Electrochemical Water Treatment Technology in Viet Nam: Achievement & Future Development

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Abstract: Electrochemical technology is a promising technique for water and wastewater treatment in the world. This technology has many advantages as compact model, saving area, highly removal efficient of non-biodegradable compounds, reduction of secondary by-products. There are some different electrochemical techniques such as electroflotation; electrocoagulation using sacrificial electrodes; direct or indirect electrochemical oxidation using Dimensionally Stable Anodes (DSA) and electrodialysis to separate ions in water. In Vietnam, some researches on water and wastewater treatment by electrochemical technology have been studied in the recently year. In this paper, notable results of studies in the application of water and wastewater treatment by electrochemical technology in Vietnam are reviewed in detail. The researching trends of the authors mainly focused on the laboratory scales and the simulated wastewater. Therefore, there are not enough scientific basis to apply for real treatment plants. In the future, electrochemical methods should be used to study the different types of wastewater at the pilot scales, especially difficult handling wastewater and toxics that biological methods cannot handle effectively.

Keywords: Water Treatment, Electrocoagulation, Electroflotation, Electrooxidation, Electrodialysis

1. Introduction

Nowadays, the industrial development has achieved many significant goals for the economic growth in Viet Nam. In addition, this process discharged waste into the environment in general and water resource in particular. At present, common methods used for treating wastewater are physical, chemical and biological methods. However, these methods have not always achieved satisfactory results with increasingly stringent discharge regulations. The wastewaters containing heavy metals, phenolic compounds, and non-biodegradable pollutants are very difficult to handle and easily retained in the receiving source, which affect human health. On other hand, traditional physical and chemical methods have relatively high process costs, ineffective or can lead to secondary pollution [1]. Therefore, how to eliminate these biodegradable components in wastewater are essential.

Over the last two decades, electrochemical technology has been studied in the field of wastewater treatment in the world. Many studies have been conducted in the laboratory and some good results have been commercialized. Electrochemical treatment has the following advantages: direct or indirect oxidation flexibility; space saving; high-performance processing and can be integrated with environmentally friendly methods [2]. Some types of wastewater have been studied such as: tanning [3, 4], pharmaceuticals [5, 6], toilet wastewater [7], textile [8, 9], sugarcane [10, 11], carwash [12], dairy [13] and some other wastewater types.

In Vietnam, over the past ten years, electrochemical techniques in water and wastewater treatment have also been studied. However, this issue is relatively novel and limited in the application. In this article, the current state and evaluation...
of water treatment results in Vietnam by electrochemical will be summarized in detail. Then the suitable future research direction in this field in Viet Nam will be proposed.

2. Electrochemical Methods for Water Treatment

2.1. Electrocoagulation

Electrocoagulation is technology that is used long-standing and popular today. In this process, when current density is applied, the coagulants are created inside the reactor using sacrificial electrodes, usually iron or aluminum. The metal ions dissolved at the electrode will react to form hydroxide, polyhydroxide compounds that produce a coagulation processing. Chemical reactions in this process can be summarized below [14]:

a. The metal is oxidized into cations at the anode, according to the equation:

\[ \text{M} \rightarrow \text{M}^{z+} + z\text{e}^- \]  \hspace{1cm} (1)

b. Water is reduced into hydrogen gas and hydroxide anions at the cathode:

\[ 3\text{H}_2\text{O} + 3\text{e}^- \rightarrow 3/2 \text{H}_2 + 3\text{OH}^- \ (E^o = 0.00\text{V/ENH}) \]  \hspace{1cm} (2)

c. General reaction:

\[ \text{M}^{z+} + z\text{OH}^- \rightarrow \text{M(OH)}_z \]  \hspace{1cm} (3)

The advantages and disadvantages of this process are shown in Table 1.

| Advantages | Disadvantages |
|------------|---------------|
| Ability to treat drinking water and wastewater | Maintenance required regularly |
| Combination of oxidation and coagulation processes | The electrode breaks down over time |
| Reduced demand for chemicals (replaced by aluminum and iron electrodes) | Wastewater must have high conductivity |
| Reduce operating costs | |
| Reduce the risk of secondary pollutants | |
| Less generated sludge | |
| Low energy requirements | |
| Ability to use solar energy | |

