SURFACE MODIFIED AND BISMUTH LOADED GRAPHITE FELTS FOR IMPROVEMENT OF ANODE ELECTRODE KINETICS IN IRON CHROMIUM REDOX FLOW BATTERY

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Abstract

In this study, anode electrodes were modified and bismuth loaded to enhance chromium redox reaction for iron chromium redox flow batteries. Graphite felt anode electrode surface was modified by two step chemical treatment. Corrosive HF was used as an etchant to increase porosity at the first step and oxidizing H2O2 was used to increase wettability of electrode at the second step. Scanning electron microscopy (SEM) and X-ray photoelectron spectroscopy (XPS) analyses were used for the physical and chemical characterization of electrode materials. Cyclic voltammetry (CV) and linear sweeping voltammetry (LSV) techniques were conducted to investigate \( \text{Cr}^{3+/2+} \) redox reaction kinetics for the modified electrodes with or without bismuth loading. Carbon nanoparticles formed on the surface and oxygen content increased. Contact angle for pristine graphite felt decreased from 132.6° to 128.5°, electrode wettability increased after treatment. Bi was loaded onto the felt electrodes, reaction became diffusion controlled. In contrast, when bismuth was loaded onto the pristine electrodes, reaction became diffusion controlled.

Keywords: Graphite Felt, Acidic Modification, \( \text{Cr}^{3+/2+} \) Electrode Kinetics, Fe/Cr Redox Flow Battery

DEMIR KROM REDOKS AKIŞ PİLLERİNDE ANOT ELEKTROT KİNETİĞİNİN İYİLEŞTİRİLMESİ İÇİN YÜZEY MODİFİKASYONU YAPILmiş VE BİZMUT YÜKLENmiş GRAFİT KEÇELER

Özett

Bu çalışmada, demir krom redoks akış pilleri için krom redoks reaksiyonunu iyileştirmek amacıyla anot elektrotuna modifikasyon yapılmış ve bizmut yüklenmiş. Grafit keçe anot elektrot yüzeyi iki aşamalı kimiyasal işlem ile modifiye edilmiştir. İlk basamakta korozif HF aşındırıcı ajan olarak kullanılmış, ikinci basamakta ise oksitleyici H2O2 sıranalabilirliğin artırılması için kullanılmıştır. Taramalı elektron mikroskopu (SEM) ve X-sınu fotoelektron spektroskopisi (XPS) analizleri elektrot malzemelerinin fiziksel ve kimiyasal karakterizasyonunda kullanılmıştır. Bizmut yüklemesinin olduğu ve olmadığı durumda \( \text{Cr}^{2+/3+} \) redoks reaksiyonu kinetiğinin analizləmsında çevrimsel voltametri (CV) ve doğrusal süpürcü voltametri teknikleri kullanılmıştır. Yüzeyde karbon nanoparçacıkları oluşmuş, oksijen içeriği artmıştır. İşlem görmemiş grafit keçe için temas açısı 132.6°den 128.5°ye düşmüş, HF ve H2O2 ile işlem sonucunda elektrot sıranalabilirliğine artmıştır. Bizmut nanoparçacıklarının dağılımı modifiye edilmiş elektrot üzerinde daha homojen hale gelmiştir. İşlem görmemiş elektrotta kayıpla, \( \text{Cr}^{3+} \) için difüzyon kataysı modifiye edilmiş elektrot üzerinde neredeyse iki katına çıkmış, yüzeydeki \( \text{Cr}^{2+} \) konsantrasyonu artmıştır. Modifiye edilmiş elektrot için bizmut çözeltide olduğunda, reaksiyon yük transferi ile kontrol edilmektedir. Aksine, bizmut keçe elektolara yükleniyorsa reaksiyon difüzyon kontrol olmaktadır.

