Enhancing hydrogen production from steam electrolysis in molten hydroxides via selection of non-precious metal electrodes

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Abstract

There are still gaps in the field of reference electrode that is needed to assist electrolysis in high temperature electrolytes (e.g. molten hydroxides) for H₂ gas production. This research aims to fill the gaps by preparing the Ni/Ni(OH)₂ reference electrode and, more importantly, testing its effectiveness against important performance factors, including ion conducting membrane (e.g. mullite tubes), internal electrolyte composition, working temperature and electrochemical control (e.g. potential scan rate). Then, this reference electrode was used to assist the study of the electrocatalytic activity of a range of cheaper working electrode materials, including stainless steel (St.st), Ni, Mo and Ag, in comparison with Pt, by means of chronoamperometry and voltammetry. The effect of introducing steam into the electrolyte (eutectic mixture of NaOH and KOH) on the electrocatalytic activity of each of these working electrodes was also studied. It was observed that
the potential of hydrogen evolution on different working electrodes followed an order of Pt > Ni > St.st > Ag > Mo (positive to negative). The performance of each working electrode was confirmed through chronoamperometry for hydrogen evolution at a constant potential of -0.7 V. It was also found in cyclic voltammetry and confirmed by chronoamperometry that the introduction of steam was apparent in increasing the current density at the cathodic limit for hydrogen evolution. It is hoped that this study will help develop non-precious metal electrodes for the production of the hydrogen fuel. In future, there will be a potential in the threshold concentration of steam for H₂ gas production.

**Keywords:** Renewable energy, Hydrogen production; Electrocatalytic activity; Water splitting; Reference electrode, Fuel cells and Chronoamperometry.

### 1 Introduction

An important pollution free fuel that can meet future needs and can also lessen the problems instigated by the consumption of conventional fuels is hydrogen (H₂). As a highly efficient fuel, H₂ can be used for power generation, transportation and heating, and has the potential to substitute, at least partly, existing fuels. It is well documented and highlighted more recently that the commonly used process for H₂ production from water, as a renewable and clean source [1], is electrolysis which splits water into its core ingredients; H₂ and oxygen (O₂) [2, 3]. Splitting of water, or more accurately steam in high temperature molten hydroxides, by means of electrolysis has great importance and advantages.
The main advantage is that in the electrolysis, heat is used as a source of energy and heat is cheaper than electricity in terms of sources and conversion (production). The conductivity of a hydroxide electrolyte at high temperatures is very good and increases with increasing temperature. The hydroxide electrolyte at high temperatures is specific to reduce the loss of energy due to the overpotential of an electrode through acceleration of the reaction kinetics [4]. All these contribute to increasing the net energy efficiency of the process. Molten hydroxides could themselves play the role of a catalyst during the reaction and thus in this technology, there is no need for precious metals as a catalyst [5]. On increasing the temperature, the decomposition voltage of a compound is usually reduced, and this phenomenon is well observed in the case of water electrolysis. In the case of a thermally insulated electrolysis cell, energy consumption is constantly minimised [6]. This can be considered for long term electrolysis. Another advantage of electrolysis at high temperatures is that the current flowing continuously through the molten electrolyte contributes to additional internal heating that is needed to compensate any heat loss that is inevitable, even in a thermally well-insulated cell.

Suitable ion conducting membrane materials are required for the better fabrication of a reference electrode. Therefore, the selection of good ion conducting materials in this field is very important especially in the case of high temperature electrolytes. The redox couple, Ag/AgCl, is commonly used in reference electrodes coupled with different ion conducting membrane materials, such as quartz, Pyrex, porcelain, and mullite [7, 8]. The Ag/AgCl couple contained in silica tube covered with graphite, or enclosed in alumina membrane [9] have also been stated as choices for high temperature molten salts. Selection of membrane materials is important for the fabrication of an
electrode in molten hydroxides. Thus, those ionic membranes with good chemical stability, reusability and reproducibility are important [10-12].

Several studies have previously investigated the use of different working electrodes such as nickel (Ni) [13, 14], platinum (Pt) [15], silver (Ag) [16], molybdenum (Mo) [17] or stainless steel (St.st) [18]. These working electrodes have the ability to conduct an adequate catalytic activity for splitting water in hydroxide electrolyte; resulting in the enhancement of reaction kinetics and a subsequent upturn in the production of H$_2$ gas. These metals were either investigated in a molten hydroxide or in an aqueous solution of hydroxide at low temperatures. These studies under different operating conditions used hydroxide and a different type of reference electrode to control the working electrode. For instance, Miles et al. [13] studied the electrochemistry of molten NaOH-KOH salt at 280 °C using platinum, nickel and silver as working electrodes against the reference electrode of Ag$^+$/Ag.