2.2. Electroflotation

Electroflotation is a process that pollutants are loaded on the reactor surface by hydrogen and oxygen microbubbles produced by electrolysis of water at the cathode and anode. Generation of the hydrogen and oxygen bubbles are simulated by the equations below:

At anode: \[ 2\text{H}_2\text{O} = \text{O}_2 + 4\text{H}^+ + 4\text{e}^- \]  \hspace{1cm} (4)
At cathode: \[ 2\text{H}^+ + 2\text{e}^- = \text{H}_2 \]  \hspace{1cm} (5)
General reaction: \[ 2\text{H}_2\text{O} = 2\text{H}_2 + \text{O}_2 \]  \hspace{1cm} (6)

Several studies have combined electroflotation and electrocoagulation in a reactor tank. The coagulation processes occur at the anodes generated aluminum hydroxide ions or iron hydroxide ions which combined to the hydrogen and oxygen bubbles are generated at the anodes and cathodes to increase the efficiency of the process. [15].

2.3. Electrooxidation

2.3.1. Direct Oxidation

In order to reach high performance of direct oxidation process, the anode needs two characteristics: high oxidation potential and corrosion resistance. The oxidation of an organic compound at the anode is simulated in three steps [2]:

a. Discharge of water forming an adsorbed hydroxyl species:

\[ \text{S[ ]} + \text{H}_2\text{O} = \text{S[OH]} + \text{H}^+ + \text{e}^- \]  \hspace{1cm} (7)

b. The adsorbed OH is the ‘activated state’ of water in O-transfer reactions to the organic molecule R:

\[ \text{S[OH]} + \text{R} = \text{S[ ]} + \text{RO} + \text{H}^+ + \text{e}^- \]  \hspace{1cm} (8)

c. Co-evolution of \text{O}_2 by oxidation of water diminishing the current efficiency:

\[ \text{S[OH]} + \text{H}_2\text{O} = \text{S[ ]} + \text{O}_2 + 3\text{H}^+ + 3\text{e}^- \]  \hspace{1cm} (9)

2.3.2. Indirect Oxidation

Oxidants such as chlorine, hypochlorite, Fenton’s reagent, peroxide, peroxodisulphate and ozone are used for indirect oxidation at the anode [14]. The chlorine and hypochlorite are often used because they are easily generated at the anodes. The indirect oxidation mechanisms of the (R) organic compounds are described by the following equations [16]:

\[ \text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{HOCl} + \text{H}^+ + \text{Cl}^- \]  \hspace{1cm} (10)
\[ \text{HOCl} \rightarrow \text{H}^+ + \text{OCl}^- \]  \hspace{1cm} (11)
\[ \text{OCl}^- + \text{R} \rightarrow \text{CO}_2 + \text{inorganic ions} + \text{H}^+ + \text{e}^- \]  \hspace{1cm} (12)
\[ \text{OCl}^- + \text{RCl} \rightarrow \text{R} \]  \hspace{1cm} (13)

2.3.3. Electroreduction

Similar to electrooxidation method, the electroreduction also has two groups: direct and indirect. The advantages of this direct process are the reducing nitrate at the cathode to generate nitrogen gas and hydroxide ions. However, some undesirable byproducts such as nitrite and ammonia are also produced during the nitrate reduction process. These two byproducts are also involved in the process in which the nitrite is reduced at the cathode, ammonia is oxidized at the anode according to the following equations [16]:

\[ \text{NO}_3^- + 3\text{H}_2\text{O} + 5\text{e}^- \rightarrow \frac{1}{2} \text{N}_2 + 6\text{OH}^- \]  \hspace{1cm} (14)
\[ \text{NO}_2^- + 2\text{H}_2\text{O} + 3\text{e}^- \rightarrow \frac{1}{2} \text{N}_2 + 4\text{OH}^- \]  \hspace{1cm} (15)
\[ 2\text{NH}_3 + 6\text{OH}^- \rightarrow \text{N}_2 + 6\text{H}_2\text{O} + 6\text{e}^- \]  \hspace{1cm} (16)

The electroreduction is often used for the treatment of
organic dyes that are insoluble in water and poor contact with the electrode surface. It dissolves some chemical substances such as Anthraquinone, Fe-TEA and Glucose to enhance the solubility of dyes and decomposition by indirect reduction. [12].

