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ABSTRACT

An elegant hanging-droplet or meniscus-based setup is proposed to carry out quantitative electrolyses from either an organic (hydroquinone) or an inorganic (permanganate) substrate. These examples validate the concept of using such easily-accessible, fast (1-3 min) and low-cost operating conditions not only for preparative applications (electrosynthesis), but also for pedagogical purposes in minute samples.

KEYWORDS
Water droplets; Electrochemical setup; Complete electrolysis; Low cost; pedagogical application.

HIGHLIGHTS

- The simplest low cost electrochemical cell: Water droplets hanging at the tip of a glass frit are used as vessels to carry out 2-compartment electrolyses
- Complete electrochemical conversions can be achieved within a minute time window
- The methodology is adapted to occasional electrochemical investigation as well as electrochemical teaching.

• GRAPHICAL ABSTRACT
1. Introduction

The miniaturization of reaction vessels is a growing concern in modern preparative and analytical chemistry. Indeed, the obvious substrate economy realized upon downsizing volumes is associated to the development of flow reactors allowing fast, successive synthetic steps [1, 2] as well as multiple analyte determination in microfluidic analyzers [3, 4].

Besides, the quest for smaller reactors may also encompass a more prospective chemistry, viz. fundamental or applied research where inexpensive, accessible and disposable general-purpose setups would allow fast validation of experiments or concepts. The intrinsic complexity of electrochemical experiments (featuring at least two electrodes in continuity in the electrolyte), especially when it comes to electrosynthesis, generally implies specific glassware and cells which become even more complex and expensive for small volumes. We recently established the possibility to perform electrochemical experiments within the simplest and cheapest available reaction vessel, i.e. a hanging droplet [5]. The originality of this work relied on the fact that micro-volume droplets were not “deposited” on a substrate or at the electrode surface, but left suspended at the fritted glass of a bridge compartment containing a conventional reference electrode (SCE). The use of water droplets as electrochemical cells is not a new concept per se. In 1992, Mc Creery et al. deposited droplets on ordered carbon/graphite electrodes to study the dependence of electron transfer kinetics on the surface defects of such electrodes [6]. More recently, a similar droplet-based setup was used by Unwin either to explore the electron transfer kinetics on highly oriented pyrolytic graphite (HOPG) electrodes [7] or to perform Scanning Electrochemical Cell Microscopy (SECCM) on samples contained in droplets or meniscus [8]. A more comprehensive survey on “droplet electrochemistry” was published by Davies et al. who performed voltammetry and electrodeposition at edge plane pyrolytic graphite electrodes within droplets immersed in an immiscible organic electrolyte [9].
In our approach, where the droplet hangs at the tip of a glass frit, the solution was not deposited on a plane surface or electrode, but featured an inverted cell configuration with respect to the above described setups. Such a device is designed both to minimize sample contamination (no other contact than the frit) and to allow electrochemistry to be performed without specific or printed working electrodes. Accordingly, both working and counter electrodes were introduced in the droplet, thus designing a classical 3-electrode setup in microliter-size volume featuring a commercial reference electrode instead of pseudo-reference wire-shape electrodes (e.g. Ag, Ag/AgCl, Pt). We showed that analytical electrochemistry (voltammetry) could be performed within the droplet, as well as short scale preparative electrochemistry with the examples of platinization and electrodeposition of gold nanocrystals onto carbon-fiber electrode surfaces. These results prompted us to go one step further in the achievement of electrolyses within a droplet by performing the complete electrochemical conversion of a solute dissolved in the droplet, therefore featuring a separator between the working and the counter electrodes in order to avoid generator-collector mode electrolysis. In this work, we have explored the possibility to carry out quantitative electrolyses on organic and inorganic species dissolved in water droplets. We notably considered the electrochemical oxidation of hydroquinone and the electrochemical reduction of permanganate in alkaline droplets. Actually, the oxidation of hydroquinone was selected as an example of electrosynthetic application achieved from controlled potential electrolyses using a 3-electrode setup while permanganate conversion in hanging droplets (galvanostatic 2-electrode setup) was proposed as an illustrative experimental entry to an electrochemistry teaching based on quantitative electrochemical transformations.

