	$E(MO-38) - E(MO-37)$ [nm]
1.97	413
1.98	409
1.99	405
2.00	401
2.04	386
2.05	388
2.10	403
	Structure B
r (angstroms)	$E(MO-38) - E(MO-37)$ [nm]
2.03	409
2.04	406
2.05	402
2.06	398
2.07	395
2.08	391
2.09	389
2.10	392
2.12	398

TABLE 3. E(MO 38) - E(MO 37) versus the Benzene-Tetracyanoethylene Intermolecular Distance

Charge Transfer Band Energy vs Benzene-TCNE Intermolecular Distance. FIGURE 4.

Figure 5. Charge Transfer Band Energy vs Benzene—TCNE Intermolecular Distance.

Acknowledgments

The authors wish to thank the Holcomb Research Institute and the Butler University Academic Grants Committee for their funding of this project.

Literature Cited

- 1. McGlynn, S.P., L.G. Vanquickenborne, M. Kinoshita, and D.G. Carrol, 1972, Introduction to Applied Quantum Chemistry, Holt Rinehart Winston, New York, N.Y., 48, 97-140.
- 2. Mobley, M.J., K.E. Rieckhoff, and E.M. Voight, 1978, Spectroscopic Studies on the Conformations of Electron Donor Acceptor Complexes of Tetra cyanoethylene, J. Physical Chem., 82, 2005-2012.