Anahtar Kelimeler: Grafit Keçe, Asidik Modifikasyon, \( \text{Cr}^{2+/3+} \) Elektrot Kinetiği, Fe/Cr Redoks Akış Pili

Cite

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1. Introduction

Due to industrial and technological developments, the world’s electrical energy demand is accelerating and usable fossil fuel reserves are decreasing considerably. Intermittent renewable energy sources such as wind and solar power cause fluctuations problem during the day depending on the weather conditions. Electrical energy storage (EES) can be considered as a satisfactory solution to compensate the energy imbalance and consequently electrical energy can be supplied when needed. It is possible to store electrical energy in different forms in different systems. Electrical energy can be stored mechanically in the form of kinetic energy, mechanically in the form of potential energy, chemically (fuel cells, flow batteries and secondary batteries) and thermally by low and high temperature processes. Among these systems, flow batteries are an alternative promising and developing electrical energy storage technology. In redox flow batteries, the energy density and the power density can be independently scaled. Energy capacity is related to the tank size and concentration of the electroactive species, while power capacity is related to active cell area and the number of cells [1,2]. One way to decrease cell size and cost of the flow battery is to develop high performance anode and cathode electrodes. Redox flow battery systems such as polysulfide bromide, Iron-Chromium, Vanadium Redox, and Hydrogen-Bromine are promising electrochemical energy storage technologies. Considering iron/chromium redox flow battery (ICRFB), iron and chromium based anolyte and catholyte solutions are used at the anode and cathode. These reactants are abundant and cheap materials, as well as ICRFBs have benefit according to economic considerations compared to vanadium redox flow batteries (VRFBs) when operated at the large scale as in smart grid applications [3].

\[
\begin{align*}
\text{Cr}^{2+} + e^- & \rightarrow \text{Cr}^{3+} & 0.41 \text{ V vs SHE} \\
\text{Fe}^{3+} + e^- & \rightarrow \text{Fe}^{2+} & 0.77 \text{ V vs SHE} \\
\text{Cr}^{2+} + \text{Fe}^{3+} & \rightarrow \text{Cr}^{3+} + \text{Fe}^{2+} & 1.18 \text{ V vs SHE}
\end{align*}
\]

Kinetics of the Cr side redox reaction is slow and H₂ evolution reaction occurs during the charging process which competes with the Cr³⁺/Cr²⁺ reduction reaction. For the Fe/Cr flow batteries [4], Electroactive well-designed electrodes can improve the anode reaction kinetics while reducing hydrogen gas output. Among the carbon based electrodes, usage of graphite felt (GF) is more common owing to 3D structure, good electrochemical activity, chemical stability, and low cost [5, 6]. Various electrode modifications such as thermal treatment [7, 8], acidic treatment [9, 10], and thermal treatment [11], electrochemical oxidation of GFs in acid solution [12], and O₂ plasma treatment [13] are considered to enhance the surface properties of the electrode. Amount of surface functional groups, wettability of the electrode and consequently catalytic activity for redox reactions can be increased with the help of these treatments. These functional groups also facilitate the Cr³⁺/Cr²⁺ redox reaction specifically [14–16]. One of the other methods to increase catalytic activity is the usage of certain metal catalysts by internal [17] or external loading [18–20] over the porous transport layer (e.g. graphite felt). Especially, bismuth shows catalytic activity for both VRFBs [17–20] and Cr³⁺/Cr²⁺ redox reaction [15, 16, 21]. Increasing the surface roughness and area are not a satisfactory solution, modification method which has a favorable effect on both surface area and catalytic activity is seen as more beneficial [22]. In other words, usage of metal catalyst with a rough or porous electrode material can be more favorable. Porosity of carbon fiber-based electrodes mainly comprises voids between fibers and their surface area is quite low. Increase in surface area and porosity is not sufficient after treatments stated above. He et al. [23] used HF/H₂O₂ treatment for GFs in VRFB. In their study, HF served as an etchant to increase available sites for redox reactions and H₂O₂ served as an oxidant. As a result, oxygen-containing functional groups and electrochemical activity of the electrode increased. Zhou et al. [24] used KOH activation for carbon paper electrode and obtained bimodal porosity. They increased the surface area of the electrode one order of magnitude compared to bare carbon paper. Zhou et al. also reported that bigger pores form pathways for electrolyte flow and smaller pores which connected to bigger ones contains active sites. In another study using KOH as the activating agent for GF, it was reported that edge carbon sites and functional groups were formed on the electrode surface, surface area was increased, and enhanced flow battery performance was achieved with the modified electrode for VRFBs [25]. Among these summarized studies HF/H₂O₂ treatment is the promising one since porosity, surface area, wettability and as a result activity increase to a sufficient level. In the present study, two step HF/H₂O₂ modification was used in order to investigate effect of HF/H₂O₂ modified GF on the performance of Cr³⁺/Cr²⁺ redox reaction for the first time. Bismuth was loaded onto the GF electrodes by electrochemical deposition method. Electrocatalytic activity of the GF was tested with cyclic voltammetry and linear sweep voltammetry for electrolyte solutions with or without bismuth addition.

