Influence of chiral compounds on the oxygen evolution reaction (OER) in the water splitting process

Mirko Gazzotti¹, Andrea Stefani¹, Marco Bonechi², Walter Giurlani², Massimo Innocenti²*, Claudio Fontanesi¹*

¹Department of Engineering ‘Enzo Ferrari’, University of Modena and Reggio Emilia, Via Vivarelli 10, 41125 Modena, Italy
²Department of Chemistry, University of Firenze, via della Lastruccia 3, 50019 Sesto Fiorentino, Firenze, Italy.

Abstract
Results are presented concerning the influence on the water splitting process of enantiopure tartaric acid present in bulk solution. Stainless steel and electrodeposited nickel are used as working electrode (WE) surface. The latter is obtained by electrodeposition on the two poles of a magnet. The influence and role played by the chiral compound in solution has been assessed by comparing the current values, in cyclic voltammetry (CV) experiments, recorded in the potential range at which oxygen evolution reaction (OER) occurs. In the case of tartaric acid and nickel WE a spin polarization of about 4 % is found. The use of the chiral environment (bulk solution) and ferromagnetic chiral Ni electrode allows for observing the OER at a more favourable potential: about 50 mV (i.e. a cathodic, less positive, shift of the potential at which the oxygen evolution is observed).

Keywords: spin dependent electrochemistry, water splitting, nickel, chirality, OER

* corresponding author:
Massimo Innocenti, m.innocenti@unifi.it
Claudio Fontanesi, claudio.fontanesi@unimore.it
1. Introduction

Spin dependent electrochemistry (SDE) is a quite recently proposed method branch of electrochemistry [1]. The peculiar characteristic of SDE is that it allows for the control, production and measurement of spin polarization currents within an electrochemical system, giving the opportunity to gain further physical insight on the influence and role of spin in the charge transport. From the technological point of view, a possible application of SDE is to enhance the efficiency of the hydrogen production via electrochemical dissociation of water: this process is commonly addressed as water splitting (WS) [2–7]. Nowadays, the energy problem is a crucial challenge to be faced. Hydrogen could be a decisive way out of it, because hydrogen is considered as the ultimate fuel for several reasons: i) it has the highest specific enthalpy of combustion of any chemical fuel ii) no carbon or $CO_2$ waste iii) water is the combustion final product.

In the present situation, the 80% of the sources of energy production consists of the traditional ones (such as oil, coal and natural gas). The perspective related to pollution issues and future shortage of oil and coal leads to search for a different type of fuel. The latter should effectively be coupled with alternative energy sources: photovoltaic, hydroelectric, wind turbine, geothermal, tidal power. All these energy sources produce electricity as the final form of energy. On the whole, energy obtained via “renewable energy” sources in 2015 represented the 16.7% of the installed power in the U.S.A. for the 13.8% of the total world-wide electricity generation [8] (including the hydrogen production that reached $10^{10}$ kg in 2015 in U.S.A.). In addition, the worldwide percentage of energy produced by renewable energy is higher than in the U.S.A., reaching the 24% of the total [5], with the 58% purchased from hydroelectric source. Within this picture, hydrogen role is growing in importance as it can be used as energy vector and/or fuel mixed with the fossil ones [9–11]. Concerning hydrogen, one of the hardest issues to be solved is its efficient storage, which has been the subject of extensive research and multiple solutions are currently viable. From classical liquid storage and high-pressure containers to “Metal hybrids” an “Metal hydride” compounds [12,13]. Moreover, an appealing positive feature of the hydrogen production obtained via water splitting is its potential use “in-situ” linked to solar energy photovoltaic electric generation in low-cost organic cells [14–16]. Here chirality comes into play in view of the spin filter effect, Chiral Induced Spin Selectivity (CISS) [17]. The positive boost due to the spin-filter effect has already been demonstrated for electrodes coated with chiral molecules [2,3,18–20]. This because ideal spin filtering would produce $OH^*$ radicals only of alpha (or only
for beta) spin state, hindering the production of $H_2O_2$ (hydrogen peroxide). The latter is a highly reactive and corrosive compound, and it can be considered a side reaction’s undesirable product of the $OH^-$ oxidation (compare equation (2)). In this work the CISS effect is exploited using chiral compounds in bulk solution, rather than adsorbed on the electrode surface. In principle, allowing to achieve a more efficient production of hydrogen through the water splitting but using a simpler and cheaper system. The water splitting process is studied as a function of different chiral additives/catalysers on both a stainless steel and nickel electrodes, the latter deposited directly on a magnetic surface.

