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Electrochemical characterization of hydroquinone derivatives with different substituents in acetonitrile

Ricardo Salazar,*a Jorge Vidal,b Maximiliano Martinez-Cifuentes,b Ramiro Araya-Maturanaa and Oney Ramirez-Rodriguez*c

The effect of carbonyl groups in the ortho position with respect to a hydroxyl group on the electrochemical oxidation of hydroquinones in acetonitrile is studied. The electrochemical response of hydroquinone on a glassy carbon electrode in 0.1 M tetrabutylammonium perchlorate was investigated in detail by voltammetry and coulometry. From these experiments, the oxidation potential was shifted to more positive values with respect to hydroquinone due to the presence of electron withdrawing groups bonded to the aromatic ring. For all compounds a diffusional behavior was observed, and the diffusion coefficient (D) of substituted hydroquinones was calculated showing higher values than found for unsubstituted hydroquinone. Theoretical calculations were carried out to gain insights into the intramolecular hydrogen bond present in these molecules affecting their electrochemical behavior. Relevant theoretical data are optimized geometrical parameters, HOMO energy, condensed radical Fukui functions (f), natural charges, Wiberg bond orders (WBO), stabilization energies caused by electron transfer, and hyperconjugation stabilization energies from the NBO analysis. In most cases, the calculations show good agreement with experimental 1H-NMR data and support the electrochemical results.

1. Introduction

The o- and p-dihydroxybenzene moieties are widely distributed in nature. They are oxidized, generally under mild conditions, to yield quinones. Cells use this type of reaction to transport an electron pair from one substance to another, for example, in mitochondrial ATP synthesis and in photosynthesis.1,2 For this electron pair from one substance to another, for example, in to yield quinones. Cells use this type of reaction to transport an

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A correlation between 1H-NMR chemical shift and IR frequency as indicators of H-bond strength, and the quinone half-wave reduction potential has been described, showing that the strong hydrogen bonds make the reduction of quinones easier.32 Many other authors have published results of hydroquinones oxidation or quinone reduction having intramolecular hydrogen bonds. In all cases it has been shown that the formation of intramolecular hydrogen bonds causes a substantial variation of the half-wave potential.31–34

The electro-oxidation of phenols and derivatives has a very complex mechanism, as shown in Scheme 1. There are many species related by electron and proton transfer which occurs as a consequence of bimolecular interactions and primary electrode processes. The experimental variables have proven to be crucial in the predominance of one species over another.35

The possible products formed from the phenoxonium ion (IV) (Scheme 2) can be divided into two groups, depending on the reaction pathway. If the substituent is OH, ortho- or

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The hyperconjugation interaction between lone pair (LP) on acceptor oxygen and sigma antibonding on donor H–O (LPO → σ*H–O') in the O–···H–O' complex has been described as a major contribution to hydrogen bond interaction obtained by NBO analysis. With second-order perturbation theory analysis, the donor-acceptor interaction (stabilization energy) can be calculated.

In this paper we study the effect of carbonyl groups in the ortho position with respect to a hydroxyl group on the electrochemical oxidation of hydroquinones in acetonitrile. The electrochemical response of hydroquinone on a glassy carbon electrode in 0.1 M tetrabutylammonium perchlorate was investigated in detail by voltammetry and coulometry. From these experiments, we determined the influence of the substituent on the electrochemical oxidation peak, the transferred electron numbers, the diffusional process and diffusion coefficient (D) of hydroquinones. The oxidation potentials were correlated with theoretical parameters.

2. Experimental

2.1. Hydroquinones

Hydroquinone (HQ), 2',5'-dihydroxyacetophenone (HQ1), 2,5-dihydroxybenzoic acid (HQ2), 2',4'-dihydroxyacetophenone (HQ6) and 3,4-dihydroxybenzoic acid (HQ8) are commercially available and were used without further purification. Ethyl 2,5-dihydroxybenzoate (HQ3) was synthesized by Fischer esterification, using HQ2 and ethanol as reagents and sulfuric acid as a catalyst; their physical constants agree with those reported in the literature. 2-Bromo-1-(2,5-dihydroxyphenyl)ethanone (HQ4) and 1-(2-chloro-3,6-dihydroxyphenyl)ethanone (HQ7) were synthesized by described procedures. The chemical structures of all the studied hydroquinones are shown in Fig. 1.