2. Experimental

PAN based GFD 2.5 EA graphite felt was supplied from SGL® group. CrCl₃·6H₂O, HF, H₂O₂ and Bi₂O₃ were purchased from Sigma Aldrich, Merck, Tekkim and Alfa Aesar respectively. Graphite felts were pretreated prior to HF/H₂O₂ treatment. Firstly, felts were immersed in acetone and kept in ultrasonic bath for 1 h and then immersed in deionized water and kept in ultrasonic
bath for 1 h. Afterwards, graphite felts were dried at 80°C followed by 2 h heat treatment at 150 °C in an oven [26], [27]. Pretreated GFs were immersed in 12 wt.% HF for 36 h, washed with deionized water until neutral pH and dried at 80°C. Oxidation of the GF was performed with 5 wt.% H₂O₂ at 180°C for 10 h in a Teflon lined stainless steel autoclave. After H₂O₂ treatment GFs were washed with deionized water and dried at room temperature. Bismuth loading of 0.005 g/cm² on graphite felt was performed by electrochemical deposition method. Pt wire, Ag/AgCl (3 M NaCl) and GF were used as counter electrode, reference electrode, and working electrode respectively. -0.5 V voltage was applied for the electrochemical deposition of bismuth. After deposition, graphite felts were thermally treated at 450°C for 0.5 h. Pristine and modified graphite felt based electrodes prepared for voltammetry tests were named as Bi/GF and Bi/HF-GF.

Scanning electron microscopy (SEM, JEOL JSM-7001F) and X-ray photoelectron spectroscopy (XPS, PHI Versaprobe 5000, reference: 284.5 eV binding energy) analysis were carried out to analyze textural and chemical properties of the prepared pristine GF, HF/GF, and Bi loaded graphite felts (Bi/GF and Bi/HF-GF). Contact angle measurement was achieved by dynamic method for ten minutes, 5μl water droplet was used on test. Electrochemical performance of felts on Cr³⁺/Cr²⁺ redox reaction was tested by cyclic voltammetry (CV) (Basi RDE-2 stand and Ivium potentiostat). Felt, platinum wire and Ag/AgCl (3 M NaCl) were used as working, counter and reference electrode respectively. Felts were immersed in ethanol for 1 h, later in deionized water for 1 h and finally in electrolyte solution before CV tests as suggested by Smith et al. [28]. GF electrodes were hanged with Pt wire like fish hook (Fig. 1). Electrolyte solutions were 1M HCl, 1M HCl + 1M CrCl₃ and 1M HCl + 1M CrCl₃ + Bi₂O₃. Scanned potential range was 1 to -1.2 V except voltammograms of bismuth free electrodes, for these electrodes scanning was conducted between 1 to -0.2 V. H₂ evolution on felts was monitored by linear sweeping voltammetry by using only acidic media (1 M HCl) in the potential range of 0 to -1.2 V. Scanning rate was kept at 100 mV/s in general and in some experiments altered to analyze electrochemical effects resulting from scan rate.

Figure 1. Electrodes used for CV, LSV and installation of working electrode.

3. Results and Discussion

3.1. Characterization of electrodes

The surface morphology of the graphite felts was detected with scanning electron microscopy analysis. Graphite felt fiber diameter for the modified graphite felt decreases as a result of corrosion effect of the HF (Figure 2 (a) and Figure 2 (c)). After HF and H₂O₂ modification carbon nanoparticles which are broken off from felt occur as reported in a previous study [23](Figure 2 (d)). The formation of such carbon particles on the electrode surface is expected to increase the surface area and porosity by the formation of indentations on the surface as a result of acid and peroxide modification. In addition to increase in surface area and porosity of the electrode, bonding of the oxygen containing groups with H₂O₂ treatment is desirable since these groups provide advantages such as wettability and catalytic active sites for chromium redox reaction [29].