The study of Ge et al. [15] involved cyclic voltammetry on a Pt or Ni wire, or a NiO pellet as the working electrode in fused NaOH at 550 °C. A Ni rod was selected as the pseudo reference electrode for the analysis of NiO reduction mechanism into the melt. Zabinski et al. [19] employed a Co-Mo-C alloy to augment the cathodic potential for electrolytic evolution of H$_2$ in a solution of 8 M NaOH at 90 °C. This was also carried out to inhibit the dissolution of Mo during open circuit dipping in the solution. Another investigation [20] coated the St.st electrode with a Ni-Mo-Fe film to enhance the catalytic activity for H$_2$ evolution in the dilute basic solution. When Ni, Co and NiCo were used as coatings to support a carbon felt electrode, this also resulted in enhanced catalytic activity for the H$_2$ evolution reaction (HER) [21].
This study is carried out to fabricate and test the Ni/Ni(OH)$_2$ reference electrode with an ion conducting mullite membrane. The reasons behind choosing Ni for the reference electrode fabrication were its range of chemical, physical, electrocatalytic, structural and corrosion resistant properties [22, 23]. Also, the electrocatalytic activity of a range of cheap working electrodes was comprehensively studied against this novel reference electrode. Then, the potentials of these cheap working electrodes for hydrogen gas production via splitting water were assessed in the presence of the eutectic mixture of NaOH-KOH (49:51, mol%) at 300 °C. The effect of steam at the electrocatalytic activity of these working electrodes has also been studied. Chronoamperometry and cyclic voltammetry were used to investigate the electrocatalytic activity of the working electrodes in this study.

2 Materials and methods

2.1 The Ni/Ni(OH)$_2$ reference electrode

The Ni/Ni(OH)$_2$ reference electrode was fabricated with a mullite tube (Multi-Lab Ltd) as the ionic membrane. The mullite tube consisted of Al$_2$O$_3$ and SiO$_2$ (36:62, mol%) with the diameter, length and thickness being 5 mm, 500 mm and 1 mm, respectively. The tube had 0.02 vol% of water absorption aptitude with 2.7 g cm$^{-3}$ of bulk density. The internal electrolyte was prepared by mixing 1.0 mol% Ni(OH)$_2$ (Arcos Organics) with the eutectic mixture of NaOH and KOH (49:51, mol%). Then this mixture was implanted into the conducting ionic mullite tube. This synthesised mixture was used internally as an electrolyte. The solubility of the Ni(OH)$_2$ in the internal electrolyte of a reference electrode is of great importance. It was described by researchers in past that the
dissolution of the Ni(OH)$_2$ in basic solution was not significant at room temperature, this trend was observed to be opposite in case of acidic solution [24].

On the other hand, Ni(OH)$_2$ has a solubility product of $6.5 \times 10^{-18}$, and this value was unaffected when noticed from the reaction of Ni(OH)$_2$ with either acid or base. Therefore, a minute amount of 1.0 mol% of Ni(OH)$_2$ was applied in the internal electrolyte. This composition was previously reported for a high H$_2$ evolution rate [25]. The hydroxide mixture (1.16 g) was placed inside the mullite tube which was positioned inside the retort, but it was quickly filled with the mixture of the salts of hydroxides when the tube outside the crucible, to avoid any absorption of moisture contents from the open air. The internal and external composition of the eutectic hydroxides should be the same. The temperature was then raised up to 300 °C to thoroughly melt the mixtures of hydroxide salts in the mullite tube membrane. The tube was filled up to the length of 12 cm. After that a Ni wire with 0.5 mm diameter and 99.98% pure temper annealed was introduced inside the tube.

The Ni wire was enclosed inside the mullite tube the left this for overnight to accomplish the melting of the salts mixture at 300 °C. Next to this, the furnace was cooled to the required temperature to solidify the molten melts mixture in the tube and sealed it. Alumina crucible with 280 mL volume and 120 mm height (Almath Crucibles Ltd) was used for the performance evaluation of the Ni reference electrode. Argon atmosphere was applied for these test by using an electrochemical analyser of Iviumn Stat multi-channel. For all these experiments 250 g of the molten hydroxides was left in the presence of 40 cm$^3$min$^{-1}$ of argon gas for 24 h and 300 °C temperature before use. The experimental setup for the designed electrodes is presented in Fig. 1
2.2 Specifications of working electrodes

The counter electrode used in this study was prepared of stainless steel rod of 304 Grade, along with the diameter of 5 mm (Unicorn Metals). Five different types of working electrodes were used including Ni, Pt, Ag, Mo, and St.st. The dimensions and properties of these working electrodes are as follows. The 99.98% pure Temper Annealed Ni working electrode was used with a 0.5 mm diameter. The Pt working electrode was 99.95% pure Temper Annealed this was also about 0.5 mm diameter. The third used working electrode was Ag with 99.99% pure Temper Annealed and 1.0 mm in diameter. The Mo working electrode was of 1.0 mm diameter and 99.95% pure Temper Annealed. The last working electrode was St.st with 0.25 mm diameter and 99.99% pure Temper Annealed. All these working electrodes were obtained from Advent Research Material.