2.4. Electrodialysis

Electrodialysis (ED) is an electrochemical process. This process involves the movement of ions through selective ion-exchange membranes, which are the results of electrostatic attraction between the ion and the two electrodes. In addition to the desalination function, ED can also remove contaminants in water as soluble ions by electrostatic attraction [17].

The advantage of ED is that the recovery rate of water is higher than the Reverse osmosis (RO) process, leading to lower wastewater treatment costs and increased water using efficiency. In addition, the life of the ion-exchange membrane is relatively high. The ED technology will require less energy than the RO when the salinity is less than 5000 mg/l [17].

3. Results of Water Treatment by Electrochemical Methods in Viet Nam

3.1. Electroflotation and Electrocoagulation

3.1.1. Electroflotation

The electroflotation technology in Vietnam was investigated for pre-treatment of wastewater before aerobic steps. The wastewater output has a COD removal efficiency as 70-80%, Nitrogen removal efficiency as 50-60%. Although this quality has not reached the standard for direct discharge to receiving sources, it is generally feasible for the thorough treatment process of pollution compounds in the next units. Dinh Quang Du et al. (2015) [18] was investigated electroflotation process for pre-treatment of pesticide production wastewater. In this study, the operating parameters of the electroflotation model include: 45° of angle electrode, aluminum electrode with 486 cm² square, 1 cm of electrode gap, retention time was 30 minutes and 12 V electrolytic voltage, H₂O₂ concentration was 80 mg/L and 30 mg/L FeCl₃ concentration. The results showed that SS, COD, BOD₅, TKN, TP removal efficiency were obtained as 83%, 76%, 57%, 53% and 82%, respectively. After the flotation process, the dissolved oxygen (DO) value in water increases. Compared with the results of using aerobic bioreactor rollers, the pre-treatment of electrofloation is more effective, which satisfies the requirements for the next biological treatment. Similarly, electrofloation process is applied at the pre-treatment stages, Le Hoang Viet et al. (2015) [19] evaluated the technical feasibility of applying this method for wastewater treatment system in catfish processing facilities. The experimental conditions are similar to the research of Dinh Quang Du. Under these experimental conditions, the SS, COD, BOD₅, TKN, TP removal efficiencies were obtained as 83.2%; 76.8%; 68.3%; 66.9% and 71.4%, respectively. The electrofloation process showed that during the reaction, the metal material on the electrode is decomposition at same performances. Therefore, the bigger the electrode size, the higher removal efficiency.

3.1.2. Electrocoagulation

Comparison of anthraquinone dye by two methods: electrocoagulation and electro-Fenton processes. The electrocoagulation uses an aluminum electrode with 0.5 mA/dm² current density, anodic area is twice cathode sizes, the stirring speed in the reaction is 1000 rpm, 0.5 g/L NaCl solution. During electrochemical reaction, aluminum anode dissolves to generate aluminum hydroxide which remove COD from wastewater. The COD removal efficiency decreased when increasing COD concentration. At the influent COD concentrations were 600 mg/l and 1040 mg/l for 97.33% and 86.92% performances respectively in 40 minutes. With the ratio of H₂O₂:FeSO₄ = 5:1 added to the Fenton reaction, the input COD concentrations were 600 mg/l and 1040 mg/l for 69.5% and 40% performances, respectively. This results illustrate when the dye concentration increase, the Fenton compounds are not sufficiently decomposed causing reduced efficiency. The comparable dyes removal time of two methods are shown in Figure 1, in the same time COD removal efficiency of electrocoagulation was higher than electro-Fenton reached approximately 90% performances after 25 minutes. The study also tested on the anthraquinone dye, the electro-Fenton processing efficiency is relatively better at 85% when the input dye concentration is 458 mg/l. The performance would depend on the types of dye and input concentration (Nguyen Thi Huong, 2009 [20]).