Figure 2. SEM micrographs of GF (a, b) and HF-GF (c, d) electrodes.

Bismuth is seen as attached to the electrode surface from electrochemically bismuth deposited electrode's SEM images (Figure 3). Bismuth surface density could be seen from mapping results and Bi particles loaded on the GF electrode are observed as agglomerated on the surface (Figure 3 (b) inset). On the other hand, bismuth particles are smaller and homogeneously dispersed on the Bi/HF-GF electrode compared to Bi/GF electrode thanks to the hydrophilic structure of the HF-GF electrode (Figure 3 (d)). In addition to hydrophilic structure, carbon nanoparticles on the HF-GF electrode are thought to allow effective distribution of bismuth on the surface.

XPS spectrum was obtained for the determination of surface chemistry and functional groups. Wide scan XPS
survey spectra between 0-1300 eV was given in Figure 4. Peak position of the carbon 1s is at 284.6 eV binding energy and surface elemental composition is found as oxygen and carbon. O1s peak intensity increases for the HF-GF electrode and oxygen percentage is 3.7% higher than the pristine GF (Table 1).

Curve fitted C1s and O1s spectrums were given in Figure 5 and carbon in C1s spectra and oxygen in O1s spectra has various electronic states. Functional groups on the surface and binding energies according to C1s spectra as follows: C=C (284.0, 283.75 eV) [30], C-C (284.7, 284.9 eV) [10], [13], [27], [31], C=O (286.0, 285.75, 285.9 eV) [10], [12], [13], [31], C=O (287.0, 288.0, 529.9, 531.1, 531.3, 531.8 eV) [10], [31]-[34] (10.31-34), carboxyl group (288.8, 289.75, 291.0 eV) [33], [35], -OH (532.8 eV) [35] and water (533.6, 534.2 eV) [31], [32].

Table 1. The contents of carbon, oxygen and surface functional groups of GF and HF-GF electrodes obtained from XPS data

|            | C1s (%) | O1s (%) |
|------------|---------|---------|
| C=C        | 15.3    | 43.1    |
| C=C        | 32.9    | 43.1    |
| C-OH       | 29.9    | 26.2    |
| O=C-OH     | 9.4     | 8.3     |
| C=O        | 6.2     | 6.2     |
| OH         | 50.6    | 50.6    |
| H2O        | 18.5    | 18.5    |
| H2O        | 30.9    | 30.9    |

Percentage of functional groups obtained after deconvolution were given in Table 1. C=C group percentage increases after chemical modification. The carbon double bond structure reduces to a single bond and thus the percentage of C=C may be higher as a result of the occurrence of oxygen containing groups. Unlike C-OH percentage, the amount of carboxyl group increases. This suggests that C-OH forms may be converted into carboxyl groups. C=O and water percentages increase for HF-GF electrode according to O1s spectra. Adsorbed water increases on the surface a result of acquired hydrophilic property of HF-GF. The presence of acidic oxygen containing groups on the surface increases the hydrophilic character of the structure, thus these groups facilitate access to the active areas for the electrolyte [10], [14], [23]. Moreover, it is known that the presence of -OH groups in the structure facilitate electron transfer [7], [9].
Wettability tests were achieved by dynamic contact angle measurement and showed in Figure 6. Initial contact angle for pristine GF is 132.6° and it decreased to 128.5° after modification. Absorption of water droplet for HF-GF electrode is faster than the pristine one with the help of oxygen containing groups on the HF-GF (Figure 6).