2. Experimental

L- (+)-tartaric acid (the natural available enantiomer), L- (−)-aspartic acid, D- (+)-glucose were purchased from Sigma Aldrich and used without further purifications. The nickel electrodeposition was carried out using a classical Watt’s Bath (WB), whose composition is: 150 g/L of nickel sulphate, nickel chloride, 37 g/L boric acid $H_3BO_3$, pH = 5. Cyclic voltammetry (CV) measurements were performed using both Autolab PGSTAT 128N and CHI660A potentiostats, employing a typical three-electrode electrochemical cell arrangement concerning the measurements involving the stainless steel working electrode. A different arrangement was adopted in the case of the Ni electrodeposited on the magnet: a Teflon cell, featuring a hole (0.8 cm diameter) in the bottom, was used in a vertical configuration where the Ni-on-the-magnet working electrode was tightened from below, a Teflon ring was used to ensure no solution leakage from the cell. Thus, Steel AISI 316L or the Ni-on-the-magnet surfaces were used as working electrodes (WE), while a Pt wire and a silver, silver chloride, KCl saturated solution (Ag/AgCl/KCl sat) electrodes were the counter (CE) and reference electrodes (RE), respectively. In order to check the quality of the nickel surface obtained via electrodeposition on the magnet, control experiments were carried out using glassy carbon, Pt and evaporated gold as working electrodes [21]. A KOH 0.1 M aqueous solution is used as base electrolyte in all reported electrochemical measurements.

3. Results

The overall water splitting reaction is, in principle, a simple redox reaction:

$$2H_2O \rightarrow 2H_2 + O_2 \quad \Delta E = 1.23 \text{ V vs RHE}$$ (1)

In fact, the reaction is much more complex, in particular concerning the water oxidation at the anode. Here, in fact, the oxygen evolution reaction (OER) occurs via a multi-step
process, which involves a rather complex reaction mechanism featuring the transfer of 4 electrons [22]. The mechanism depends on a number of chemical as well as physical parameters, and in particular on the solution pH. In the following equations, (2)–(5), the overall half-cell reactions are reported:

- Alkaline solution:
  \[4OH^- \rightarrow O_2 + 2H_2O + 4e^- \quad E_a^0 = 0.401 \text{ V vs NHE} \quad (2)\]
  \[4H_2O + 4e^- \rightarrow 2H_2 + 4OH^- \quad E_c^0 = -0.828 \text{ V vs NHE} \quad (3)\]

- Acid solution:
  \[2H_2O \rightarrow O_2 + 4H^+ + 4e^- \quad E_a^0 = 1.229 \text{ V vs NHE} \quad (4)\]
  \[4H^+ + 4e^- \rightarrow H_2 \quad E_c^0 = 0.00 \text{ V vs NHE} \quad (5)\]

In principle, the potentials associated to reactions (2) and (4) are the thermodynamic values relevant to the OER, but beyond the thermodynamic potential it is necessary to apply an overpotential to observe the gas evolution. The key to improve the process efficiency is to minimize the overpotential [2–4]. The following sequence of elementary reaction steps represents the step-by-step overall reaction that occurs at the anode in an acid solution (where \(M\) represent the anode metallic conductive substrate) [22–25].

\[
\begin{align*}
M + H_2O_{(bulk)} & \rightarrow (M)OH_{(ads)} + H^+_{(sol)} + e^- \quad \Delta G_1 \\
(M)OH_{(ads)} & \rightarrow (M)O_{(ads)} + H^+_{(sol)} + e^- \quad \Delta G_2 \\
(M)O_{(ads)} + H_2O_{(bulk)} & \rightarrow (M)OOH_{(ads)} + H^+_{(sol)} + e^- \quad \Delta G_3 \\
(M)OOH_{(ads)} & \rightarrow M + O_2(gas) + H^+_{(sol)} + e^- \quad \Delta G_4
\end{align*}
\]

\(\Delta G_i\) is the Gibb’s free energy of each elementary steps. As reported in literature [28,29], \(\Delta G_{OER}\) is the max of the four \(\Delta G_i\) values, namely \(\Delta G_3\). \(\Delta G_i\) can be calculated with the formula \(\Delta G^0 = -nFE\) if all elementary steps are known and so the \(\Delta E^0\) of the redox couple involved [26–28]. Those values are useful in compiling the Latimer diagram of the reaction, a compact diagram of each step involved in the redox chain, reported in Scheme 1, where numbers in the purple arrows represent the \(\Delta E^0\) of the reactions from (6) to (9). The \(\Delta E^0\) of the total reaction is the results of the equation \(\frac{\sum_i n_i \Delta E_i^0}{\sum_i n_i}\) where \(n_i\) represent the number of electrons exchanged during each step.
Scheme 1 – Water electrolysis: Latimer reaction mechanism.