The performance of the working electrodes was carried out in a cylindrical alumina crucible using the same protocol as conducted for the reference electrode. Though, it was not easy in practice to attain the exact requisite temperature because the electrolyte temperature is dependant on furnace temperature. So the temperature variation can be controlled by the furnace temperature. The furnace controller of temperature had an accuracy of ±1 °C. In addition to the furnace temperature electrolyte temperature was also affected by some other factors. Including the furnace insulation effectiveness and the ambient temperature.

2.3 Electrochemical investigation

The electrochemical methods used in this investigation were cyclic voltammetry and chronoamperometry. These techniques were used to study the behaviour of the working electrodes
in the molten salts at variable working conditions [26]. The measurements were made between one
of the working electrodes (e.g. Ni, Pt, Ag, Mo, St.st) and the designed reference electrode of Ni.
The depth of immersion for the working electrodes was ~14 mm inside the electrolyte. Cyclic
voltammetry (CV) measurements were noted from negative circuit potential to a positive one.
These type of analyses are very important that have already been used in different studies [27, 28].
CV investigations were also conducted at different temperatures and in the presence of steam
inside molten salts. Introduction of the steam at 7.28 cm$^3$ min$^{-1}$ flow rate was fixed, mixing with
argon gas that itself flows at 40 cm$^3$ min$^{-1}$.

The mixture of argon and steam was effervesced inside the molten salts. CV plots are plotted as
current density versus potential. Table 1 shows the working electrodes used in this study, their
diameters and calculated surface area respectively. The information regarding the different
operating temperatures included as supplementary material. The surface area of the working
electrodes can be calculated using Eq. (1).

A = \pi \times D \times h + \frac{1}{4} \times \pi \times D^2 \tag{1}

where A: the surface area of the immersion part inside the melt (cm$^2$), \pi: mathematical constant
(3.141), D: diameter of the working electrode (cm), and h: the immersion depth of the wire inside
the electrolyte (cm).

3 Results and discussion
In this section cyclic voltammetry scans were performed for different working electrodes (e.g. Ni, Pt, Ag, Mo and St.st) in eutectic molten hydroxide as explored below:

### 3.1 Cyclic voltammetry investigation of working electrodes

#### 3.1.1 Ni working electrode

To determine the functioning of different metal electrodes against the designed reference electrode cyclic voltammetry analyses were performed [29, 30]. For this, in the first run blank, Ni wire as a working electrode is used at a temperature of 300 °C and 100 mVs$^{-1}$ scan rate using the prepared nickel reference electrode in molten hydroxide. Fig. 2(a) displays the obtained cyclic voltammetry full scan. Fig. 2(b) shows cyclic voltammetry scanned from -0.8 V to -0.1 V vs reference electrode. The latter only focuses on the reduction limit.

The number of redox peaks can be easily noted as shown in Fig. 2(a). The C2 peak is the cathodic current credited to the oxide’s film reduction [31], made on Ni wire surface, while the A2 peak is anodic current attributed to its oxidation. In addition, the reduction potential started at -0.465V likely corresponds to the evolution of H$_2$ gas [15, 32] and the resultant chemical process is shown in Fig. 2(a) and expressed as reaction (2). The peak A1 is assigned to the generation of oxygen gas as seen in Fig. 2(a) and represented as reaction (3).

\[ 2 \text{H}_2\text{O} + 2e^- \rightarrow \text{H}_2(g) + 2 \text{OH}^- \]  

(2)

\[ 2 \text{OH}^- \rightarrow \frac{1}{2} \text{O}_2(g) + \text{H}_2\text{O} + 2e^- \]  

(3)
The peak C2 in Fig. 2(a) has a potential of -0.37 V which is scarcely noticeable compared to peak A2 in the first potential cycle (Fig. 2(a)). The redox reactions occurred at the Ni electrode surface in the eutectic molten hydroxides can be credited to this phenomenon [15]. The peak A2 denotes the oxidation of Ni wire [33], which may cause the accumulation of oxide layer on the nickel surface. Consequently, the peak C2 observed because of the reduction of this oxide layer. Furthermore, on limiting the applied voltage between -0.8 V and -0.1 V, the C2 peak vanishes as in Fig. 2(b) due to the lack of oxidation of the nickel wire.