3.2. Application of electrodes for Cr side of Fe/Cr flow battery

Cyclic voltammetry was used to investigate the reduction and oxidation behavior of chromium ions and bismuth ions, proton adsorption and desorption on the electrode surface. Voltammograms obtained with the GF and HF-GF electrodes in 1 M HCl electrolyte were given in Figure 7. Potential sweeping was performed from the positive potential to the negative potential at the scanning rate of 100 mV/s between 1.0 V and -0.2 V to observe proton adsorption and desorption on the surface. Peaks for the proton adsorption and desorption are seen at 0.075 V and 0.875 V for the GF electrode and 0.175 V and 0.825 V for the HF-GF electrode respectively [36]. It is seen that the peaks observed for HF-GF electrode approach each other and the potential difference between peaks (ΔE_p) decreases as 0.15 V compared to GF electrode. This shows that the proton adsorption desorption becomes more reversible on the HF-GF electrode. The increase in peak current density for the HF-GF electrode is related to the higher proton concentration on the surface of the HF-GF electrode, which has hydrophilic structure [10].
the HF-GF electrode and the peak position is seen at more negative value. It is thought that, bismuth peak is more severe hence surface of the fibers is covered with bismuth for Bi/HF-GF electrode. In addition, this peak covers the potential value of Cr$^{3+}$ anodic peak. Although the Cr anodic peak cannot be distinguished on Bi/GF and Bi/HF-GF electrodes clearly, the current density value around 0.5 V is higher on the Bi/HF-GF electrode. According to the Randles-Sevck equation (Equation 1), the diffusion coefficient can be obtained from the slope of the line when scan rate is plotted against the current [11], [37]. In this equation A is electrode area, D is diffusion coefficient, n is number of transferred electrons, $\theta$ is scan rate, $C_0$ is concentration, $\alpha$ is charge transfer coefficient (0.5) and $I_p$ is peak current.

$$I_p = \frac{2.69 \times 10^5 A D^{1/2} n^{3/2} \theta^{1/2} C}{10^6}$$ (1)

The CV curves for GF and HF-GF electrodes at different scan rates in 0.5 M CrCl$_3$ + 1 M HCl solution were given in Figure 9. The high intensity of the Cr$^{3+}$ anodic peak on the HF-GF electrode at different scan rates indicates that the polarization is lower on HF-GF electrode. The linearity of the lines shows that the reaction is controlled by charge transfer for both GF and HF-GF electrodes [31]. In the peak current against the square root of scan rate graphs, the slope of the line is higher in the HF-GF electrode and diffusion coefficient on the HF-GF electrode is twice that of the GF electrode. Diffusion coefficients of Cr$^{2+}$ on GF and HF-GF electrodes are 2.3x10$^{-6}$ cm$^2$/s, 4.7x10$^{-6}$ cm$^2$/s respectively. Cr peak on the GF electrode could not be distinguished in CV curves taken at different scan rates in the bismuth containing 0.5 M CrCl$_3$ + 1 M HCl solution, so the graph could not be plotted. Cr peak could not be seen due to ineffectiveness of Cr$^{2+}$ ions diffusion on GF electrode. On the HF-GF electrode, the graph is plotted because the current density value for Cr$^{3+}$ is higher and it can be seen that the slope of the line in this graph is decreased compared to the bismuth free condition. The diffusion constant for Cr$^{2+}$ ions is approximately four times lower (0.8x10$^{-6}$ cm$^2$/s) than the bismuth free condition when bismuth ions are present in solution. On the other hand, charge transfer becomes easier in the bismuth containing electrolyte solution [31]. Although the peak separation could not be made for Cr when the bismuth catalyst is in solution, the current density value decreases linearly with the decrease in the voltage scan rate. On the contrary, when the bismuth catalyst is loaded to electrode, the decrease in peak current does not show a linearity. This showed that reaction becomes diffusion controlled when the bismuth catalyst is loaded on the felt.

Cyclic voltammetry curves (CVs) for HF-GF working electrode in both bismuth containing and bismuth free chromium electrolyte and Bi/HF-GF working electrode in bismuth free chromium electrolyte were given in Figure 8 (b). In the CV curve obtained in bismuth-free conditions, Cr$^{2+}$/Cr$^{3+}$ anodic peak is seen around 0.5 V. The current density value of this peak increases by 20 mA/cm$^2$ compared to the untreated felt on the HF-GF electrode. In addition, there is a relatively small hysteresis opening around -0.2 V. The hysteresis in this region may be due to the interaction between hydrogen and acidic groups on the HF-GF electrode or may be due to the Cr$^{3+}$ reduction reaction. In the case which bismuth catalyst is in solution, bismuth cathodic peak and anodic peak are seen around -0.2 V and 0.05 V respectively and their positions shift to 0.1 V right compared to GF electrode as seen. The fact that the anodic peak shifts to the right side reduces the reversibility, this does not seem to be a problem since the oxidation of bismuth is not desirable.