The energy for the reaction can be provided by different sources, e.g. with the use of light, often coupled with catalytic substrate, like TiO$_2$ [29–31]. Remarkably, the final product at the anode, i.e. oxygen, is produced in its triplet state, which is the most stable molecular oxygen species. Here spin dependent electrochemistry can play a role. Thus, experiments of electrochemical water splitting were carried out using two different anodes: i) a stainless steel AISI 316 L electrode ii) Ni electrodeposited directly on the North (or South) pole surface of a permanent magnet (Ni-on-the-magnet electrode). Cyclic voltammetry results are presented focusing on the response of the anodic OER. In the case of the AISI 316L steel, results were collected using, L-(+)-tartaric acid, D-(−)-aspartic acid and D-(+)-glucose. In the case of Ni-on-the-magnet electrode, CV measurements were still performed with bulk concentrations of D-(+)-glucose, L-(+)-tartaric acid and D-(−)-tartaric acid, and particular focus was also payed to potential sweeps experiments in presence of L-(+)-tartaric acid. On the whole, the experimental electrochemical outcome allows to discuss the role played on the OER efficiency by suitable spin injection; in relation to the chirality of compounds present in solution.

3.1 Steel AISI 316L

3.1.1 L-(+)-tartaric acid

Figure 1 shows cyclic voltammetry curves, obtained as a function of different concentrations of bulk L-(+)-tartaric acid, on the stainless steel working electrode. In both the forward and backward curves, the current is nearly negligible in the 0.0 to 0.6 V range. In the case of the base electrolyte solution (blue curve, Figure 1), at potential values larger than 0.6 V the current starts to increase almost linearly until reaching a final 2.5 mA value at 0.8 V. Tartaric acid 0.5 mM solution (green curve, Figure 1) shows a lower current, 2.4 mA, at 0.8 V. Tartaric acid 5 mM solution (red curve, Figure 1) shows an initial higher current but at the potential where the OER occurs (around 0.8 V) the current is lower when compared with the previous ones. Tartaric acid 25 mM solution (black curve, Figure
1) shows the lowest current, 1.2 mA, at 0.8 V. The current appears to be correlated to the concentration of the chiral compound with an inverse proportionality. The inset of Figure 1 presents a more detailed CV, in the 0.5 to 0.75 V potential range. The inset shows a small but neat difference between the forward and backward scans. The current for the KOH base electrolyte solution is always higher than the one measured in presence of tartaric acid.

![Cyclic voltammery curves recorded on a steel AISI 316L WE. 50 mV/s potential scan rate, Pt sheet as CE and Ag/AgCl/KCl_sat as RE. KOH 0.1 M aqueous solution is the base electrolyte. L-(+)-tartaric acid concentration: (blue) 0 mM, (green) 0.5 mM, (red) 5 mM, (black) 25 mM. The inset shows the CV detail in the 0.5 and 0.75 V potential range.](image)

**Figure 1** — Cyclic voltammery curves recorded on a steel AISI 316L WE. 50 mV/s potential scan rate, Pt sheet as CE and Ag/AgCl/KCl_sat as RE. KOH 0.1 M aqueous solution is the base electrolyte. L-(+)-tartaric acid concentration: (blue) 0 mM, (green) 0.5 mM, (red) 5 mM, (black) 25 mM. The inset shows the CV detail in the 0.5 and 0.75 V potential range.

### 3.1.2 L-(−)-aspartic acid

Figure 2 shows CVs in solutions with different concentrations of L-(−)-aspartic acid. Qualitatively, CVs pattern is quite similar to that shown in Figure 1. Quantitatively, all the tartaric acid solutions at different concentration feature a lower current with respect to that of the base electrolyte. Note that, no peaks are evident in the backward scan, indicating that any reaction (OER) occurring during the forward scan (oxidation regime) is irreversible.
Figure 2 – Cyclic voltammetry curves recorded on a steel AISI 316L WE. 50 mV/s potential scan rate, Pt sheet as CE and Ag/AgCl/KCl_{sat} as RE. KOH 0.1 M aqueous solution is the base electrolyte. L-(–)-aspartic acid concentration: (red) 0 mM, (blue) 1 mM, (light blue) 10 mM, (green) 50 mM, (black) 100 mM.

3.1.3 D-(+)-glucose

Figure 3 shows CVs as function of the concentration of D-(+)-glucose. A systematic variation in the current as a function of the D-(+)-glucose concentration can be noted. The maximum efficiency, obtained comparing the current response at fixed potential values, is found for the 10 mM D-(+)-glucose concentration (green curve, Figure 3). The KOH base electrolyte solution is characterized by the lowest current values (red curve, Figure 3).