The disappearance of the reduction peak C2 as shown in Fig. 2(a), reveals that no distinctive reduction peak in this potential scan was confirmed when the scan was limited as shown in Fig. 2(b). For the understanding of this reaction, the potential of H₂O and NiO decomposition at 300 °C temperature was measured by the help of HSC 6.0 software. The decomposition reactions are (4 and 5). It reveals from these findings that the two decomposition potentials are quite close, showing that those reactions may occur at the same time during the cathodic sweep. This finding agrees with [15] who stated that the decomposition potentials of water and nickel oxide at 550 °C were very close.

\[ \text{H}_2\text{O} \rightarrow \text{H}_2(g) + \frac{1}{2}\text{O}_2(g) \quad (E = -1.013 \text{ V}) \quad (4) \]

\[ 2\text{NiO} \rightarrow 2\text{Ni} + \text{O}_2(g) \quad (E = -0.965 \text{ V}) \quad (5) \]

Fig. 2(c) shows the CV plot of Ni working and reference electrode with and without the presence of steam at a temperature of 300 °C and 100 mV s⁻¹ scan rate. No substantial change in the anodic peak A2 (oxidation of Ni) is witnessed with the introduction of steam inside the molten melt as
presented in Fig. 2(c). At cathodic limit, an increase in the current density is noted that increases from -1.6 A cm\(^{-2}\) to -2.09 A cm\(^{-2}\) with the existence of steam. In order to exhibit the effect of the presence of steam in increasing the evolution of H\(_2\) gas, the potential scan is limited between -0.8 and -0.3 V as shown in Fig. 2(d). It is obvious from the figure that the reduction potential for the evolution of H\(_2\) gas is the same with and without steam. However, the effect of steam can be recognised by enhancing the current density limit for the evolution of hydrogen gas. This increase represents the amount of steam that exists with molten melt and contributed to an increase in the yield of the hydrogen gas.

3.1.2 Pt working electrode

In electrochemical studies Pt has great importance because of its good stable characteristics, therefore here Pt working electrode in molten hydroxide is used to achieve reliable CV scans against the designed Ni reference electrode [15]. Fig. 3(a) shows the CV plot of the Pt working electrode in the eutectic molten melt at the same conditions of temperature and scan rate like the previous Ni working electrode. It can be observed from Fig. 3(a) that the anodic limit A1 is observed because of the oxidation of the eutectic molten hydroxides [34] as in reaction (3). While the water reduction, made at the anode, derives the cathodic limit C1 [34] as in reaction (6).

\[
\text{H}_2\text{O} + \text{e}^- \rightarrow \frac{1}{2}\text{H}_2(\text{g}) + \text{OH}^-
\]  

Therefore the corresponding potential for hydrogen gas evolution at C2 is -0.44 V. To focus the scan on the cathodic limit the applied potential is limitised between -0.8 and -0.3 V as shown in Fig. 3(b). As because of this potential limitation, the scan is still stable but the reduction potential shifts negatively to a value of about 0.04 V smaller than the full scan. Furthermore, the potential
scan rate varies for the purpose of testation of the platinum working electrode in the settled conditions of temperature and molten hydroxide.

Furthermore, to the above factors, it is imperative as well to study the effect of steam in the eutectic fused salts with Pt wire working electrode. The reason for investigating this factor is to understand its effect on the behaviour of the platinum working electrode. The cyclic voltammogram scan can translate this change to the behaviour of the electrode. Fig. 3(c) shows the CV scan of platinum as a working electrode with and without the presence of steam. The cyclic voltammetry scan compares the influence of the presences of steam inside the eutectic molten hydroxide with the cyclic voltammetry scan without steam as shown in Fig. 3(c) at a temperature of 300 °C and a scan rate of 100 mV s⁻¹. The presence of steam inside the eutectic molten hydroxide directly affects the obtained cyclic voltammetry scan by increasing the flow of current density at the cathodic limit from 1.16 mA cm⁻² at C1 to 1.82 mA cm⁻² at C1'. This attributed to the increase steam bubbles around the cathode which upon electrolysis generate more hydrogen gas this is also well mentioned in literature [35].

For more clarification, the cyclic voltammetry scans are limited between -0.8 to -0.3 V with and without the presence of steam in the eutectic molten hydroxide, as shown in Fig. 3(d). There is no change in the cyclic voltammetry scan due to this limitation. The presence of steam merely influences the scan by increasing the current density limit of C1 to C1' which represents the reaction of the evolution of hydrogen gas. Even though the platinum electrode is a stable metal in a eutectic molten hydroxide at high temperatures; it still has limited use in the industry because it is classified as a precious metal [36].
3.1.3 Ag working electrode

The blank CV scan is also recorded for a silver working electrode in the studied molten melts, at a 300 °C temperature and 100 mV s\(^{-1}\) scan rate as shown in Fig. 4(a). Literature is scarce regarding the exact nature of the silver wire reaction in the hydroxide. For example, the research directed by Miles et al. [13] reported that the reaction of the silver wire in the molten hydroxides may involve \(\text{Ag}^+ \rightarrow \text{Ag}, \ \text{AgO}_2 \rightarrow \text{Ag}^+\text{O}_2\) or some other silver electrode reaction. In this experiment of Ag working electrode with Ni/Ni(OH)\(_2\) reference similar findings have been noted as of [13].

