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# A Theoretical Study of Charge-Transfer Properties of a new Material involving Naphthalenyl Unit



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#### **ABSTRACT**

In this study, the charge transport properties of 4- [(E)- [(2-hydroxy-1-naphthalenyl) methylene] amino] -3-methyl benzoic acid (HNMB) molecule were determined by using Marcus and DFT methods. In addition to this, reorganization energies (\lambda e and λh), the ionization potentials (IPs) and the electron affinities (EAs) are also reported. It is understood from the results that HNMB has suitable photovoltaic properties in terms of solar cells.

### ARTICLE INFO

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Marcus method, charge transport properties, reorganization energy, solar cells

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In this report, the charge transfer properties for HNMB sample was investigated by Amsterdam density functional (ADF2019) [14].

#### 1. INTRODUCTION

In the current scenario, solar energy has received a lot of attention due to its increased energy need and availability worldwide. This energy need is aimed to be met from organic-based solar cells due to the high production costs of silicon-based solar cells [1-3]. In addition, there are results showing that the conjugate molecule used in the solar cell increases the structureproperty relationship efficiency [4]. In addition, whether the molecule is suitable for photovoltaic cells is also linked to the relationship between structure and property.

HNMB is a commercial molecule important for mechanochromatic luminescent and OLED materials. In addition, HNMB was synthesized by Zhang et al [5], and its emission properties and reliability in pressure sensor applications were investigated.

The HNMB molecule is a polycyclic aromatic hydrocarbon containing naphthalene. Naphthaleneaddition polycyclic aromatic hydrocarbons are available in many experimental [6-9] and theoretical studies [10-13].

## 2. METHOD

ADF is a successful computational chemistry software for calculating molecules in terms of structure, electronics, optics and more. Firstly, the dimer structure of the subject compound has been optimized in the ADF program and is given in Fig. 1. Next, the charge transfer integrals of HNMB molecule optimized at B3LYP/TZP level was calculated.

Charge transfer rate (W) of HNMB is found by following equation [15-18], known as the Marcus-Hush equation.

$$W = \frac{V^2}{\hbar} \left(\frac{\pi}{\lambda k_B T}\right)^{1/2} \exp\left(-\frac{\lambda}{4k_B T}\right) \tag{1}$$

where  $\lambda$  is the reorganization energy, T is the temperature, V is the effective charge transfer integral and  $\hbar$  and  $k_B$  are the Planck and Boltzmann constants, respectively.

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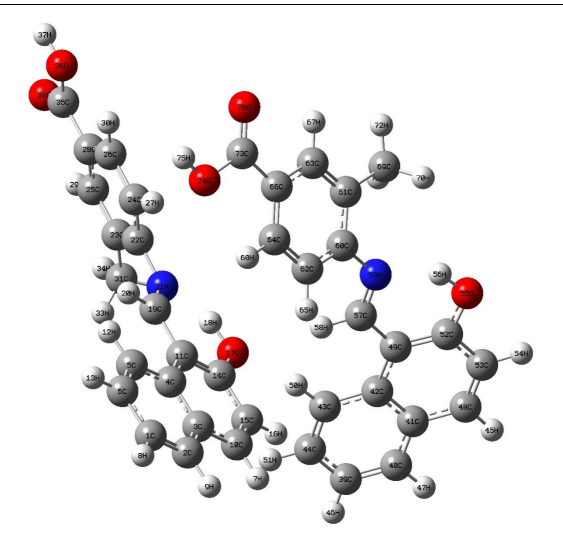


Fig. 1. Optimized structure of HNMB dimer

 $V_{ij}$  representing the electronic coupling between intermolecules; depending on the spatial overlap  $(S_{ij})$ , charge transfer integrals  $(J_{ij})$  and site energies  $(e_i(j))$ , respectively,

$$V = \frac{J_{ij} - S_{ij}(e_i + e_j)/2}{1 - S_{ij}^2}$$
 (2)

Electron (or hole) reorganization energies,  $\lambda_e$  (or  $\lambda_h$ ) can be written as follows:

$$\lambda_h = \lambda_1 + \lambda_2 = [E^+(g^0) - E^+(g^+)] + [E^0(g^+) - E^0(g^0)]$$

$$[(E^+(g^0) - E^0(g^0)] - [(E^+(g^+) - E^0(M^+))]$$
(3)

and

$$\lambda_e = \lambda_3 + \lambda_4 [E^0(g^-) - E^0(g^0)] + [E^-(g^0) - E^-(g^-)] [E^0(g^-) - E^-(g^-)] - [E^0(g^0) - E^-(g^0)]$$
(4)

 $E^0(g^0)$  is the energy of the neutral molecule in neutral geometry,  $E^{+/-}(g^{+/-})$  is ion energy (cation/anion) in ionic

geometry,  $E^{+/-}(g^0)$  is ion energy (cation/anion) in neutral geometry, and  $E^0(g^{+/-})$  in ionic geometry, the energies of the neutral molecule are shown in equations 3 and 4.