Synthesized compounds. 1-(2,5-Dihydroxyphenyl)-2-iodoethanone (HQ5)

Four hundred milligrams (1.73 mmol) of 2-bromo-1-(2,5-dihydroxyphenyl)ethanone (HQ4) were dissolved in 50 mL of acetone and then 649 mg (4.3 mmol) of sodium iodide dissolved in 20 mL of acetone was added. The reaction mixture was stirred for 2 hours and was then heated under reflux for 30 minutes. The solvent was removed under vacuum and the residue was purified by flash column chromatography on silica gel using a 2 : 1 mixture of hexane–ethyl acetate. Yellow solid. Yield: 80%. MP: 120.5–121 °C. IR (KBr, cm⁻¹) 3287, 3221, 3179, 1642, 1618, 1569, 1476, 1425, 1367, 1309, 1263, 1209, 1086, 1011, 924, 833, 829, 789, 647. 1H-NMR (CDCl3, 300 MHz) δH 4.32 (s, 2H, CH2), 4.75 (s, 1H, HO–C5), 7.08 (dd, J = 2.9, 8.9 Hz, 1H, H–C4), 7.18 (d, J = 2.9 Hz, 1H, H–C6), 11.46 (s, 1H, HO–C2). 13C-NMR (75 MHz, DMSO-d6) δC 67.5, 115.32, 118.54, 118.64, 124.49, 149.51, 153.21, 197.27. HRMS (EI, 70 eV) M+ m/z 277.94423 – calculated for C10H7IO3: 277.94399.

2.2. Electrochemical experiments

2.2.1. Electrolytic medium: acetonitrile containing 0.1 M tetrabutylammonium perchlorate (TBAP). The working concentrations of each hydroquinone varied between 0.1 mM and 2 mM.
2.2.2. Differential pulse voltammetry (dpv), cyclic voltammetry (cv) and linear sweep voltammetry (lsv). They were carried out with a CH Instrument 760-C electrochemical work station. All the voltammetric experiments were carried out with 1.0 mM solutions of each hydroquinone. A stationary glassy carbon electrode (GCE, CH Instrument, with an area of 0.0707 cm²) was used as a working electrode for the dpv and cv experiments. For hydrodynamic experiments, a rotating disk glassy carbon electrode was employed (CH Instrument, with an area of 0.0707 cm²). The surface of the disk was polished to a mirror finish with 0.1 μm alumina powder before use and after each measurement. Platinum wire was used as an auxiliary electrode, and all potentials were measured against a nonaqueous Ag/Ag⁺ reference electrode CH Instrument 112. For all experiments, the resistance was compensated automatically.

2.2.3. Coulometric analyses. Studies on exhaustive electrolysis were carried out for two hours at constant electrode potential (+0.2 V peak potential found by dpv showed in Table 1) in a divided cell, on a glassy carbon mesh electrode using 30 mL of 1 × 10⁻⁵ M solutions of the compounds. A three-electrode circuit with a reference non aqueous Ag/Ag⁺ and platinum wire as a counter electrode, and all potentials were measured against a nonaqueous Ag/AgCl sat) reference electrode CH Instrument 112. For all experiments, the resistance was compensated automatically.