A couple of redox peaks observed in the CV scan of Ag electrolysis as displayed in Fig. 4(a). Therefore, the anodic peak A2 may be ascribed to the oxidation whereas the cathodic peak C2 is because of the reduction reaction. These peaks noted because of the oxide layer made on the surface of the silver wire. Moreover, the cathodically augmented current of C1 at -0.52 V corresponds to the evolution of \(\text{H}_2\) gas, these results are in connection with [32]. Equivalent to this, cathodic chemical reaction shown in equation (2). Reaction (3) represents the generation of oxygen gas as appears in peak A1 of Fig. 4(a). Formation of steam/ or water molecule noted from these mentioned equations.

Subsequently, the cyclic voltammetry scan is limited between -0.8 and -0.3V as shown in Fig. 4(b) to emphasise on the cathodic limit of the evolution of hydrogen gas. It can be observed from Fig. 4(b) that during this limited scanning, there is no considerable change in the starting point of the potential at the cathodic limit. Moreover, it can be noted that the current density at C1’ decreased. The current density at the cathodic peak C1 as shown in Fig. 4(a) is roughly about -0.61 A cm\(^{-2}\), decreasing to approximately -0.18 A cm\(^{-2}\) at C1’ Fig. 4(b) during the limited scan potential. The
disappearance of the oxidation peak A2 and the reduction peak C2 during the potential scan limitisation is because of the scan starts at potential more negative than the oxidation potential of silver wire. In this case, the decrease in the current density of C1' should be proportional to the silver oxide’s reduction to silver. Therefore, it should be mentioned that the silver oxide does not completely reduce at C2, but its reduction is completed at C1 simultaneously with the evolution of hydrogen reaction as seen in Fig. 4(a). This conclusion is based on a noticeable increase in the current density, which otherwise decreases when the potential scan is limited.

To understand the influence of steam in molten salts on the hydrogen gas evolution with Ag working electrode experiments performed at 300 °C and 100 mV s⁻¹ Fig. 4(d). These experiments were performed on the cathodic limit. The presence of steam with molten melt did not affect the reduction potential of the evolution of hydrogen gas reaction as shown in Fig. 4(d). Thus, the same scan was produced without steam as well. This stable behaviour of the scan can be attributed to how the silver working electrode works in the molten melts against the Ni working electrode. It can also describe in this way that Ag working electrode remains stable and did not show any active response against the steam. Ag stability is somehow opposite to electrocatalytic activity. Moreover, this stable behaviour is associated with the noble metal properties of the Ag [37, 38]. Nobel metals (Ag, Cu and Au) are least reactive against acids that is why use frequently for ornamental purposes due to lower reactivity. So this stable behaviour of the Ag metal in steam is directly linked with its noble or inert metal characteristics, further steam is amphoteric (acts as both acid and base) in nature.

3.1.4  **Mo working electrode**
Molybdenum is one of the transition metals. It has a good electrocatalytic capacity for enhancing the activity of other metals such as nickel [20]. Due to the electroactivity of the molybdenum, it was investigated as a working electrode in this study. Fig. 5(a) and b show the CV of the molybdenum working electrode vs Ni reference electrode at same conditions of temperature and scan rate. It can be observed from Fig. 5(a), that the electrochemical stability window is between A and A’ (approximately 0.33 V). The increase in the cathode current density corresponds to the evolution of H₂, similar current density increase is detected at C1 and -0.8 V [32]. Equivalent to this, a chemical equation is shown in reaction (2). While anodically an increase in the current at A1 corresponds to the oxygen gas evolution as can be observed in reaction (3).

The subsequent cyclic voltammetry scan is limited between -0.8 and -0.2V as shown in Fig. 5(b) to focus on the evolution of the hydrogen gas reaction. Therefore, no change in the reduction peak can be noticed if the scan range is limited to the potential located after the potential of the oxidation peak A’ reaction takes place. The effect of the presences of steam inside the molten salts on the molybdenum working electrode electrocatalytic behaviour for increasing the hydrogen and oxygen gas evolution is also considered. Fig. 5(c) and Fig. 5(d) show the CV scans of the molybdenum working electrode against the Ni/Ni(OH)₂ reference electrode with and without steam with eutectic melt.

It is obvious from Fig. 5(c) that the effect of the presence of steam is apparent in increasing the current density of the evolution of hydrogen gas at the cathodic limit C1’, and the simultaneous evolution of oxygen gas at the anodic limit A1’. At the cathodic limit, the current enhances from -0.12 A cm⁻² without the presence of steam at point C1 to -0.49 A cm⁻² with the presence of steam
at point C1'. At the anodic limit, the current rises from 0.14 A cm$^{-2}$ without the presence of steam at point A1 to 0.465 A cm$^{-2}$ with the presence of steam at point A', these results are in agreement with the literature [39].

In order to understand the effect of the steam’s presence on increasing the current density of the evolution of hydrogen gas reaction, the potential voltammetry scan is limited between -0.8 and -0.3 V respectively as shown in Fig. 5(d). It is very clear from constraining the scan range that there is a considerable effect of the presence of steam in increasing the current density of the evolution of hydrogen gas reaction. It increases from -0.164 A cm$^{-2}$ at C1 to -0.51 A cm$^{-2}$ at C1'. This result is shown in Fig. 5(d) through the increase of the molybdenum metal activity with steam and makes a significant change on the H$_2$ evolution.