Figure 3 – Cyclic voltammetry curves recorded on a steel AISI 316L WE. 50 mV/s potential scan rate, Pt sheet as CE and Ag/AgCl/KCl_{sat} as RE. KOH 0.1 M aqueous solution is the base electrolyte. D-(+)-glucose concentration: (red) 0 mM, (light blue) 1 mM, (green) 10 mM, (black) 100 mM.
3.2 Nickel electrodeposited on magnet

3.2.1 D-(+)-glucose

Figure 4 shows the cyclic voltammetry of Ni electrodeposited on the north surface of a permanent magnet used as working electrode in two different solutions: D-(+)-glucose 0.1 M in base electrolyte (KOH 0.1 M) and bare base electrolyte. The experimental set-up is the one with the Teflon cell described in Section 2 (2. Experimental). Note that Error! Reference source not found.4 and Error! Reference source not found.5 values are normalized to point out the differences between the current in the solution with the chiral compound and the one with only the support electrolyte KOH. The value that has been used to normalize the data was the current associated to the oxidation of the Ni surface, otherwise the current peak, in the forward scan, in the base electrolyte curve: an opportune dividing coefficient is adopted in order to make that peak current value equal to 1. In the case of pure electrolyte KOH solution (red curve, Figure 4) the Ni peaks are evident, with the oxidation one happening at 0.55 V in the forward scan while during the backward scan the reduction peak splits in two smaller peaks, one at 0.39 V and the other at 0.46 V. Otherwise, the curve describing the current in the solution containing the chiral compound is completely different, resulting more like the one found with the stainless steel electrode (Figure 3). In particular, the D-(+)-Glucose curve doesn’t have the peaks of the Ni nor in forward nor in backward scan. The absence of peaks in the glucose’s curve means that the redox process became irreversible if the glucose is added to the KOH solution. The anodic peak current in the solution containing the chiral compound results being 50% higher than the one measured with only the support electrolyte.

![Normalized cyclic voltammetry curves recorded with electrodeposited Ni on the North pole of the magnet. Red curve: KOH 0.1 M aqueous solution. Black curve: 0.1](image)

**Figure 4** – Normalized cyclic voltammetry curves recorded with electrodeposited Ni on the North pole of the magnet. **Red curve:** KOH 0.1 M aqueous solution. **Black curve:** 0.1
M of D-(-)-Glucose in KOH 0.1 M aqueous solution. 50 mV/s potential scan rate, a Pt as CE, Ag/AgCl/KCl sat as RE.

Error! Reference source not found.5 shows the cyclic voltammetry measured on Ni electrodeposited on the south pole of the permanent magnet used as working electrode; those experiments were carried out with the same solutions of Error! Reference source not found.4, in the same operative conditions. As in Error! Reference source not found., the D-(-)-Glucose’s curve doesn’t have the characteristic peaks associated to the oxidation and the reduction of the Ni.

For what concerns the support electrolyte, the graph shows one peak at 0.55 V in oxidation and one peak and a shoulder in reduction at 0.37 V and 0.45 V respectively, coherent to what has been already seen in Error! Reference source not found.4. As for Error! Reference source not found.4, the ordinate axis has been normalized using the value of the current associated to the oxidation peak of the support electrolyte, so that peak would result having value equals to 1 on the normalized axis. In comparison with Figure 4, the current of the solution containing the chiral compound is lower in the south pole, resulting having the maximum current in oxidation almost equal to the one measured with the support electrolyte.

Figure 5 – Normalized cyclic voltammetry curves recorded with electrodeposited Ni on the south pole of the magnet. Red curve: KOH 0.1 M aqueous solution. Black curve: 0.1 M of D-(-)-Glucose in KOH 0.1 M aqueous solution. 50 mV/s potential scan rate, Pt as CE, Ag/AgCl/KCl as RE.

Error! Reference source not found. shows the comparison between CVs collected with a Ni electrode on top of north and south magnet polarities, recorded in a wider potential range. Remarkably, at 1.2 V it is present a peak for both the curves and the north pole peak current of the oxidation process is larger than the south pole corresponding value.
Figure 6 – Comparison between cyclic voltammetry curves recorded with electrodeposited Ni on both poles of the magnet. **Red curve:** south pole with 0.1 M of D- (+)-Glucose in KOH 0.1 M aqueous solution. **Black curve:** north pole with 0.1 M of D- (+)-Glucose in KOH 0.1 M aqueous solution. 50 mV/s potential scan rate, Pt as CE, Ag/AgCl/KCl as RE.

### 3.3 Stepped sweeps

For a more accurate visual detection of the actual OER starting potential, “stepped sweeps” technique measurements were performed. In fact, differently from the previous CVs, these experiments feature incremental values of potential which is maintained constant for a chosen amount of time. These measurements are reported in Figures 7 and 8, showing the anodic current recorded as a function of time during the potential ramp for two different electrodes.