2.2.4. Characterization of the synthesized compounds. Melting points were uncorrected and measured on a Büchi SMP-20 or a Gallenkamp melting point apparatus. Infrared spectra were recorded on a Nicolet 510P FT-IR spectrophotometer for KBr discs, and the frequencies are given in cm⁻¹. NMR spectra were obtained on a Bruker AVANCE DRX 300 instrument at 300.13 and 75.5 MHz for 1H and 13C NMR, respectively. Chemical shifts (δ) are reported in parts per million downfield from TMS for 1H NMR, or relative to residual solvent signals (CHCl₃, 7.26 ppm for 1H NMR, DMSO-d₆, 39.52 ppm for 13C NMR spectra). 13C NMR spectra were acquired on a broad-band decoupled mode. Silica gel 60 (230–400 mesh ASTM) and DC-Alufolien 60 F₂₅₄ were used for flash-column chromatography and analytical TLC, respectively.

2.2.5. Theoretical calculations. The calculations were carried out using the Gaussian 03 program package. Geometries were optimized at the B3LYP/6-311++G** level for C, H, O, Cl, and Br, and at the B3LYP/SDB-cc-pVTZ level for I. No imaginary frequencies were found at the optimized molecular geometries, indicating that they are real minima of the potential energy surface. NBO calculation was carried out with the 6-311G** basis set to avoid the problems associated with diffuse functions in this kind of calculation.

3. Results and discussion

3.1. Differential pulse voltammetry (dpv) results

Dpv results reveal that HQ and HQ derivatives exhibited one well-defined anodic peak at potentials higher than +0.8 V versus Ag/AgCl sat) (Fig. 2 and Table 1). Oxidation peak potential values of HQ derivatives were shifted towards more positive values compared with HQ (Eₚ, HQ = +0.88 V). The results of the oxidations of HQ and HQ derivatives are shown in Fig. 2 and Table 1.

Oxidation of these hydroquinones is dependent on two factors: the electronic effects of substituents on the aromatic ring, and the formation of an intramolecular hydrogen bond (IHB) between a hydroxyl group and a substituent in the ortho-position (Fig. 1). The oxidation potential of the substituted HQs is increased compared to HQ due to the presence of electron withdrawing groups bonded to the aromatic ring, because they decrease its electron density as well as that on the hydroxyl.

![Chemical structures of the investigated compounds.](image-url)
oxygen atom, hindering the loss of one electron from it. Additionally, the potential depends on the strength of the intramolecular hydrogen bond of the phenolic hydroxyl. A strong hydrogen bond facilitates oxidation because it increases the electron density on the oxygen atom, facilitating the transfer of electrons.

To explore and quantify both the effects of a substituent on the ring and the IHB, and establish their relationship with the oxidation potential we used DFT calculations (Table 2).

3.2. DFT calculations

The optimized geometric parameters, HOMO energy and condensed radical Fukui functions \( f^+ \) are summarized in Table 3, respectively. The natural charges \( q \) and the Wiberg bond order (WBO) from the NBO analysis for the optimized geometries from HQ to HQ8 are presented in Tables 4 and 5, respectively. The stabilization energies for selected NBO donor–acceptor pairs in the HQs, given by second order perturbation energies of the Fock matrix in the NBO basis, are presented in Table 6. According to these results, substituents on the aromatic ring, for HQ1 to HQ8, increase the O1–H1 distance compared to HQ. The differences in the O3···H1 distances and in the O1–H1–O3 angles between HQs (Table 3) are indicative of differences in the IHB strength. On the other hand, the condensed radical Fukui function indicates the most susceptible site for a radical attack leading to a loss or a gain of an electron. With this criterion we found that, for all HQs, O1 is the more reactive center for electron abstraction. Besides, the reduction of the natural charge on O1 should increase their oxidation potential compared to HQ. Our results show that HQ1 to HQ7 show a reduced natural charge on O1 compared with HQ, in agreement with the experimental data. On the other hand, the WBO reflect the change caused by aromatic ring substituents on the intramolecular hydrogen bond, which also affects the ability of the HQs to transfer electrons. The WBO for O3···H1 changes from HQ1 to HQ7 depending on the nature of the carbonyl group and the aromatic ring substituents.