![Figure 1. COD removal performance of dyes wastewater (1. Electrocoagulation; 2. Electro-Fenton process [16]).](image-url)
Electrocoagulation was investigated in combination with micro electrocoagulation to remove simultaneously Pb^{2+}, Cu^{2+}, and F\textsuperscript{-} ions in wastewater by V'o An Khue (2014)\cite{23}. The optimum parameters for this method are as follows: initial pH value 5, 5V of voltage, removal time is 30 minutes, 60g of Fe-C, 20-27 mesh of size Fe-C. Under this condition and the initial concentration of ions is 50mg/L, Pb^{2+}, Zn\textsuperscript{2+}, Cu\textsuperscript{2+} and F\textsuperscript{-} removal efficiency are obtained 99.76%, 99.26%, 99.78%, 94.03%, respectively. After treatment, the wastewater is very clean and water quality is better than discharge standard, could be discharged directly to the receiving source. The mechanism of fluoride removal is mainly due to the formation of AlF\textsubscript{3}(OH)\textsubscript{3-x} precipitate and a small fraction of the fluoride absorption of iron hydroxide.

Nguyen Ngoc Anh et al. (2010)\cite{24} was carried out the integration of two technologies including USBF (Up-flow Sludge Blanket Filtration) and EC (Electrocoagulation) into once of aquatic wastewater treatment system. A total of eight experiments were conducted, and determined to parameters for design and operation of batch electrochemical tank combined to the USBF with substrates inside and without substrates. The results indicated that at 10 h of total retention time, the USBF performances without substrates were obtained as: 89.96%, 96.33%, 97.52%, 89.34%, 71.95% for SS, COD, BOD\textsubscript{5}, TKN, TP of parameters, respectively. The SS, COD, BOD\textsubscript{5}, TKN parameters meet QCVN 11: 2008/BTNMT (column A). The optimum conditions of this process are as follows: Anode should be made of aluminum material and cathode should be made of iron material, batch retention time for electrocoagulation tank works at 45 minutes, the distance between the two electrodes is 2 cm, electrode area: 100 cm\textsuperscript{2}, 24 V of voltages, 1.6 A of current density.

### 3.2. Electrooxidation

Nguyen Thi Lan Phuong et al. (2016)\cite{25} studied on wastewater containing reactive dye by electrochemical method with 304 stainless steel anode. Under conditions of pH = 6, 30 mA/cm\textsuperscript{2} of current density. After only 6 minutes of treatment, this method had COD and color effluents were achieved the QCVN 13: 2008/BTNMT (column B) standard with color influents as a range of 34-8453 Pt-Co and COD < 170 mg/L. The color removal efficiency during this study was relatively high, but the COD removal efficiency was dependent on the original concentration. The input COD concentration of 806 mg/L was achieved 17% removal efficiency. However, the input COD concentration less than 806 mg/L was achieved 70%. With anode and cathode as 304 stainless steel, the oxidation processes occur directly on the electrode surface or indirectly in solution through the OH\textsuperscript{-} radicals. In addition, the 304 stainless steel electrode also was dissipated during the process to form Fe\textsuperscript{3+} ions which involved in the electrochemical and electro-Fenton processes increasing the color removal efficiency of dye wastewater. The results of UV-Vis spectroscopy of influent and effluent showed that the azo groups and aromaticity of dye dissolved into simpler compounds, resulting in loss of color and reduced toxicity of the reactive dye.

Research on producing new electrodes on Titanium subract is also being done in Vietnam. Under the manufacturing conditions are a mixture of 0.5 M SnCl\textsubscript{4} + 0.5% SbCl\textsubscript{3} in C\textsubscript{2}H\textsubscript{5}OH + HCl solutions with 1:1 of rate and they are heated to 450°C. Dinh Thi Mai Thanh et al. (2007)\cite{26} have successfully manufactured the working electrode of SnO\textsubscript{2}-SbO\textsubscript{2}/Ti. Study on the Phenol removal efficiency in wastewater, after 7 hours of electrolysis with 10 mA/cm\textsuperscript{2} of current density was obtained low performance at 39%, 62.7% of COD removal efficiency. Compared with some other electrode materials, the treatment efficiencies of this electrode type are not high. Based on this research, Chu Thi Thu Hien (2014)\cite{27} has successfully developed the Ti/SnO\textsubscript{2}-SbO\textsubscript{2}/PbO\textsubscript{2} inert anodic electrode by combining the decomposition of SnCl\textsubscript{4}, SbCl\textsubscript{3} solutions at 480°C temperature and was covered by PbO\textsubscript{2} at 120 minutes. Electrode life is approximately 290 hours. The simulated phenol of 500 mg/L concentrations showed the highest efficiency of electrochemical treatment as conditions: pH = 8.50, COD = 1.50 g/L Na\textsubscript{2}SO\textsubscript{4} concentrations, 1% NaCl solution. The phenol removal efficiency was 99.60% at 360 minutes. The results of electrode application have been manufactured to treat real textile wastewater that COD removal efficiency was 95.17% after 360 minutes with
constant current intensity of 50 mA/cm\(^2\). The Cl\(^-\) ions not only a reducing passivity role and activating the electrode but also contributes to the formation of ClO\(^-\), ClO\(_3\^-\) and HO\(^*\) radicals are phenolic oxidation agents and intermediate products, according to the equations below:

\[
\text{ClO}^-(ad) + \text{H}_2\text{O}(ad) \rightarrow \text{Cl}^- + 2\text{HO}^* \quad (17)
\]

\[
\text{ClO}_3^-(ad) + 2\text{H}_2\text{O}(ad) \rightarrow \text{Cl}^- + 4\text{HO}^* \quad (18)
\]

Similarly, research by Nguyen Ngoc Kien (2013) [28] produced PbO\(_2\) anode on carbon graphite subtract combined with 304 stainless steel cathode. The optimum conditions for the synthesis of anode in this study are: 20 ml/l of HNO\(_3\) solution, 0.6 M of Pb(NO\(_3\))\(_2\) concentration, 0.4 M Cu(NO\(_3\))\(_2\), 1 g/l of gelatin, 25\(^\circ\)C - 30\(^\circ\)C temperature, the current density for electrodialysis is 40 mA/cm\(^2\). The study investigated the phenol removal ability as conditions: pH = 8.0, 7.5 g/l of NaCl solution, 1000 mg/l of input phenol concentration. The results showed that at the current density of \(i = 75\) mA/cm\(^2\) which reached 92.7% removal efficiency at 210 minutes. The results of the X-ray diffraction pattern of the PbO\(_2\) electrode show in figure 2 illustrated the line represents the unaltered tetrahedral structure, which indicates that the strength of the electrode is very high.

![Figure 2. X-ray diffraction pattern of PbO\(_2\) sample (A. Before reaction B. After reaction [28]).](image-url)
The issue of ammonium removal of wastewater is great interested today. In this field, Pham Cong Minh et al. (2015) [29] has conducted to research on removing of ammonium (NH\textsubscript{3}-N) for synthetic wastewater by indirect electrochemical oxidation using Platinum anode and stainless steel for cathode. Synthetic wastewater was prepared in a laboratory by dissolving 99% (NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4} solution. The surface areas of each electrode are 100 cm\textsuperscript{2}. The model uses circulating pump with 5 liters per minute and stirred at 150 rpm. The study conducted to the ammonium removal in direct and indirect electrolyte conditions. The direct electrolysis (input wastewater without chloride addition) of ammonium removal efficiency is very low, approximately 8% after 80 minutes. Meanwhile indirect electrolysis (chloride added to wastewater) concentrated that chloride productions are generated at the anode while presence of chloride ions and chlorination of the critical point between chlorine and ammonium to converting to nitrate and N\textsubscript{2}. Ammonium removal efficiency was above 99% with indirect electrolysis conditions as: 25 mg/l of input ammonium concentration, 0.03% of NaCl solution, pH\textsubscript{±} = 7.5, 20 mA/cm\textsuperscript{2} of current density with removal time at 70 minutes.