#### 3.3.1 Stepped sweeps on AISI 316 L

Stepped sweeps were performed using the AISI 316 L steel sheet as working electrode, with the L- (+)-tartaric acid present in bulk solution. The applied potential programme as a function of time features a first potential ramp from 0 to 0.75 V at a 50 mVs⁻¹ scan rate, followed by eight +0.01 V potential steps, up to 0.83 V. The potential was maintained constant for 10 s before the next step. Tartaric acid concentration 0.5 and 5 mM in bulk solution was used. A rather large increase in the current was found, which should reflect a more efficient way to produce oxygen. Indeed, visual observation of the anode shows that the oxygen evolution occurs at more negative (smaller) potentials, anticipating the OER between 20 to 30 mV with respect to the pure KOH base electrolyte solution (the relevant table (Table 1SI) is present in the Supporting Information).
Figure 7 – Stepped sweeps curves recorded on a steel AISI 316L WE, Pt sheet as CE and Ag/AgCl/KCl\textsubscript{sat} as RE. KOH 0.1 M aqueous solution is the base electrolyte. A potential ramp from 0 to 0.75 V at a 50 mV/s was applied, followed by eight +0.01 V potential steps, up to 0.83 V. The potential was maintained constant for 10 s between each step. L-(+)-tartaric acid concentration: (blue dash) 0 mM, (green) 0.5 mM, (red) 5 mM

3.3.2 Stepped sweeps Ni-on-magnet

Stepped potential sweeps experiments were recorded also exploiting a ferromagnetic electrode in tight contact with a magnet: the Ni electrodeposited on magnet (Ni-on-magnet) electrode. This to possibly maximize the spin-injection efficiency. Figure 8 shows the anodic current recorded as a function of time: measurements have been carried out with L-(+)-tartaric acid present in bulk solution, as well as with the pure 0.1 M KOH base electrolyte for reference purposes. The applied potential programme as a function of time features a first potential ramp from 0 to 0.70 V at a 50 mVs\textsuperscript{-1} scan rate, followed by +0.01 V potential steps, maintained constant for 15 s. The tartaric acid concentration is 0.5 mM in bulk solution. Results show two different situations depending on the orientation of the substrate magnet. In fact, when the Nickel sheet lays on the south pole of the magnet, the oxidation current results slightly higher for the TA solution than the KOH alone; moreover, a huge current increase can be observed between the chiral solution and the basic one when the North pole of the magnet is used. A significant increase in the current is found, which reflects a more efficient way to produce oxygen. Indeed, for both the North and the South magnet pole visual observation of the anode
shows that the oxygen evolution occurs at more negative (smaller) potentials, anticipating the OER between 30 to 50 mV with respect to the pure KOH base electrolyte solution (Table 1SI in the Supporting Information).

![Graph](image)

**Figure 8** – Stepped sweeps curves recorded on a Ni deposited on magnet working electrode, a Pt sheet is the CE, Ag/AgCl/KCl is the RE. KOH 0.1 M aqueous solution the base electrolyte. First a potential ramp from 0 to 0.70 V at a 50 mV/s was applied, followed by eight +0.01 V potential steps, up to 0.78 V. The potential was maintained constant for 15 s between each step. L-(+) -tartaric acid concentration: (blue dash) South mag orientation 0 mM, (red dash) South mag orientation 0 mM, (light blue) South mag orientation 0.5 mM, (orange) North mag orientation 0.5mM