Compounds HQ1–3 show a similar \( E_p \) (range ±8 mV), and these three molecules show an electron- withdrawing group (acetyl, carboxyl and ethoxycarbonyl, respectively) in the ortho-position with respect to a phenol function (Fig. 1). The presence of these groups causes a decrease of the electron density on O1, as reflected in their natural charge for HQ1–3 compared with HQ, and hence making them more difficult to oxidize. The oxidation potential is shifted about 140–150 mV higher than that of HQ.

The IHB O1–H1···O3 with acetyl in HQ1 is stronger than with carboxyl in HQ2 and with ethoxycarbonyl in HQ3, as reflected in the shorter distance and higher WBO for O3···H1 in HQ1 compared to HQ2 and HQ3. The latter is also reflected by hyperconjugative interaction between LPs of O3 and \( \sigma^* \) of O1–H1, with the stabilization energy higher for HQ1 (22.53 kcal mol\(^{-1}\)) compared to HQ2 (15.78 kcal mol\(^{-1}\)) and HQ3 (16.73 kcal mol\(^{-1}\)), indicating a strong IHB for the first case. These results are consistent with a higher \(^1\)H-NMR chemical shift for H1 in HQ1 (\( \delta_{H1} 11.81 \) ppm) compared with HQ3 (\( \delta_{H1} 10.48 \) ppm), and this greater deshielding of H1 in HQ1 also reveals a strong IHB. For this reason, taking into account the IHB strength, HQ1 should be oxidized more easily. However, it is possible also to argue that the greater strength of the hydrogen bond of the keto group, which facilitates oxidation, is counteracted by its greater electron- withdrawing effect, which hinders the transfer of electrons from the hydroxyl oxygen atom. The latter is supported by a higher stabilization energy for the LPs of O1 and \( \sigma^* \) and \( \pi^* \) for C1–C\( _2 \), which indicate the degree of delocalization of O1 on the aromatic ring. For HQ1 LP of O1 and \( \sigma^* \) and \( \pi^* \) for C1–C\( _2 \) interact with a stabilization energy of 43.59 kcal mol\(^{-1}\), while for HQ2 and HQ3 the stabilization energies are 42.72 and 42.08 kcal mol\(^{-1}\). The presence of these opposite operating effects explains the very similar potential of the three compounds.

On the other hand, HQ4 and HQ5 have an acetyl group the same as HQ1, but additionally, they possess a halogen atom in an \( \alpha \)-position to the carbonyl group. HQ4 has a bromine atom and HQ5 an iodine atom instead (Fig. 1). They show a difference of approximately 50 mV with respect to HQ1 (Table 1). A plausible explanation takes into account the differential electron withdrawing effect of the halogen atoms, due to their different electronegativities, on the carbonyl group. A higher electron density on the carbonyl group causes a strengthening of the IHB. The latter is reflected by the differences in the distance and WBO of O3···H1, because in HQ1 \( d_{O3 \cdot \cdot \cdot H1} \) is 1.696 Å and WBO is 0.076, while in HQ4 and HQ5 \( d_{O3 \cdot \cdot \cdot H1} \) it increases to 1.711 Å and 1.707 Å, and the WBO is reduced to 0.072 and 0.070, respectively, all of which indicates a weakening of the IHB in HQ4 and HQ5. Accordingly, hyperconjugative interaction between LPs for O3 and \( \sigma^* \) for O1–H1 indicates a weaker IHB for HQ4 and HQ5 compared with HQ1. The chemical shift of H1 is 11.81 ppm for HQ1, 11.35 ppm for HQ4 and 11.46 ppm for HQ5, showing the weakening of the IHB, in agreement with the theoretical parameters. Fig. 3 shows a good correlation between the chemical shift for chelated hydrogen (H1) and the Wiberg bond order (WBO) O3···H1. Moreover, the electron-withdrawing effect of \(\alpha\)-haloacetyl groups is greater than that of the acetyl group, which is reflected by an increase of the stabilization energy for delocalization of O1 into C1–C\( _2 \),
Table 2  The optimized structures, HOMO and LUMO orbitals of the investigated HQs