2. Experimental section

2.1 Procedure

The original electrochemical setup described in our previous paper [5], which only consisted in a vertical glass frit (porosity 4) has been completed by a second U-shaped glass frit to host the reference electrode. All glassware was purchased from Ellipse Labo, then cut or curved to the desired shape by
a glassblower and then fitted with a thread to host a 15 mm-diameter cap. The scheme and pictures of the corresponding setup are shown in Figure 1 for either a 2-electrode setup (hanging droplet) or a 3-electrode setup (meniscus placed between the two frits). Working electrode 2 was a gold square grid (5 mm x 5 mm) and the counter electrode, placed in compartment 3, was a stainless steel grid. Compartments 3 and 4 were carefully filled with the same electrolytes than those of the droplets using 5 mL plastic syringes equipped with needles in order to avoid air bubbles to form in the vicinity of the frit. Once the compartments were filled with electrolyte, a 50 or 75 µL droplet was placed at the bottom of compartment 3 using an Eppendorf pipette fitted with a 200 µL plastic tip. For 3-electrode electrolyses, the U-shaped compartment 4 was lifted up towards the hanging droplet until it finally formed a meniscus 1, the original droplet being immobilized between compartments 3 and 4 (see Figure 1 for a detailed view). The electrochemical monitoring of hydroquinone and benzoquinone concentrations along the electrolysis was carried out by cyclic voltammetry at a homemade Pt disk microelectrode (100 µm diameter, see right photograph in Fig 1) introduced in the droplet. In both potentiostatic and galvanostatic mode electrolyses using potassium chloride 0.1 mol.L⁻¹ or sodium hydroxide 1 mol.L⁻¹ as electrolytes, the potential difference between the working (gold grid) and counter (stainless steel) electrodes was in a 4 to 6 V window (current range 1-5 mA), mostly accounting for the resistance introduced by the glass frit separator. Nevertheless, this potential difference allow galvanostatic electrolyses to be performed with unspecific apparatus, such as a simple power supply instead of a potentiostat.

2.2 Chemicals

For the hydroquinone oxidation experiments, the electrolyte consisted of a 0.1 mol.L⁻¹ solution of potassium chloride (Sigma Aldrich, > 99%) in deionized water, while permanganate reduction was achieved in 1 mol.L⁻¹ sodium hydroxide (Acros organics, > 98.5 %) in deionized water. Hydroquinone (HQ) was purchased from VWR chemicals and freshly recrystallized in deionized water. Benzoquinone
(BQ) was purchased from Sigma Aldrich and recrystallized from petroleum ether. Potassium permanganate was purchased from VWR chemicals (> 99%).

2.3 Instrumentation

All electrolyses and cyclic voltammetry were performed at room temperature using an Autolab potentiostat (PGSTAT 20). UV-Vis spectra were recorded on a Perkin Elmer Lambda 45 spectrophotometer.

![Diagram of the setup used in this work](image)

**Figure 1**

Scheme and photographs of the setup used in this work

3. Results and discussion

3.1 Electrochemical conversion of hydroquinone (HQ) to 1,4-benzoquinone (BQ)

Hydroquinones are ubiquitous compounds which namely enable electron transfer along the photosynthetic chain [10], are used as precursors in the synthesis of quinones [11] or prevent radical polymerization of vinyl monomers [12]. On the other hand, hydroquinones are toxic [13] and their content in wastewater is scrutinized [14, 15]. Therefore, the conversion of hydroquinone (benzene-
1,4-diol, HQ) appeared as a meaningful example to estimate the efficiency of an electrochemical oxidation carried out from a droplet containing this organic water-soluble molecule. The controlled potential electrolysis of HQ to benzoquinone (BQ) was achieved in only 60 s in an aqueous droplet solution of potassium chloride (0.1 mol.L⁻¹) using the 3-electrode setup. The oxidation potential was set at 1.7 V/SCE, i.e. the more positive value to ensure fast electron transfer without significant O₂ or Cl₂ evolution at the gold electrode. Note that under these conditions, as verified in our previous work [5] no water evaporation was observed from the droplet. In Figure 2, The conversion of HQ and formation of BQ were monitored by in situ cyclic voltammetry using a 100 µm Pt microelectrode (the electrolysis was stopped while recording the cyclic voltammograms (CVs)) as shown in Figure 2A and 2B, respectively. Under our conditions (unbuffered aqueous solution), HQ oxidation peak originally at 0.52 V/SCE shifted towards more positive potentials values while electrolysis proceeded.

![Figure 2](image-url)

**Figure 2**

Electrochemical oxidation of hydroquinone HQ (5 mmol.L⁻¹ contained in a 50 µL aqueous 0.1 mol.L⁻¹ KCl droplet) at 1.7 V vs. SCE using the 3-electrode configuration. A : Evolution of the CVs of HQ as a function of the charge (scan rate 0.2 V.s⁻¹). B : Evolution of the CVs of BQ as a function of the charge (scan rate 0.2 V.s⁻¹). C : Experimental (points) and theoretical (for 100% faradaic efficiency, solid line) concentration of BQ formed as a function of charge – experimental concentration estimated from the calibration of the plateau reduction currents of BQ (like in panel B) with an authentic sample of BQ (5 mmol.L⁻¹). D: UV-vis spectra of HQ (5 mM) before (blue curve) and after (green curve) electrolysis of
the droplet (47 mC). The black curve is the UV-vis spectra of an authentic BQ sample performed at a concentration (5 mM) corresponding to a 100% conversion of HQ.

