Electrochemical Acetoxylation Of Nitrobenzene At Graphite And Platinum Electrodes

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Abstract:- Anodic acetoxylation of nitrobenzene has been carried out on graphite and platinum electrodes in glacial acetic acid and sodium acetate. On graphite electrode p-acetoxy nitrobenzene is the major product along with o-acetoxynitrobenzene in minor amount and dimer in traces. By using platinum electrode, the major product is dimer along with p-acetoxynitrobenzene in traces. The most probable mechanism has been proposed.

Keywords:- Anodic acetoxylation, Nitrobenzene, Graphite and platinum electrodes.

I INTRODUCTION

Anodic acetoxylation of aromatic compounds in glacial acetic acid using graphite and boron doped diamond have been studied by Fankhauser and his Co-workers¹. A number of other workers hold similar such views²⁻⁴. In the present work, anodic acetoxylation of nitrobenzene has been carried out on graphite and platinum electrodes in glacial acetic acid and sodium acetate.

Materials

II EXPERIMENTAL

Nitrobenzene(BDH) was distilled under vaccum at 11mmof Hg.The purity was checked by physical parameters and confirmed by TLC.

III ELECTROCHEMICAL PROCESS

The anodic acetoxylation reaction was carried out galvanostatically in a divided cell and locally fabricated 5v-100mA regulated DC power supply. The potentials were measured as cell voltages. Current voltage studies were performed with graphite and platinum anode in glacial acetic acid and sodium acetate.

The anolyte contained 0.1M substrate in 5% glacial acetic acid 20% sodium acetate mixture. Assuming 2electron process to occur depending on the intensity of current realized at the given potential, the duration of current passed was fixed whenever current decrease was unduly large either the polarity was reversed or potential was taken to zero volt for a short period as practiced in established electro synthetic works ^{5,6}.

IV PRODUCT ANALYSIS

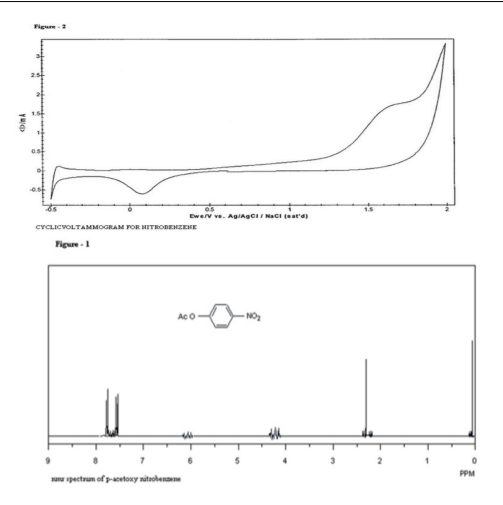
The organic product mixture was separated by ether extraction and resolved on TLC test plate, using 10:1 benzene-ether mixture as elutent. The products were identified by means of co-TLC, physical constants, chemical analysis and pmr spectral analysis.

V RESULTS AND DISCUSSION

The current voltage curves for the system suggests, the working potential range on graphite anode was observed to be 1.5v to 1.6v. The working potential range on platinum anode was observed to be 1.7v to 1.85v.By using graphite electrode, p-acetoxy nitrobenzene as the major product along with o-acetoxynitrobenzene in minor amount and dimer in traces. On platinum electrode, dimer as the major product along with p-acetoxy nitrobenzene in traces.

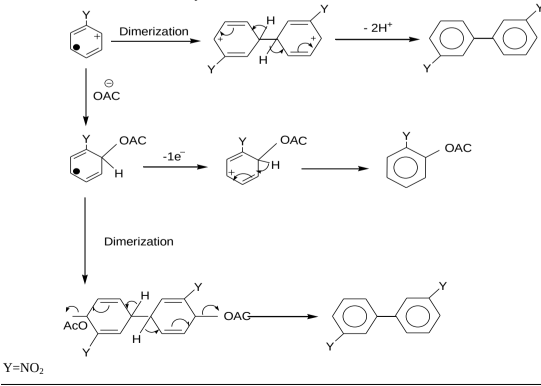
In anodic acetoxylation reactions, graphite anode facilitates adsorption controlled phenomenon, on platinum anode diffusion controlled process occurs. By using graphite or platinum loss of 1 electron occur from the system initially to form radical cation.

On graphite anode, radical cation gets chemisorbed and it is induced by nucleophilic attack of OAC⁻. This results in nucleophilic addition at para position By using platinum anode different situation arises As soon as radical ion is formed it gets diffused into the bulk solution away from the electrical double layer. The labile radical ion undergo dimerisation and it might be exposed to nucleophilic attack by OAC⁻ in the solution.

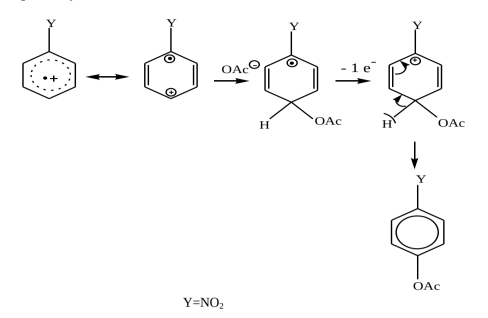


The following mechanism has been proposed for acetoxylation of nitrobenzene At platinum electrode.

Formation of dimer and o-acetoxy nitrobenzene.



At Graphite Electrode. Formation of p-acetoxynitrobenzene.



Such a multiplex approach finds references in the earlier literature. On platinum the attack of the neutral substrate molecule on the cation radical to yield the dimerized product has been favorably proposed^{7-8.}

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