<table>
<thead>
<tr>
<th>Optimized structure</th>
<th>HOMO</th>
<th>LUMO</th>
</tr>
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<tbody>
<tr>
<td>HQ</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HQ1</td>
<td></td>
<td></td>
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<tr>
<td>HQ2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HQ3</td>
<td></td>
<td></td>
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<tr>
<td>HQ4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HQ5</td>
<td></td>
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<tr>
<td>HQ6</td>
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</tbody>
</table>

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The electronegative chlorine atom bonded in the ortho-position to the carbonyl group in HQ7 reduces the natural charge on oxygen atom 1 and increases the WBO of O3/C1/C1/C1H1. Moreover, the electronic delocalization from chlorine to the acetyl group causes an increased hydrogen bond strength compared with HQ1, a evidence db yt he smaller stabilization energy of the LP of O3 interaction and σ* of O1–H1. All these factors explain why the oxidation potential of HQ7 is 0.11 V lower than that of HQ1.

HQ1 and HQ2 are isomers of HQ6 and HQ8, respectively. HQ1 and HQ2 are p-hydroquinones, while HQ6 is a resorcinol and HQ8 is a catechol (o-hydroquinone). The IHB is stronger in HQ6 compared to HQ1, according to the O3···H1 distance and the WBO. Stabilization energy for hyperconjugative interaction also indicates a strong IHB in HQ6. The strong electronic delocalization from the hydroxyl group located in the para-position from 31.87 kcal mol$^{-1}$ for HQ to 44.46 kcal mol$^{-1}$ and 44.42 kcal mol$^{-1}$ for HQ4 and HQ5, respectively. Therefore, the potential increases as a result of the decrease in the electron density of the aromatic ring. It is also reflected by an increase of the natural charge on O1.

Table 2 (continued)

<table>
<thead>
<tr>
<th>Optimized structure</th>
<th>HOMO</th>
<th>LUMO</th>
</tr>
</thead>
<tbody>
<tr>
<td>HQ7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HQ8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* B3LYP/6-311++G(d,p) is used for C, H, O, Cl and Br atoms. SDB-cc-pVTZ is used for I atom.

Table 3 Geometric parameters and condensed radical Fukui functions

<table>
<thead>
<tr>
<th></th>
<th>d$_{O3}$</th>
<th>d$_{O1-H1}$</th>
<th>L$_{O3-H1-O3}$</th>
<th>E$_{HOMO}$</th>
<th>f$_{O1}$</th>
<th>f$_{O2}$</th>
<th>E$_{p}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HQ</td>
<td>0.962</td>
<td></td>
<td>0.21647</td>
<td>7.50</td>
<td>7.50</td>
<td>0.880</td>
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</tr>
<tr>
<td>HQ1</td>
<td>1.696</td>
<td>0.986</td>
<td>146.310</td>
<td>2.2618</td>
<td>9.18</td>
<td>6.96</td>
<td>1.028</td>
</tr>
<tr>
<td>HQ2</td>
<td>1.774</td>
<td>0.980</td>
<td>144.449</td>
<td>2.2814</td>
<td>8.73</td>
<td>7.35</td>
<td>1.024</td>
</tr>
<tr>
<td>HQ3</td>
<td>1.760</td>
<td>0.981</td>
<td>145.066</td>
<td>2.2206</td>
<td>8.84</td>
<td>7.18</td>
<td>1.032</td>
</tr>
<tr>
<td>HQ4</td>
<td>1.711</td>
<td>0.984</td>
<td>145.361</td>
<td>2.3478</td>
<td>9.13</td>
<td>7.09</td>
<td>1.080</td>
</tr>
<tr>
<td>HQ5</td>
<td>1.707</td>
<td>0.984</td>
<td>145.397</td>
<td>2.3378</td>
<td>9.13</td>
<td>7.10</td>
<td>1.076</td>
</tr>
<tr>
<td>HQ6</td>
<td>1.669</td>
<td>0.991</td>
<td>147.646</td>
<td>2.4226</td>
<td>8.64</td>
<td>3.70</td>
<td>1.632</td>
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<tr>
<td>HQ7</td>
<td>1.591</td>
<td>0.993</td>
<td>147.341</td>
<td>2.3160</td>
<td>9.16</td>
<td>6.87</td>
<td>0.920</td>
</tr>
<tr>
<td>HQ8</td>
<td>2.156</td>
<td>0.966</td>
<td>122.717</td>
<td>2.3982</td>
<td>9.07</td>
<td>6.27</td>
<td>1.064</td>
</tr>
</tbody>
</table>