### 3.1.5 St.st working electrode

Stainless steel (302) is composed of iron, nickel, chromium, manganese, silicon, carbon, phosphorus and sulphur. It was used in this study as a working electrode to examine its stability and productivity in the molten salts. Fig. 6(a) shows a cyclic voltammetry scan at an operating temperature of 300 °C and a scan rate of 100 mV s$^{-1}$. At the anodic limit A1, the corresponding peak is due to the oxidation of the melt (2 OH$^-$ → 0.5 O$_2$ (g) + H$_2$O + 2 e$^-$) while the reduction of the water formed at the anodic limit is seen at the cathodic limit C1. The corresponding reaction of the reduction of water becomes is shown in reaction (6).

In the case of the oxidation peak A2, it corresponds to the oxidation occurring on the surface of the stainless steel working electrode and the potential of oxidation observed at -0.33 V.
Subsequently, the CV scan is limited to a range between -0.8 V and -0.3 V in order to focus the scan on the cathodic limit for the HER as shown in Fig. 6(b). No change in the reduction potential which starts at -0.5 V, and the current at the cathodic limit C1 which approximately equals -1.4 A cm$^{-2}$; is discernible. The oxidation peak A2 disappears when the CV scan is limited, even though the potential of the return scan is positive prior to A2 peak.

Fig. 6(c) and d show the cyclic voltammetry of the stainless steel with and without the presence of steam with molten salts and same working conditions of temperature at 300 °C and scan rate of 100 mV s$^{-1}$. No significant change can be observed from the figure regarding the presence of steam with hydroxide salts at the cathodic limit C1 for the evolution of hydrogen reaction as shown in Fig. 6(c). At the oxidation peak, the current density increased from 0.12 A cm$^{-2}$ without the presence of steam (A2’) to 0.23 A cm$^{-2}$ with the presence of steam (A2). This increase in the current density from A2’ to A2 is responsible for increasing the surface area of the oxide metal.

The effect of changing the operating temperature of the eutectic molten hydroxide on the working electrode kinetics activity is shown in Fig. S1 was also studied. The studied temperatures were 225 and 300 °C respectively. It can be observed that the evolution of hydrogen gas becomes more efficient and sees an increase with increasing temperature for all working electrodes.

### 3.2 Working electrode’s performance evaluation

The stability of the reference electrode in different working conditions and its working companion against other reference electrode has already been verified in previous studies [30, 40]. The stability, reusability and reproducibility of the Ni/Ni(OH)$_2$ reference electrode have also been reported with good experimentation. This electrode has worked with stability and reproducibility
for almost 9 days. Furthermore, a comparison of the reference electrode with other conventional ones (quasi Pt and Ag) has also been made.

After studying the kinetic reaction of each working electrode separately in the eutectic molten hydroxide at different operating conditions respectively; it is imperative to compare their performance. This is essential for discerning which electrode provides more affordable, durable, stable kinetics; and also fast catalytic response for the HER. The comparison focuses mainly on the cathodic limit of the HER. Cyclic voltammetry scans of the different working electrodes (i.e. Ni, Pt, Ag, Mo, St.st) are compared in the eutectic molten hydroxide at a temperature of 300 °C, a potential scan rate of 100 mVs⁻¹ and an argon gas atmosphere as shown in Fig. 7.

It is obvious from Fig. 7 that each working electrode has a unique reduction potential value. It can, therefore, be observed from the above figure that the platinum working electrode had more positive reduction potential value (approximately -0.47 V) followed by the reduction potential of the nickel working electrode (approximately -0.49s V) and then the reduction potential of the stainless steel working electrode at -0.51 V. The reduction potential values of the silver and molybdenum working electrodes occurred at the lower end of the comparison at -0.53 V and -0.56 V respectively.

The results of this study are in close comparison with the literature [41] in which Ag/AgCl was used as a reference electrode. In this study, current density at the cathodic limit is the highest for nickel working electrode followed by the stainless steel and platinum working electrodes respectively. However, silver and molybdenum have the lowest current density respectively. Table
2, displays the reduction potential and the current density at the cathodic limit as observed from the above figure for different working electrodes.

A high current density measured at the cathodic limit means a high HER. This HER is influenced directly by the electrocatalytic activity of the working electrode inside the eutectic molten hydroxide. As mentioned and tabulated in Table 2. The highest hydrogen evolution reaction that can be achieved at the cathodic limit is done by using the nickel working electrode followed by stainless steel, platinum, silver and finally molybdenum. Therefore, the nickel wire had a higher electrode activity in comparison to the other working electrodes. This behaviour reinforces nickel as a popular choice in electrochemical processes as cathode material for the hydrogen gas evolution reaction. However, some studies such as [42] have revealed that nickel can be deactivated during H₂ generation in alkaline water electrolysis and the metal requires the V₂O₅ addition to the electrolyte to cause reactivation. These observed results were repeated for three times for all working electrodes, no change on the observed results was experienced.