Ionization potentials (IPa/IPv) and electron affinities (EAa/Eav) significantly affect the injection ability of the sample [19-22]. The vertical and adiabatic ionization potentials (IP $_{\rm v}$  /IP $_{\rm a}$ ), the vertical and adiabatic electron affinities (Ea $_{\rm v}$ /EA $_{\rm a}$ ) of the HNMB were obtained as follows:

$$IP(v) = E^{+}(g^{0}) - E^{0}(g^{0})$$
(5)

$$IP(a) = E^{+}(g^{+}) - E^{0}(g^{0})$$
 (6)

$$EA(v) = E^{0}(g^{0}) - E^{-}(g^{0})$$
(7)

### 3. RESULTS AND DISCUSSION

The calculated reorganization energies of HNMB molecule are tabulated in Table 1. These calculated values are factors affecting the performance of solar cells. For

example, the mobility of electrons and holes is closely related to the reorganizational energy. The reorganization energy of an electron or hole varies inversely with the transport rate. It can be seen from Table 1 that HNMB can be used as electron transport material in solar cells.

The bigger EA and smaller IP means better electron and hole transport [23]. Table 1 shows all

adiabatic/vertical EAa/v and IPa/v values of HNMB. The EAa/v and IPa/v values of HNMB 6.416/5.544 and 0.639/0.884 eV. Therefore, the HNMB molecule can form suitable for the electron injection layer.

**Table 1.** The calculated reorganization energies (in eV).

Molecule	λ <sub>electron</sub>	$\lambda_{hole}$	IPa	IP <sub>v</sub>	EAa	EA <sub>v</sub>
HNMB	0.682	0.951	0.639	0.884	6.416	5.544

In order to calculate effective transfer integrals, the HNMB dimer structure was created in the anisotropic position of the molecule. Transfer integral results of this geometry are given in Table 2. The larger the transfer integral and the greater the charge transfer will be. The electron transfer integral of HNMB is higher than its hole transfer integral. So, it can be said that HNMB is a good candidate as electron transfer material.

Charge transfer rates ( $W_{electron}$  and  $W_{hole}$ ) are calculated using equation 1 for HNMB in parallel geometry at 298.15 K and are tabulated in Table 2. HNMB molecule has a high electron transfer rate ( $260 \times 10^9 \text{ s}^{-1}$ ).

**Table 2.** The charge transfer integrals (in eV) and the charge transfer rates (in s<sup>-1</sup>).

Molecule	Velectron	Vhole	Welectron	Whole
HNMB	0.01757	0.00838	260x10 <sup>9</sup>	140x10 <sup>9</sup>

#### 4. CONCLUSION

In the present work, the charge transfer properties of HNMB are obtained by theoretical calculations. Both reorganization energies and Marcus-Hush theory results proved that HNMB is a good electron transfer material. Thus, with these results, HNMB can be used as electron-transporting layers in solar cells and OLED structures.

#### **REFERENCES**

- [1] C.W. Tang, Two-layer organic photovoltaic cell, Applied Physics Letters, 1968, **48**, 2, 183-185
- [2] N. S. Sariciftci, Plastic photovoltaic devices, Materials Today, 2004, 7, 9, 36–40.
- [3] A. Aboulouard, A. G. Gürek, and M. E. Idrissi, Computational study of organic small molecules based on imidazolinone for photovoltaic applications, Energy Sources, Part A: Recovery, Utilization, and Environmental Effects, 2020, 1827092.
- [4] Y. J. Cheng, Yang S. H., Hsu, C. S., Synthesis of Conjugated Polymers for Organic Solar Cell

Applications, Chemical Reviews, 2009, **109**, 11, 5868-5923.