The complex electrochemistry of the hydroquinone/quinone redox couple is well documented [16, 17], even in unbuffered solutions [18] but mechanistic considerations related to the pH dependence of hydroquinone electrooxidation were out of the scope of this work. Actually, HQ oxidation did not provide a linear relationship between the peak current and HQ concentration, while BQ displayed a stable and linear plateau-shaped reduction wave at -0.48 V/SCE which allowed its concentration to be monitored by cyclic voltammetry (Figure 2B and 2C). The nearly quantitative conversion of HQ to BQ during electrolysis was assessed by the overlay of experimental and expected faradaic formation of BQ (Figure 2C) and by the UV-vis spectra recorded before and after electrolysis (Figure 2D). This confirms that no secondary electrode reactions (like oxygen evolution) were not observed in our conditions. Besides, comparison between the UV-vis spectra obtained after electrolysis and the one recorded for an authentic BQ sample at a concentration of 5 mM (i.e., the same concentration used for the starting HQ) confirmed the full conversion of HQ into BQ (Figure 2D). Moreover, even if HQ did not exhibit a stable oxidation peak in our conditions, the evolution of the CVs in Figure 1A were consistent with the conversion of HQ.

3.2 Electrochemical reduction of permanganate in alkaline aqueous solutions

The objective of this work was also to test a qualitative and visual electrochemical transformation that could be performed in water droplets. Actually, manganese high oxidation states are very interesting with this respect, especially in alkaline media where permanganate MnO$_4^-$ (purple), manganate MnO$_{4}^{2-}$ (green) and even hypomanganate MnO$_{4}^{3-}$ (grey-blue) ions can be formed. Such transformations are currently realized by undergraduate students in test tubes [19] and the corresponding oxidation states described structurally and thermodynamically [20, 21]. Since i) the standard potential values of the MnO$_4^-$/MnO$_{4}^{2-}$ and MnO$_{4}^{2-}$/MnO$_{4}^{3-}$ in alkaline solutions (0.558 V/SHE and 0.285 V/SHE respectively [22]) establish the thermodynamic stability of the intermediate oxidation state and ii) the electrochemical successive generation of MnO$_{4}^{2-}$ and MnO$_{4}^{3-}$ from MnO$_4^-$ has been described at the analytical scale (cyclic voltammetry) in 1 mol.L$^{-1}$ NaOH aqueous solutions [23] the quantitative electrochemical generation of Mn (VI) and Mn (V) anions from an alkaline permanganate droplet could be successful. In order to make the electrolysis as simple as possible, we used the 2-electrode setup with the hanging droplet under galvanostatic conditions (no reference electrode connected). The results of the one- and two-electron reduction of a 10$^{-2}$ mol.L$^{-1}$ potassium permanganate droplet in aqueous 1 mol.L$^{-1}$ sodium hydroxide solution are presented in Figure 3. The color evolution could not
be directly observed during electrolysis (under conventional lighting the droplet was too dark), but was very apparent once the droplet diluted (Figure 3A). Both the color and the UV-vis spectrum (maximum absorption at 620 nm, in agreement with literature [24]) clearly indicated a rather quantitative 1 F/mol conversion of permanganate according to \( \text{MnO}_4^- + e^- \rightarrow \text{MnO}_4^{2-} \). Upon continuing the electrochemical reduction to a final 2 F/mol relative charge vs. initial permanganate, the green intermediate solution became light blue, slowly turning to blue-grey. The UV-vis spectrum at 2 F/mol, recorded 2 minutes after the end of the electrolysis (blue line in Figure 3B), did not exhibit the expected maximum at 670 nm for \( \text{MnO}_4^{3-} \) [22], but displayed most of the characteristics of permanganate absorption (compare with the purple curve in Figure 3B). Actually, and as already reported [19], this color change is consistent with the instability of the hypomanganate ion in the presence of oxygen (our experiments were performed under aerobic conditions) as well as its synproportionation according to \( \text{MnO}_4^- + \text{MnO}_4^{3-} \rightarrow 2 \text{MnO}_4^{2-} \).

![Figure 3](image)

**Figure 3**

Electrochemical reduction of permanganate (10 mmol.L\(^{-1}\) contained in a 75 µL aqueous 1 mol.L\(^{-1}\) NaOH droplet) at 1 mA using the 2 electrode configuration. **A:** Photographs of the 75 µL droplet diluted in 1 mL deionized water before electrolysis (0), after 75 mC (1 F/mol) and 150 mC (2F / mol). **B:** UV-vis absorption spectra of the same diluted droplets recorded 2 min after the electrolyses.

4. Conclusion

One of the simplest and smallest setup allowed the achievement of fast and complete electrochemical conversions, from examples taken among organic substrates or inorganic species. Electrochemical preparations in droplets from the same setup should also be accessible in organic solvents, upon confinement within a chamber allowing saturated solvent vapor pressure, as described for analytical applications [5]. The possibility to carry out either controlled-potential electrolyses (3 electrodes) or galvanostatic ones (2 electrodes) make such a fast, low-amount / low cost electrochemistry available and attractive for novice or occasional operators as well as for teachers looking for simple electrochemical experiments.
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6. References