### 3.3 Hydrogen evolution reaction (HER)

Fig. 8(a) shows the obtained current-time chronoamperometry at a constant potential of all tested working electrodes in the eutectic molten hydroxide during 10 minutes of the HER. This test was executed at an operating temperature of 300 °C and 40 cm³/min⁻¹ argon atmosphere. The chronoamperograms show that the electrodes change during the first stages of HER, accomplishing a near stationary state. Their reactivity is retained along the noted time period, with platinum followed by nickel being by far, the best one material among the tried (tested) materials and displaying the highest current density values in comparison to stainless steel, silver and
molybdenum. This result confirms that the blank metal of platinum and nickel working electrodes in the eutectic molten hydroxide respectively have a better performance for splitting steam to produce hydrogen gas. The stainless steel working electrode is third in order for hydrogen gas production.

The performance of the different working electrodes was also tested with the presence of steam inside the eutectic molten hydroxide and at an operating temperature of 300 °C, as shown in Fig. 8(b). It is obvious from the figure that the attained current density value of different working electrodes (without steam) slightly increased with the presence of steam inside the eutectic molten hydroxide. It can also be observed from the above figure that a significant increase in the current density of stainless steel as the working electrode, occurs in the presence of steam. This increase indicates that the electro-catalytic activity of stainless steel under these condition mirrors the value of nickel metal.

On the other hand, platinum still ranks as the most electro-active for the hydrogen evolution reaction. Despite this, its use was generally limited in history because it is classified as a precious metal in comparison to the others. Similar to steam introduction to increase HER strategy, doping strategy of nanosheets and other conducting materials with heteroatom to increase the electrocatalytic activity and resultantly increase HER, was also applied in literature [43-45] with a positive outcome. In addition to nanosheets, nanocrystals of trimetallic alloy [46] were also used for HER with high catalytic power. In this study and in other mentioned ones the main focus is the electrocatalytic activity of the materials/ or electrodes which directly plays a key role in HER.
Conclusions

The aims behind this detailed research were to find cheaper, electrocatalytic working electrodes, vs a novel Ni/Ni(OH)$_2$ reference electrode, that can be used to increase the feasibility of hydrogen gas production in eutectic molten hydroxide (NaOH-KOH, 49–51 mol%), at 300 °C temperature.

The most important findings that can be drawn from the results are:

- The reduction potential of the hydrogen evolution reaction using different working electrodes was in the order of (more positive to negative reduction potential): Pt > Ni > St.st > Ag > Mo. The performance of each working electrode for the hydrogen evolution reaction was confirmed through chronoamperometry tests at a constant potential of -0.7 V. These tests confirm the stability and productivity of each working electrode. The produced chronoamperograms found that the platinum had the highest current density followed by nickel, stainless steel, silver and then molybdenum at the constant potential of -0.7 V.

- It was also found from the cyclic voltammograms that the presence of steam inside the eutectic molten hydroxide is apparent in increasing the current density at the cathodic limit for the hydrogen evolution reaction. However, the starting point of reduction potential for the hydrogen evolution reaction was still approximately the same with and without the presence of steam inside the eutectic molten hydroxide.

- The effect of increasing the operating temperature of the eutectic molten hydroxide influenced the performed cyclic voltammetry scans. This effect appeared to clearly shift the reduction potential in a positive direction at high temperatures. This positive shift was applicable for all tested working electrodes. The shift in the reduction potential with an increase in the operating temperature was approximately 0.1 V for all tested working
electrodes. This was despite the fact that each one had a different reduction potential for the hydrogen evolution reaction.

Acknowledgement

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**List of Tables**

Table 1. Working electrode’s surface area specifications.

<table>
<thead>
<tr>
<th>Working electrode</th>
<th>Diameter (cm)</th>
<th>Surface area (cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nickel</td>
<td>0.05</td>
<td>0.22</td>
</tr>
<tr>
<td>Platinum</td>
<td>0.05</td>
<td>0.22</td>
</tr>
<tr>
<td>Silver</td>
<td>0.10</td>
<td>0.44</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>0.10</td>
<td>0.44</td>
</tr>
<tr>
<td>Stainless steel</td>
<td>0.025</td>
<td>0.11</td>
</tr>
</tbody>
</table>
**Table 2.** Reduction potential and the current limit at cathodic limit.