- [5] M. Zhang, L. Zhao, R. Zhao, Z. Li, Y. Liu, Y. Duan, Tianyu Han, A mechanochromic luminescent material with aggregation-induced emission: Application for pressure sensing and mapping, Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy 2019, 220 117125.
- [6] Chu T-C, Buras ZJ, Smith MC, Uwagwu AB, Green WH.From benzene to naphthalene: direct measurement of reactions and intermediates of phenyl radicals and acetylene., 2019 PCCP. 21, 22248-22258.
- [7] Yang T, Troy TP, Xu B, et al. Hydrogenabstraction/acetyleneaddition exposed. Angew Chem, Int Ed. 2016, 55, 14983-14987.
- [8] Parker DSN, Kaiser RI, Troy TP, Ahmed M. Hydrogen abstraction/acetylene addition revealed. Angew Chem, Int Ed. 2014, 53, 7740-7744.
- [9] Yu T, Lin MC. Kinetics of phenyl radical reactions studied by the cavity-ring-down method. J AmChemSoc. 1993, 115, 4371-4372.
- [10] Liu P, Li Z, Bennett A, Lin H, Sarathy SM, Roberts WL. The site effect on PAHs formation in HACAbased mass growth process. Combust Flame. 2019, 199, 54-68.
- [11] Mebel AM, Georgievskii Y, Jasper AW, Klippenstein SJ. Temperature- and pressure-dependent rate coefficients for the HACA pathways from benzene to naphthalene. Proc Combust Inst. 2017, **36**, 919-926.
- [12] Tokmakov IV, Lin MC. Reaction of phenyl radicals with acetylene: quantum chemical investigation of

- the mechanism and master equation analysis of the kinetics. J Am Chem Soc. 2003, **125**, 11397-11408.
- [13] Richter H, Mazyar OA, Sumathi R, Green WH, Howard JB, Bozzelli JW. Detailed kinetic study of the growth of small polycyclic aromatic hydrocarbons. 1. 1-naphthyl + ethyne. J Phys Chem A. 2001, **105**, 1561-1573.
- [14] ADF2019, SCM, Theoretical Chemistry, Vrije Universiteit, Amsterdam, The Netherlands, http://www.scm.com
- [15] S. H. Wen, A. Li, J. L. Song, W. Q. Deng, K. L. Han, W. A. Goddard, First-Principles Investigation of Anistropic Hole Mobilities in Organic Semiconductors, J. Phys. Chem. B 2009, 113, 8813.
- [16] S. Chai, S. H. Wen, J. D. Huang, K. L. Han, Density functional theory study on electron and hole transport properties of organic pentacene derivatives with electron-withdrawing substituent, J. Comput. Chem., 2011, 32, 3218.
- [17] W. Q. Deng, L. Sun, J. D. Huang, S. Chai, S. H. Wen, K. L. Han, Quantitative prediction of charge mobilities of π-stacked systems by first-principles simulation, Nat. Protoc. 2015, 10, 632.
- [18] J. D. Huang, S. H. Wen, K. L. Han, First-principles investigation of the electronic and conducting

- properties of oligothienoacenes and their derivatives, Chem.-Asian J. 2012, 7, 1032.
- [19] Y. Wen and Y. Liu, Recent Progress in n-Channel Organic Thin-Film Transistors, Adv. Mater., 2010, 22 12.
- [20] L. Wang, B. Xu, J. Zhang, Y. Dong, S. Wen, H. Zhang and W. Tian, Theoretical investigation of electronic structure and charge transport property of 9,10-distyrylanthracene (DSA) derivatives with high solid-state luminescent efficiency, Phys. Chem. Chem. Phys. 2013, 7, 15.
- [21] García, G., Moral M., Garzón A., Granadino-Roldán J.M., Navarro, A., Fernández-Gómez M., Poly(arylenethynyl-thienoacenes) as candidates for organic semiconducting materials. A DFT insight, Org Electron, 2012, 13 12.
- [22] Li, Y., Zou, L.Y., Ren, A.M. and Feng, J.K., Theoretical study on the electronic structures and photophysical properties of a series of dithienylbenzothiazole derivatives, Comput. Theor. Chem., 2012, **981**, 14.
- [23] Rohloff, R., Kotadiya, N.B., Craciun, N.I., Blom, Wetzelaer, G.A.H., Electron and hole transport in the organic small molecule α-NPD, Appl. Phys. Lett. 2017, 110, 073301.