When the Bi/HF-GF and Bi/GF electrodes compared, the intensity of the bismuth anodic peak is much higher on the HF-GF electrode and the peak position is seen at more negative value. It is thought that, bismuth peak is more severe hence surface of the fibers is covered with bismuth for Bi/HF-GF electrode. In addition, this peak covers the potential value of Cr$^{3+}$ anodic peak. Although the Cr anodic peak cannot be distinguished on Bi/GF and Bi/HF-GF electrodes clearly, the current density value around 0.5 V is higher on the Bi/HF-GF electrode. According to the Randles-Sevck equation (Equation 1), the diffusion coefficient can be obtained from the slope of the line when scan rate is plotted against the current [11], [37]. In this equation $A$ is electrode area, $D$ is diffusion coefficient, $n$ is number of transferred electrons, $\theta$ is scan rate, $C_0$ is concentration, $\alpha$ is charge transfer coefficient (0.5) and $I_p$ is peak current.

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Linear sweeping voltammetry graphs for monitoring hydrogen evolution on GF, HF-GF, Bi/GF and Bi/HF-GF electrodes show that hydrogen evolution is suppressed better on HF-GF electrode (Figure 10). This is directly related to the oxygen containing groups formed on the
HF-GF electrode. Formation of hydrogen is inhibited by bonding protons with oxygen groups located on the electrode surface. The proton concentration on the surface of the HF-GF electrode is higher as previously mentioned (Figure 7). Metallic bismuth and protons form BiH$_3$ solid as given below. As a result, bismuth inhibits evolution of hydrogen gas by forming BiH$_3$ solids [19], [20], [32].

\[
\begin{align*}
\text{Bi}^{3+} + 3e^- &\rightarrow \text{Bi} & 0.308 \text{ V vs SHE} \\
& & 0.109 \text{ V vs. Ag/AgCl (3M NaCl)} \\
\text{Bi} + 3\text{H}^+ + 3e^- &\rightarrow \text{BiH}_3 & -0.8 \text{ V vs SHE} \\
& & -1.0 \text{ V vs. Ag/AgCl (3M NaCl)}
\end{align*}
\]

Bi/HF-GF electrode is better than Bi/GF electrode to suppress hydrogen evolution. The change in current density around -1.0 V is considered to be caused by formation of BiH$_3$ solid on the Bi/HF-GF and Bi/GF electrodes. Although the formation of hydrogen gas on the bismuth free electrodes appear to be less in the LSV curves, it is thought that, bismuth loaded electrodes may be more effective to suppress hydrogen evolution for long term operations.
Figure 9. CV curves of GF (a), HF-GF (b) in 1M HCl + 0.5M CrCl$_3$ electrolyte, GF (c), HF-GF (d) in 1M HCl + 0.5M CrCl$_3$ + 0.005 g Bi/cm$^2$ electrolyte, Bi/GF (e), Bi/HF-GF (f) in 1M HCl + 0.5M CrCl$_3$ electrolyte at different scan rates.
4. Conclusion

In summary, corrosive HF and oxidizing H₂O₂ were used to obtain a porous and hydrophilic electrode structure. The effect of electrode structure and double acting bismuth material which shows feature as catalyst and hydrogen evolution suppressor agent on the Cr²⁺/Cr³⁺ redox pair were investigated. Contact angle for graphite felt decreased to 128.5° after modification. The wettability of the felt treated with HF and H₂O₂ increased and as a result, reactant concentration on the surface increased. Hydrogen evolution decreased compared to other electrodes with the oxygen containing groups formed on the surface of the HF-GF electrode. However, bismuth loaded HF/GF is going to be better to prevent hydrogen formation since BiH₃ solid consists on the electrode. GF and HF-GF electrodes are charge transfer controlled but the diffusion coefficient for the HF-GF electrode is twice compared to GF electrode. Although the peak separation for Cr could not be seen clearly when the bismuth catalyst was in electrolyte, the current density value decreased linearly with the decrease in the voltage scan rate. This indicates that the reaction is still under charge transfer control. On the contrary, when the bismuth Catalyst is loaded onto the felt, the reaction is controlled by diffusion.

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