* Distances in Angström; angles in degrees, energies in a.u.; potentials ($E_p$) in volts; $f_{O1}$ radical Fukui function.

Table 4 Natural charges of the selected atoms in HQ compounds

<table>
<thead>
<tr>
<th></th>
<th>HQ</th>
<th>HQ1</th>
<th>HQ2</th>
<th>HQ3</th>
<th>HQ4</th>
<th>HQ5</th>
<th>HQ6</th>
<th>HQ7</th>
<th>HQ8</th>
</tr>
</thead>
<tbody>
<tr>
<td>O1</td>
<td>-0.678</td>
<td>-0.672</td>
<td>-0.670</td>
<td>-0.673</td>
<td>-0.662</td>
<td>-0.663</td>
<td>-0.665</td>
<td>-0.661</td>
<td>-0.666</td>
</tr>
<tr>
<td>H1</td>
<td>0.459</td>
<td>0.502</td>
<td>0.494</td>
<td>0.493</td>
<td>0.494</td>
<td>0.494</td>
<td>0.493</td>
<td>0.491</td>
<td>0.477</td>
</tr>
<tr>
<td>O2</td>
<td>-0.677</td>
<td>-0.675</td>
<td>-0.677</td>
<td>-0.677</td>
<td>-0.674</td>
<td>-0.674</td>
<td>-0.657</td>
<td>-0.672</td>
<td>-0.696</td>
</tr>
<tr>
<td>H2</td>
<td>0.466</td>
<td>0.462</td>
<td>0.461</td>
<td>0.462</td>
<td>0.462</td>
<td>0.462</td>
<td>0.462</td>
<td>0.474</td>
<td>0.477</td>
</tr>
<tr>
<td>O3</td>
<td>-0.611</td>
<td>-0.651</td>
<td>-0.654</td>
<td>-0.585</td>
<td>-0.591</td>
<td>-0.620</td>
<td>-0.609</td>
<td>-0.609</td>
<td></td>
</tr>
</tbody>
</table>
relative to the acetyl group leads to a large increase in electron density of the carbonyl oxygen, thereby enhancing the hydrogen bond that facilitates oxidation. Despite the above, its reduction potential is greater than that of the other HQs. This can be explained by stabilization of semiquinone radicals from hydroquinones, which is not present in the radical derivate from resorcinols, as is the case of this molecule.

On the other hand, HQ8 exhibits a weak IHB between H1⋯O2, with a distance of 2.156 Å and a WBO of 0.006. Non hyperconjugative stabilization energy was found for this interaction. Although HQ8 has weaker IHB, it does not have a high reduction potential, which can be explained in terms of product stabilization. The p-quinones are more stable than the o-quinones because the carbonyl groups are further apart, thus the electron densities of the two carbon–oxygen π bonds do not affect each other; the carbonyl oxygen atoms, with a high electron density, are not close; and the carbonyl carbon atoms, electron deficient, are not neighbors. The greater stability of catechols compared to hydroquinones, and of p-quinones compared to o-quinones, determines that HQ2 is oxidized more easily than HQ8. Zhu et al. report a comparison of hydride affinities between o-quinones and p-quinones, and demonstrate, using calculation methods, that p-benzoquinone is 5.5 kcal mol$^{-1}$ more stable than o-benzoquinone, while the catechol monoanion is 7.3 kcal mol$^{-1}$ more stable than the hydroquinone monoanion.$^3$

