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#### MS15 P13

Structures of Some Hydroxysalicylaldehyde Schiff Bases <u>Arzu Ozek <sup>a</sup></u>, Çigdem Albayrak<sup>°</sup>, Mustafa OdabayghP, nul Odorat Büylikgüngdr<sup>'</sup> "OndoAu-Mayu Unie, Deportment of Physics, Samsun-Turkey. <sup>°</sup>Ondokuz Mays Univ., Department of Chemistry Samsun-Turkey E-mail :arzuozek@omu.edu.tr

## Keywords: tautomerism, hydrogen bonds, diffraction structure Analysis

The *molecules* of lhe three compounds,  $C_{14}H_{12}BrN_1O_2$  with the orto, meta and para positions of Br (I,II and III, respectively), adopt the phenol-imine taulomeric form with strong intmmoleculm 0-H ..N hydrogen bonds.



In the componds, the phenol-imine tautomer, is favored over the keto-amine haut, as indicated by the C6-01, C8-N I, CI-C8, and CIC6 bond lengths. A similer situation was observer] for 2-(3-methoxysalicylideneamino)-IHbenzimidazole monohydrate (I)- ihe OI-C6 bond lengths are approximately the sent, indicating single-bond, hantent, whereas lhe C8-NI bond ]coprins are indicative of significant deublebond character in (I), (II) and (III).

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#### **MS15 P14**

A Preference for the Thione Structure – Lithium 2-Thiooxo-1,2-dihydropyridine-1-olate Ethanol Solvate Ingrid Svoboda.<sup>a\*</sup> Hartmut Fuess,<sup>a</sup> Nina Schneiders, and Jens Hartung<sup>b</sup> <sup>a</sup>Department of Material Science TU Darmstadt, Germany. <sup>b</sup>Department of Organic Chemistry, Kaiserslautern, Germany. E-mail: svoboda@tu-darmstadt.de

# Keywords: Thiol Thione Tautormerism, Lithium Compound, Thiohydroxamate, Pyridinethione.

2-Thiooxo-1,2-dihydropyridine-1-olate is an ambident nucleophile that is preferentially alkylated at sulfur in the presence of hard countercations, such as Na<sup>+</sup> [1]. The reactivity of the title compound, however, does not fit into this general scheme. Its inherent low reactivity toward strong electrophiles in association with a slight preference for the O-alkylation prompted us to explore its solid state structure at 300 K. Diffraction experiments performed at 100 K and 150 K surprisingly did not afford data sets of an improved quality. The compound crystallizes in  $P2_1/c$  (Z = 4). Two infinite chains of Li and O give rise to  $Li_2O_2$ rhombi, which are tilted by approximately 90 ° toward one another in an accordion-like manner. Two modes of Li binding are seen in the crystal. In the first, Li is surrounded by two S- and three O-atoms leading to distorted trigonal bipyramidal coordination sphere. In the second, Li binding of four O-atoms in a distorted tetrahedral manner is seen. One corner of this tetrahedron is occupied by an ethanol solvate molecule. The structure of the 2-thiooxo-1,2-dihydropyridine-1-olate entity of the molecule is distinctively different from the one reported for the heterocyclic cores of 2-alkylsulfanyl pyridine-1-oxides [2], bis[2-thiooxo-1,2-dihydropyridine-1-olato]nickel [3] and the corresponding zink complex [4]. The correlation of data with those reported for *N*-alkoxypyridine-2(1*H*)-thiones, on the other hand, fits much closer thus pointing to a preference for the thione formula of the title compound in the solid state [5].

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#### MS15 P15

Synthesis, Structural Characterization, Electrochemical, Catalytic, Antimicrobial and Thermal Properties of the Polymeric Metal Complexes Özlem <u>Yılmaz</u><sup>a</sup>, Ertan Şahin<sup>b</sup>, Mehmet Tümer<sup>a</sup>, Mehmet Aslantas<sup>c</sup>, <sup>a</sup>Chemistry Department, Faculty of Arts and Sciences, University of KSU, 46100, Kahramanmaras, TURKEY. <sup>b</sup>Department of Chemistry, Faculty of Arts and Sciences, Ataturtk University, 25240 Erzurum, TURKEY. <sup>c</sup>Physics Department, Faculty of Arts and Sciences, University of KSU, 46100, Kahramanmaras, Turkey. E-mail: <u>mtumer@ksu.edu.tr</u>

## Keywords: X-ray crystallography, electrochemical, catalytic reactions

In this  $4 - \{(E) - [(4 - \{[(1E) - (4 - \{(E) - (E) - (E$ study. [(aminophenyl)imino]methyl}phenyl)methylene]amino}p henyl)imino]methyl}ben-zaldehyde (L) ligand was prepared from the reaction between terephthalaldehyde with 1,4-di-aminobenzene and then its metal complexes were obtained. After the oxidation reactions, in the solvent mixture, the oxidation product 3,3'-5,5'-tetra-tert-butyl-4,4'diphenoquinone (TTBDQ) was obtained as a single crystal. The oxidation product (TTBDQ), C<sub>28</sub>H<sub>40</sub>O<sub>2</sub>, crystallizes in the space group P-1 with one-half molecule in the asymmetric unit and the other half generated by an inversion centre. The diphenoquinone moiety is planar within  $\pm 0.016(3)$ Å. The thermal studies for the ligand and its complexes studied. In addition, the electrochemical and antimicrobial properties of the compounds were investigated.

### MS15 P16

**Cocrystals of oxalic acid with tricyclic quinazolone derivatives.**<u>Akmal Tojiboev</u><sup>a</sup>, Kambarali Turgunov<sup>a</sup>, Bahodir Tashkhodjaev<sup>a</sup>. <sup>a</sup>*S.Yunusov Institute of Chemistry of Plant Substances, Tashkent, Uzbekistan.* E-mail: <u>a tojiboev@yahoo.com</u>

### Keywords: crystal engineering, cocrystals, quinazolone

The design of organic solids by crystal engineering is presently of high interest [1]. Tricyclic quinazoline derivatives are potentially valuable synthons in crystal engineering: their electron donating nitrogen atom holds