4. Discussion

It is known that the presence in bulk solution of chiral compounds is able to “induce” an enantio-selective or generate local spatial chirality effects: see for instance results obtained in chiral ionic liquids and enantio-recognition effects in achiral ionic liquids, this latter induced by the presence of suitable bulk chiral compounds (even in a low concentration) [32]. Thus, in this work we exploit this idea to induce chirality effects by simply performing electrochemical experiments in the presence of bulk chiral compounds. As a whole, our results concerning the water electrolysis are consistent with such a picture already present in the literature. Concerning the CVs recorded by using a stainless steel anode, the L-(+)-tartaric acid and L-(−)-aspartic acid exert a blocking activity on the electrode. The anodic current is found to decrease as the concentration of the acid is increased (compare CVs shown in Figure 1 and Figure 2). Probably this effect
is due to simple coulombic attraction between the anode (charged positively) and the relevant acid anion, which can be adsorbed on the surface eventually leading to a decrease in the overall current. A different electrochemical behaviour is found when the D-(+)-glucose is present in the solution, Figure 3. The current is always found larger than that of the base electrolyte, and in terms of efficiency the current increases as a function of the concentration until 10 mM, then starts to decrease for larger concentrations (compare black line, 100 mM, Figure 3). A different picture emerges when the ferromagnetic Ni surface, deposited on a magnet acting as substrate, is used as the anode. In the case of the L-(+)-tartaric acid, Error! Reference source not found.8, the current recorded in response to potential sweeps in the presence of the bulk chiral compound is always larger than that of the base electrolyte. In this peculiar case, the effect of the presence of the chiral compound in bulk solution has been investigated by comparing the peak current values as a function of the magnet orientation (North vs. South), showing significant differences in the values of the current obtained at a fixed potential. Such an analysis is a crucial issue in unravelling the role of the spin. Moreover, to obtain a complete picture, the electrochemical results can be compared for the two enantiomers. Altogether four different situations are to be quantitatively compared: 1) L-(+)-tartaric acid North 2) L-(+)-tartaric acid south 3) D-(−)-tartaric acid north 4) D-(−)-tartaric acid south. Table 2SI, 3SI, 4SI and Table 5SI report, in the most synthetic way, the current results obtained performing CVs in all four cases. Moreover, Table 6SI and Table 7SI aim to present in a clear and simple way, as much as possible, the catalytic effect observed in the different combination of enantiomers and the magnet pole. In particular, the sign found in column titled “sgn (J_{\text{ratio North}} - J_{\text{ratio South}})” (Table 6SI and Table 7SI) exhibit what is the most efficient combination. Remarkably, the L-(+)-tartaric acid North is more effective than L-(+)-tartaric acid South in a consistent way, while the D-(−)-tartaric acid south combination is found more effective than the D-(−)-tartaric acid North (coherently with the results of the stepped sweeps previously presented in Section 3.3.2). In the case of the glucose, only the D-(+)-glucose enantiomer was examined and it is found that the current for the D-(+)-glucose north combination is larger than the D-(+)-glucose south one, Error! Reference source not found.4 and Figure 5. Please note that due to fluctuations in the base electrolyte CVs, in Error! Reference source not found.4 and Error! Reference source not found.5 normalized data are presented. Table 8SI and Table 9SI report the percentage of spin polarization of currents in the experiments made on the
different Ni surface deposited on poles’ magnet. Those values are calculated with the formula (10).

\[
\frac{\frac{I(\text{Tartr})}{I(\text{KOH})}_{\text{North}} - \frac{I(\text{Tartr})}{I(\text{KOH})}_{\text{South}}}{\frac{I(\text{Tartr})}{I(\text{KOH})}_{\text{North}} + \frac{I(\text{Tartr})}{I(\text{KOH})}_{\text{South}}} = SP\%
\]

5. Conclusion

In this work it was explored the influence of chiral compounds in bulk solution on the hydrogen production process pursued via water electrolysis, often addressed as “water splitting”. To this end two different electrode materials (steel and Ni) and three organic chiral compounds (tartaric acid, aspartic acid, glucose) have been selected. Assessment of the catalytic activity of the different combinations of organic compound and electrode material was performed by measuring CVs under controlled conditions. The KOH 0.1 M aqueous solution base electrolyte was selected as the experimental reference situation to evaluate the catalytic effect on the water splitting process. Chiral compounds have been selected because they are known to enhance the OER efficiency due to spin effects [2–7]. In this work the chiral compounds are present in bulk solution rather than adsorbed on the electrode surface. The results are encouraging both on steel and on Ni as well. The L-(+)-tartaric acid yields a moderate potential shifting effect (50 mV in the most favourable case), just on the shoulder preceding the current ramp for the OER. On the contrary, the aspartic acid does not seem to exert any prominent effect. In the case of the Ni working electrode the effect of the magnetic field, to select “up” and “down” spin injection, was investigated. Here the most interesting results are obtained. The comparison of the oxidation current peak, essentially due to the oxygen evolution reaction, yields a final consistent picture. Where the L-(+)-tartaric acid is found to enhance the evolution when the north-pole of the magnet is placed in direct contact with the Ni anode surface, the situation is reversed, i.e. larger efficiency for the D-(−)-tartaric acid coupled with the south magnetic field orientation. By and large, it appears that the results obtained using bulk chiral compounds yield a consistent spin filtering effect. On the whole, an average 4 % spin polarization value is obtained (average obtained by the SP values of each measurement), a value which is definitively less than the 15 % to 20 % SP range obtained in the case of well-ordered adsorbed monolayers directly on top of the electrode surface. In this case, using bulk compounds the preparation of the systems is by far easier and less
demanding. Without using exotic materials (for example cadmium based nanoparticles) which are also cheap and non-strategic.

All in all, the results obtained especially on the Ni as a function of the magnet orientation, relate well with results previously presented in the literature. This gives further impulse to the scientific research in the field of spin effects in the OER process.

Acknowledgements
The authors acknowledge financial support for this research by University of Modena and Reggio Emilia (Department of Engineering ‘Enzo Ferrari’), through “Photo-electrochemical water splitting” (FAR 2019) project.