Lam Hoa Hung et al. (2017) [30] investigated the p-Nitrophenol ability of degradation by electro-Fenton process using bar graphite electrode. Pt/Ti anode type of grid was used and a graphite rod with 9.5 mm diameter of cathode. This study was carried out considering of reducing Fe\textsuperscript{3+} to Fe\textsuperscript{2+} ions and deoxygenation generated to H\textsubscript{2}O\textsubscript{2} under electrolysis conditions are separated by salt bridges of 1.0M NaCl solution. The result of the reducing process Fe\textsuperscript{3+} to Fe\textsuperscript{2+} on the graphite electrode occurs relatively easily, the reaction was occurred when applied to 1.0 V voltage and saturated at 3.0 V. However, Fe\textsuperscript{3+} ions removal efficiency decreased over time as a result of under-voltage condition. When the voltage is constant, the current density supplied to electrodes decrease over time, so the equilibrium voltage at the electrodes surface fall. However, at 75 minutes electrolysis, the Fe\textsuperscript{2+} ion concentration was accumulated at 2.74x10\textsuperscript{-5} M which are enough to join the electro-Fenton reaction. The H\textsubscript{2}O\textsubscript{2} is generated more difficult because the activity of graphite electrodes is relatively low. The voltage applied to generate the Fe\textsuperscript{2+} ion concentration has removed form 1280 mg/l to 160 mg/l at 35 minutes. This study showed that when pH value increases to causing H\textsuperscript{+} ion concentration decrease, the H\textsubscript{2}O\textsubscript{2} producing reaction is good but Fe\textsuperscript{2+} ion concentration reducing. Therefore, the OH\textsuperscript{*} radical is generated that caused to decrease the organic decomposition efficiency. The research has compared the performance between three methods: electrochemical, electro-Fenton and Fenton’s reagent technologies on paper industry wastewater. The results showed that high-to-low removal efficiency are electro-Fenton, Fenton’s reagent and electrochemical respectively. Electrochemical method was the least performance because the only oxidation agent is electrolysis process but the graphite electrode has low decomposition efficiency. Fenton's reagent method has the disadvantage that generated Fe\textsuperscript{2+} is low because the H\textsubscript{2}O\textsubscript{2} concentration decreases over time. The presence of strong oxidizing agents are OH\textsuperscript{-} and HO\textsuperscript{2-} radicals so removal efficiency of Fenton electrochemical is higher than other two methods. Nguyen Duc Dat Duc et al. (2016) [31] conducted the same study to use graphite anode to remove color of real textile wastewater by electro-Fenton process. The results showed that optimal conditions of the process were as: pH = 3.11, 1.82 mM Fe\textsuperscript{2+} (this concentration is higher than the research of Dinh Thi Mai Thanh, 2009), 19 V of current density. The color removal efficiency reached over 96% at 30 minutes and color concentrations reduce from 1500 Pt-Co to 46 Pt-Co. This research was contradictory to the study by Lam Hoa Hung (2017) when the voltages above 7 V do not increase the p-nitrophenol treatment efficiency.

Azodyes from textile wastewater most commonly have complex carbonaceous structures that are difficult to decompose using traditional methods such as physiological and biological methods. Nguyen Thi Le Hien et al. (2009) [33] carried out removing Red metal by electro-Fenton method with working anode is graphite covered Cu\textsubscript{1.5}Mn\textsubscript{1.5}O\textsubscript{4} and without covered Cu\textsubscript{1.5}Mn\textsubscript{1.5}O\textsubscript{4} (5 cm\textsuperscript{2} electrode area). Cathode is platinum grip and Calomel saturated KCl is reference electrode. In this study, the reaction was concentrated in the cathode with deoxidation process produces H\textsubscript{2}O\textsubscript{2}. Fenton’s reagent process is formed from H\textsubscript{2}O\textsubscript{2} combined to Fe\textsuperscript{2+} ion which create to OH\textsuperscript{*} radicals to destroy pollutants. Under optimal conditions as: -0.6 V/SCE cathodic voltage, oxygen flow as 1 L/m, pH =3, 1 mM Fe\textsubscript{2}SO\textsubscript{4}, 0.1 M Na\textsubscript{2}SO\textsubscript{4} of electrolyte solution, 0.185 mM of Red - Methyl Effluent COD concentration is 50 mg/l after at 24 hours electrolysis when the input COD concentration of 1300 mg/l. Investigation effect of H\textsubscript{2}O\textsubscript{2} generated by cathode with aeration and no aeration. The results showed that reducing of dissolved oxygen available in water without aeration is not high performance. However, performance unchanged if aeration is greater than the saturation level of reaction at 1 L/m flow rate. A mount of Fe\textsuperscript{2+} added also has a great influence on processing. The amount of
Fe\(^{2+}\) less than 1 mM concentration is not enough to total creating H\(_2\)O\(_2\) generated, but if Fe\(^{3+}\) more 1 mM that decreases efficiency. A mount of Fe\(^{3+}\) ion concentration redundancy is oxidized to create Fe\(^{3+}\) ion at anodic electrode, the pair of Fe\(^{3+}/\text{Fe}^{2+}\) oxidation-reduction reaction redundancy will occur consecutive reaction on anodic and cathodic electrodes that decrease electrical efficiency. The results illustrated to \(\text{Cu}_2\text{Mn}_4\text{O}_9\) C/Ppy/\(\text{Cu}_2\text{Mn}_4\text{O}_9\) anode has the highest processing efficiency. In the same study, Nguyen Thi Le Hien et al. (2010) [34] repeated the study on Orange-Methyl by C/Ppy/\(\text{Cu}_2\text{Mn}_4\text{O}_9\) anode. The results showed that, at 1 mA/cm\(^2\) of current density, pH = 3, aeration flow rate of 0.5 L/m\(^2\), 1 mM Fe\(^{2+}\). Orange-Methyl has lost its color completely after 15 hours treatment. The UV-Vis spectroscopy at the treatment sites showed that peak characterization of the Orange-Methyl was lost. At 229 nm wavelength there is a new peak which indicates the presence of OH\(^{+}\) radicals involved in the decomposition of the azo color group.