### 3.3. Determination of transferred electrons by coulometric studies

As mentioned above, the oxidation reaction of hydroquinones and catechols in acetonitrile follows an electrochemical–chemical–electrochemical (ECE) sequence under a mixed kinetic control by electron and proton transfer. The determination of the electrons transferred during the oxidation of all HQs (Fig. 1), were based on the exhaustive electrolysis at controlled-potential using a three compartment cell. Glassy carbon mesh was used as a working electrode, a platinum wire as a counter electrode and an Ag/AgCl as a reference electrode. To calculate the number of electrons was considered the sum of the final electric charge ($Q_{\text{corrected}}$) for successive electrolysis. The number of electrons transferred was calculated for each mole of HQ since the overall net charge, and using Faraday equation ($Q_{\text{net}} = n \times F \times e$, where $n$ = number of moles, $F$ = Faraday constant (96 500 C mol$^{-1}$))
and \( e = \text{number of electrons} \). The electrolysis was carried out by applying 100 mV more than the oxidation potential peak obtained by VPD (Table 1). Coulometric studies on the oxidation of each compound revealed an average of 1.9 \( \pm \) 0.1 transferred electrons. Table 1 summarizes peak oxidation potentials and number of electrons obtained for different derivatives in an acetonitrile medium. Thus, an influence by substituent on the transferred electrons in the redox process was not observed.

3.4. Cyclic voltammetry characterization

The electrochemical behavior of HQ and HQ substituted was carefully investigated at a bare GCE in acetonitrile containing 0.1 M tetrabutylammonium perchlorate (TBAP) using cyclic voltammetry. These experiments were carried out at different sweep rates ranging from 0.01 to 1.5 V s\(^{-1}\).

Fig. 4 shows the CV curves of 1.0 mM of HQ2. The separation between the anodic potential peak (\( E_{pa} \)) and the cathodic potential peak (\( E_{pc} \)) is, on the average, higher than 0.45 V for all the compounds studied. So, under these conditions, anodic signals were electrochemically irreversible as can be seen in Fig. 4A, and the peak current enhancements with increased scan rate for all compounds.\(^{51}\)

Fig. 4B shows a linear relationship of \( i_p \) vs. the square root of the scan rate over the whole 0.001–1.5 V s\(^{-1}\) range, strongly suggesting that the redox reactions of HQ and HQ substituted are diffusion-controlled.\(^{52}\) Plots of \( \log i_p \) vs. \( \log v \) had slopes close to 0.5 for both signals, confirming that currents were diffusion-controlled (Fig. 4C). The same analysis was carried out for all hydroquinones showing similar behaviors, a diffusion-controlled redox process. On the other hand, oxidation and reduction potential values were dependent on the sweep rates, supporting the electrochemically irreversible character of the processes of all compounds in this medium.\(^{51}\) Fig. 5A confirms this phenomenon for HQ3 because, in all cases, both anodic and cathodic processes (\( E_{pa}, E_{pc} \) and \( \Delta E_p \)) show good

![Fig. 4](image1.png)

**Fig. 4** (A) Cyclic voltammograms of 1.0 mM HQ2 in acetonitrile + 0.1 M tetrabutylammonium perchlorate at different scan rates. (B) Linear relationship of \( i_p \) vs. \( V^{1/2} \) of graph A. (C) \( \log i \) vs. \( \log v \) for the anodic process of A.

![Fig. 5](image2.png)

**Fig. 5** (A) Cyclic voltammograms of 1 mM of HQ3 in acetonitrile + 0.1 M tetrabutylammonium perchlorate at different scan rates. (B) Linear relationship between \( E_p \) and Ln \( v \) for A. (C) Linear relationship between \( \Delta E_p \) and \( \log v \) for A.
A reasonable explanation of this decrease of the diffusion coefficient, like the rest of the compounds, has one oxidation derivative, like the rest of the compounds, has one oxidation wave which did not show significant differences with the others. This supports the fact that the oxidation process involved a similar number of transferred electrons, and that it was also diffusion-controlled as mentioned before.