References
[1] D. Mishra, T.Z. Markus, R. Naaman, M. Kettner, B. Göhler, H. Zacharias, N. Friedman, M. Sheves, C. Fontanesi, Spin-dependent electron transmission through bacteriorhodopsin embedded in purple membrane, Proc. Natl. Acad. Sci. 110 (2013) 14872–14876. https://doi.org/10.1073/pnas.1311493110.
[2] W. Mtangi, V. Kiran, C. Fontanesi, R. Naaman, Role of the Electron Spin Polarization in Water Splitting, J. Phys. Chem. Lett. 6 (2015) 4916–4922. https://doi.org/10.1021/acs.jpclett.5b02419.
[3] W. Mtangi, F. Tassinari, K. Vankayala, A. Vargas Jentzsch, B. Adelizzi, A.R.A. Palmans, C. Fontanesi, E.W. Meijer, R. Naaman, Control of Electrons’ Spin Eliminates Hydrogen Peroxide Formation During Water Splitting, J. Am. Chem. Soc. 139 (2017) 2794–2798. https://doi.org/10.1021/jacs.6b12971.
[4] C. Fontanesi, R. Naaman, W. Mtangi, Water splitting method and system, WO 2016056011 A1, 2016.
[5] X. Liang, J. Liu, D. Zeng, C. Li, S. Chen, H. Li, Hydrogen generation promoted by photocatalytic oxidation of ascorbate and glucose at a cadmium sulfide electrode, Electrochimica Acta. 198 (2016) 40–48. https://doi.org/10.1016/j.electacta.2016.03.023.
[6] F.A. Garcés-Pineda, M. Blasco-Ahicart, D. Nieto-Castro, N. López, J.R. Galán-Mascarós, Direct magnetic enhancement of electrocatalytic water oxidation in alkaline media, Nat. Energy. 4 (2019) 519–525. https://doi.org/10.1038/s41560-019-0404-4.
[7] S. Haschke, D. Pankin, Y. Petrov, S. Bochmann, A. Manshina, J. Bachmann, Design Rules for Oxygen Evolution Catalysis at Porous Iron Oxide Electrodes: A 1000-Fold Current Density Increase, ChemSusChem. 10 (2017) 3644–3651. https://doi.org/10.1002/cssc.201701068.
[8] P. Beiter, T. Tian, 2015 Renewable Energy Data Book, National Renewable Energy Lab. (NREL), Golden, CO (United States), 2016. https://doi.org/10.2172/1339347.
[9] K. AlRafea, M. Fowler, A. Elkamel, A. Hajimiragha, Integration of renewable energy sources into combined cycle power plants through electrolysis generated hydrogen in a new designed energy hub, Int. J. Hydrog. Energy. 41 (2016) 16718–16728. https://doi.org/10.1016/j.ijhydene.2016.06.256.
[10] G. Krajačić, R. Martins, A. Busuttil, N. Duić, M. da Graça Carvalho, Hydrogen as an energy vector in the islands’ energy supply, Int. J. Hydrog. Energy. 33 (2008) 1091–1103. https://doi.org/10.1016/j.ijhydene.2007.12.025.

[11] L. Ntziachristos, C. Kouridis, Z. Samaras, K. Pattas, A wind-power fuel-cell hybrid system study on the non-interconnected Aegean islands grid, Renew. Energy. 30 (2005) 1471–1487. https://doi.org/10.1016/j.renene.2004.11.007.

[12] A. Züttel, Materials for hydrogen storage, Mater. Today. 6 (2003) 24–33. https://doi.org/10.1016/S1369-7021(03)00922-2.

[13] S. Niaż, T. Manzoor, A.H. Pandith, Hydrogen storage: Materials, methods and perspectives, Renew. Sustain. Energy Rev. 50 (2015) 457–469. https://doi.org/10.1016/j.rser.2015.05.011.

[14] P. Morvillo, F. Parenti, R. Diana, C. Fontanesi, A. Mucci, F. Tassinari, L. Schenetti, A novel copolymer from benzodithiophene and alkylsulfanyl-bithiophene: Synthesis, characterization and application in polymer solar cells, Sol. Energy Mater. Sol. Cells. 104 (2012) 45–52. https://doi.org/10.1016/j.solmat.2012.04.044.

[15] P. Morvillo, R. Diana, C. Fontanesi, R. Ricciardi, M. Lanzi, A. Mucci, F. Tassinari, L. Schenetti, C. Minarini, F. Parenti, Low band gap polymers for application in solar cells: synthesis and characterization of thienothiophene-thiophene copolymers, Polym. Chem. (2013). https://doi.org/10.1039/C3PY01618H.

[16] S.J. Appleyard, Simple photovoltaic cells for exploring solar energy concepts, Phys. Educ. 41 (2006) 409.

[17] K. Ray, S.P. Ananthavel, D.H. Waldeck, R. Naaman, Asymmetric Scattering of Polarized Electrons by Organized Organic Films of Chiral Molecules, Science. 283 (1999) 814–816. https://doi.org/10.1126/science.283.5403.814.