<table>
<thead>
<tr>
<th>Working electrode</th>
<th>Temperature (°C)</th>
<th>Reduction potential $E_{\text{red}}$ (V)</th>
<th>Current density $j$ (A cm$^{-2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni</td>
<td>300</td>
<td>-0.49</td>
<td>-1.67</td>
</tr>
<tr>
<td>Pt</td>
<td>300</td>
<td>-0.47</td>
<td>-1.23</td>
</tr>
<tr>
<td>Ag</td>
<td>300</td>
<td>-0.53</td>
<td>-0.20</td>
</tr>
<tr>
<td>Mo</td>
<td>300</td>
<td>-0.55</td>
<td>-0.16</td>
</tr>
<tr>
<td>St.st</td>
<td>300</td>
<td>-0.51</td>
<td>-1.41</td>
</tr>
</tbody>
</table>
Fig. 1. Experimental setup: (1) Argon inlet, (2) Steam inlet, (3) Reference electrode, (4) Working electrode, (5) Counter electrode, (6) Argon outlet, (7) Reaction vessel, (8) Corundum crucible and (9) Molten salt.
Fig. 2. Voltammetric peaks of a 0.5 mm nickel working electrode in the eutectic molten hydroxide at a temperature of 300 °C. RE: Ni/Ni(OH)₂; CE: 5 mm stainless steel rod; atmosphere of Ar gas at 40 cm³/min; the immersion depth: 14 mm; scan rate: 100 mVs⁻¹, (a) Scan negatively between -1.0 and 1.0 V, (b) Limiting the scan between -0.8 and -0.1 V, (c) Scan negatively between -0.8 and 1.0 V for steam analysis, (d) Limiting the scan between -0.8 and -0.3 V for steam analysis.
Fig. 3. Voltammetric peaks of a 0.5 mm Platinum working electrode in the eutectic molten hydroxide at a temperature of 300 °C. RE: Ni/Ni(OH)$_2$; CE: 5 mm St.st rod; an Ar gas atmosphere at 40 cm$^3$min$^{-1}$; the immersion depth: 14 mm; Scan rate: 100 mV s$^{-1}$; (a) Scan negatively between -0.8 and 0.5 V, (b) Limiting the scan between -0.8 and -0.3 V, (c) Scan negatively between -0.8 and 0.5 V for steam analysis, (d) Limiting the scan between -0.8 and -0.3 V for steam analysis.
Fig. 4. Voltammetric peaks of a 1.0 mm silver working electrode in the eutectic molten hydroxide at scan rate of 100 mVs\(^{-1}\) and operating temperature of 300 °C. RE: Ni/Ni(OH)\(_2\); CE: 5 mm St.st rod; an Ar gas atmosphere of 40 cm\(^3\) min\(^{-1}\); the immersion depth: 14 mm; a) Scan negatively from -0.8 to 1.0 V, (b) Limiting the scan between -0.8 and -0.3 V, (c) Scan negatively from -0.8 to 1.0 V for steam analysis, (d) Steam analysis by limiting the scan between -0.8 and -0.3 V.
**Fig. 5.** Voltammetric peaks of a 1.0 mm molybdenum working electrode in the eutectic molten hydroxide at a temperature of 300 °C and a scan rate of 100 mV s⁻¹. RE: Ni/Ni(OH)₂; CE: 5mm St.st rod; an Ar gas atmosphere at 40 cm³min⁻¹; the immersion depth: 14 mm: (a) Scan negatively from -0.8 to 0.1 V, (b) Limiting the scan between -0.8 and -0.2 V, (c) Scan negatively from -0.8 to 0.1 V for steam analysis, (d) Limiting the scan between -0.8 and -0.3 V for steam analysis.
Fig. 6. Voltammetric peaks of a 0.25 mm stainless steel working electrode in the eutectic molten hydroxide at a temperature of 300 °C. RE: Ni/Ni(OH)$_2$; CE: 5 mm St.st rod; an atmosphere of Ar at 40 cm$^3$ min$^{-1}$; the immersion depth: 14 mm; Scan rate: 100 mV s$^{-1}$. (a) Scan negatively from -0.8 to 0.3 V, (b) Limiting the scan between -0.8 and -0.3 V (c) Scan negatively from -0.8 to 0.3 V for steam analysis, (d) Steam analysis by limiting the scan between -0.8 and -0.3 V.
Fig. 7. Comparison of cyclic voltammograms of Ni, Pt, Ag, Mo, St.st working electrode in the eutectic molten hydroxide at a temperature of 300 °C and scan rate of 100 mV s\(^{-1}\). RE: 0.5 mm of Ni/Ni(OH)\(_2\) and CE: 5 mm of St.st; immersion depth: 14 mm, an Ar gas atmosphere: 40 cm\(^3\) min\(^{-1}\).
Fig. 8. Chronoamperograms of the hydrogen evolution reaction for all working electrodes (Ni, Pt, Ag, Mo, St.st) in the eutectic molten hydroxide at a temperature of 300 °C, and at an applied potential of -0.7 V
during 10 min; (a) Without steam and argon gas atmosphere at 40 cm$^3$min$^{-1}$, (b) With steam and argon gas atmosphere at 40 cm$^3$min$^{-1}$.