The relationship between the limiting currents and the square root of the rotation rate was linear for all compounds (Fig. 6, A), and the diffusion-controlled current at different rotating rates. (B) Levich treatment of A.

Fig. 6 (A) Typical hydrodynamic voltammetry on a glassy carbon rotating disk electrode of HQ6 in acetonitrile + 0.1 M tetrabutylammonium perchlorate at different rotating rates. (B) Levich treatment of A.

3.5. Hydrodynamic voltammetry

Fig. 6 shows typical voltammograms on a graphite rotating disk electrode corresponding to 1.0 mM solutions of HQ5. This derivative, like the rest of the compounds, has one oxidation wave which did not show significant differences with the others. This supports the fact that the oxidation process involved a similar number of transferred electrons, and that it was also diffusion-controlled as mentioned before.

The relationship between the limiting currents and the square root of the rotation rate was linear for all compounds (Fig. 6), in agreement with Levich. From these plots, the diffusion coefficients $D$ were calculated (Table 1) and some differences were found between the derivatives, all of which showed lower $D$ values than HQ: $D_{HQ} = 3.42 \pm 0.05 \times 10^{-5}$ (cm$^2$ s$^{-1}$). A reasonable explanation of this decrease of the $D$ values in HQ derivatives may be attributed to the active presence of substituents on the speed of diffusion to the surface electrode, influenced by a larger molecular volume. These results were confirmed by determination of geometrical parameters and condensed radical Fukui functions. The optimized geometrical parameters, HOMO energy, and condensed Fukui functions are summarized in Table 3. From the calculated results, it is found that a substituent in the aromatic ring increases the O1–H1 distance for HQ1 to HQ8, compared to HQ. The differences between the O3···H1 distances and the O1–H1–O3 angles among the HQs are indicative of differences in the intramolecular hydrogen bond strength, as we mentioned above. These results suggest that the $D$ value depends on both the nature and the position of the substituent on the aromatic ring, also confirming that bigger molecules have smaller diffusion coefficients.

The presence of bromine (HQ4) and iodine (HQ5) atoms in the ortho-position (Fig. 1) increases the molecular volume, decreasing the $D$ value. No significant differences in $D$ values were found when the substituent was a ketone (HQ1), a carboxylic acid (HQ2) or an ester (HQ3). In a decreasing order, coefficients obtained from the hydrodynamic experiments were $D_{HQ} > D_{HQ7} > D_{HQ8} > D_{HQ3} > D_{HQ1} > D_{HQ8} > D_{HQ4} > D_{HQ5} > D_{HQ6}$. The determined $D$ values show a similar tendency as reported by Valencia et al. for quinines and aromatic compounds in acetonitrile.

4. Conclusions

Electrochemical oxidation of studied hydroquinone and hydroquinone derivatives in acetonitrile involves 2-protons and 2-electrons to give the quinone derivative as a final product. The electrochemical process is electrochemically irreversible with a $\Delta F_p$ higher than 0.45 V for all the compounds studied. The presence of substituents on the aromatic ring (carbonyl groups in the ortho position for example) shifted the redox potential to more anodic values respect to HQ because the presence of withdrawing groups decreases the electron density on the hydroxyl oxygen atom. A decrease in the $D$ value with respect to hydroquinone was observed. Differences in the molecular structure of these hydroquinones significantly influence the characteristics of the C–O···H–O IHB present in them, affecting the electrochemical behavior. NBO calculations gave an insight into the electronic characteristics of the IHB and helped us to rationalize the relationship between the substituent on the aromatic ring, the presence of an IHB, and the oxidation potential of the hydroquinones studied in this work.

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References