[18] I. Carmeli, K.S. Kumar, O. Heifler, C. Carmeli, R. Naaman, Spin Selectivity in Electron Transfer in Photosystem I, Angew. Chem. Int. Ed. 53 (2014) 8953–8958. https://doi.org/10.1002/anie.201404382.

[19] R. Naaman, D.H. Waldeck, Chiral-Induced Spin Selectivity Effect, J. Phys. Chem. Lett. 3 (2012) 2178–2187. https://doi.org/10.1021/jz300793y.

[20] F. Tassinari, S.P. Mathew, C. Fontanesi, L. Schenetti, R. Naaman, Electric-Field-Driven Alignment of Chiral Conductive Polymer Thin Films, Langmuir. 30 (2014) 4838–4843. https://doi.org/10.1021/la500657e.

[21] M. Gazzotti, S. Arnaboldi, S. Grecchi, R. Giovanardi, M. Cannio, L. Pasquali, A. Giacomino, O. Abollino, C. Fontanesi, Spin-dependent electrochemistry: Enantioselectivity driven by chiral-induced spin selectivity effect, Electrochimica Acta. 286 (2018) 271–278. https://doi.org/10.1016/j.electacta.2018.08.023.

[22] X. Sala, S. Maji, R. Bofill, J. García-Antón, L. Escriche, A. Llobet, Molecular Water Oxidation Mechanisms Followed by Transition Metals: State of the Art, Acc. Chem. Res. 47 (2014) 504–516. https://doi.org/10.1021/ar400169p.

[23] G.V. Buxton, C.L. Greenstock, W.P. Helman, A.B. Ross, Critical Review of rate constants for reactions of hydrated electrons, hydrogen atoms and hydroxyl radicals (·OH/O− in Aqueous Solution, J. Phys. Chem. Ref. Data. 17 (1988) 513–886. https://doi.org/10.1063/1.555805.

[24] H. Dau, C. Limberg, T. Reier, M. Risch, S. Roggan, P. Strasser, The Mechanism of Water Oxidation: From Electrolysis via Homogeneous to Biological Catalysis, ChemCatChem. 2 (2010) 724–761. https://doi.org/10.1002/cctc.201000126.

[25] R. Giovanardi, C. Fontanesi, W. Dallabarba, Adsorption of organic compounds at the aluminium oxide/aqueous solution interface during the aluminium anodizing
process, Electrochimica Acta. 56 (2011) 3128–3138. https://doi.org/10.1016/j.electacta.2011.01.065.

[26] C. Fontanesi, R. Benassi, R. Giovanardi, M. Marcaccio, F. Paolucci, S. Roffia, Computational electrochemistry. Ab initio calculation of solvent effect in the multiple electroreduction of polypyridinic compounds, J. Mol. Struct. 612 (2002) 277–286.

[27] C. Fontanesi, Theoretical study of the dissociative process of the 4-chlorotoluene radical anion, J. Mol. Struct. THEOCHEM. 392 (1997) 87–94. https://doi.org/10.1016/S0166-1280(97)90379-1.

[28] M. Marcaccio, F. Paolucci, C. Fontanesi, G. Fioravanti, S. Zanarini, Electrochemistry and spectroelectrochemistry of polypyridine ligands: A theoretical approach, Inorganica Chim. Acta. 360 (2007) 1154–1162.

[29] A. Pozio, A. Masci, M. Pasquali, Nickel-TiO2 nanotube anode for photoelectrolysers, Sol. Energy. 136 (2016) 590–596. https://doi.org/10.1016/j.solener.2016.07.040.

[30] Y. Kuang, T. Yamada, K. Domen, Surface and Interface Engineering for Photoelectrochemical Water Oxidation, Joule. (n.d.). https://doi.org/10.1016/j.joule.2017.08.004.

[31] M. Rudolph, D. Stanescu, J. Alvarez, E. Foy, J.-P. Kleider, H. Magnan, T. Minea, N. Herlin-Boime, B. Bouchet-Fabre, M.-C. Hugon, The role of oxygen in magnetron-sputtered Ta3N5 thin films for the photoelectrolysis of water, Surf. Coat. Technol. 324 (2017) 620–625. https://doi.org/10.1016/j.surfcoat.2016.09.007.

[32] S. Rizzo, S. Arnaboldi, V. Mihali, R. Cirilli, A. Forni, A. Gennaro, A.A. Isse, M. Pierini, P.R. Mussini, F. Sannicolò, “Inherently Chiral” Ionic-Liquid Media: Effective Chiral Electroanalysis on Achiral Electrodes, Angew. Chem. Int. Ed. 56 (2017) 2079–2082. https://doi.org/10.1002/anie.201607344.