3.3. Electrodialysis

Electrodialysis is a relatively high efficiency of desalination technology in parallel with Reverse Osmosis technology (RO). However, there are some disadvantages compared to RO technology, so it is not widely to use. Nguyen Hoai Chau et al. (2005) [35] have applied this technology to remove salinity, hardness and suspended solid in ground water with the experimental model scale of 100 l/h capacity. The research has compared the performance of three different membranes: Vietnamese membrane made from cellulose acetate; Russian membranes and US membranes. The cylindrical membrane is 200 mm of diameter and 12.56 cm\(^2\) area of each. The working electrode is made of Titanium covered an anti-corrosive layer. The results showed that three membranes had similar performance that reached as 83%, 66%, 65% performance for hardness, suspended solid, desalination removal efficiencies respectively. This study demonstrates that electrodialysis technology has potential in field of desalination with low power consumption, not over 1 kWh per 1 kg salt to be removed. However, this method still has some limitations that affect the performance by osmosis and electro-osmosis process, the fouling membranes by presence of organic and inorganic substances.

In addition to desalination, the electrodialysis is also investigated for the treatment of nitrogen compounds in groundwater. Model of 30–60 l/h capacity included titanium electrodes covered by noble metal oxide layer for anti-corrosive, 53 VDC of voltage, a range of 0.75–1.5A current density. The results illustrated over 99% ammonium removal efficiency when input ammonium concentration was 10.4 mg/l. The nitrite and nitrate removal efficiency were 80% and 60%, respectively. The power consumption for this process ranged from 1.2 to 2.5 kWh/m\(^2\) (Nguyen Thu Tra, 2005 [36]).

4. Conclusion

In conclusion, the electrochemical methods in Vietnam to treat water and wastewater have been investigated and applied but quantities and quality also exceeded. Those technologies have also been studied to combine with other methods that achieved the desired treatment performance. However, the researching trends of the authors mainly focused on the laboratory model scales and the simulated raw water. Therefore, overall efficiency of specific indicators have not been evaluated and applied of real wastewaters. However, the results also provided the technical and economic feasibility of this approach. Previous studies focused on electrocoagulation, although this approach is cheap costs but it was generated secondary sludge which is harmful to environment. So research on electrochemical oxidation should be applied in the future, especially the advanced electrodes that the world is using today. In the future, electrochemical methods should be used to study the different types of wastewater at the pilot model scales, especially difficult handling wastewater and toxics that biological methods cannot handle effectively. In addition, it is necessary to find the optimal operating parameters to saving treatment costs as well as to reusing resources for sustainable development. On the other hand, the electrodialysis technology for desalination is applied to treat salty water in the Mekong Delta also interested in situation affected by climate change. That is also mainly research trends to environmental technology faculty of the Vietnamese – German University at the present.

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