Selective Leaching of Vanadium from Boiler Oiled Ash Residue Using Sodium Carbonate-Bicarbonate Binary Solution

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Abstract
The present work aims to optimize an efficient, economic and environmental hydrothermal method to leach vanadium from boiler ash residue using sodium carbonate and bicarbonate mixture solution. For this purpose, many factors include liquid/solid phase ratio, leaching temperature, leaching time, oxidant addition and its percent and ash fineness are studied and optimized. From the obtained results, it is concluded that, maximum vanadium leaching (84.3%) is achieved from grinded ash (74 µm) using carbonate-bicarbonate binary solution at liquid/ash ratio of 20/1 in the presence of KMnO4 as oxidant. The obtained leach liquor is then subjected to pH readjusted at 5-6. This is followed by ammonium chloride (NH4Cl) addition under heating (40°C) and stirring to selective precipitation the leached vanadium species as ammonium meta-metavanadate (NH4VO3). The precipitate is dried at 100°C and then roasted at 750°C to obtain V2O5 oxide.

Keywords: Vanadium; Hydrothermal leaching; Boiler ash; Carbonate; Bi-carbonate

Introduction
Vanadium as metal and oxide are very important in many applications especially in steel alloys, aerospace applications, ceramics and glass besides as catalysts in chemical industries [1]. In nuclear applications, the short half-life, good neutron capture, low induced activation, high strength at elevated temperatures, superior ductility at low temperatures besides resistance to irradiation damage makes vanadium alloys (V-Ti-Cr) an excellent material for the inner structure of the fusion reactors in addition in purification of uranium [2]. Vanadium naturally exist in carnotite [K2(UO2)2(VO4)2·3H2O], vanadinite [Pb3(VO4)2Cl] and patronite [VS4] ores in addition vanadium-bearing magnetite ores, but vanadium recovery from these ores is not efficient for the previous using. So, the oil ashes as additional vanadium source is interested [3-5].
The oil ashes referred the residues produced from combustion crude oils as fuel in power plants (several million tons are generated annually) but small amounts of these ashes are reused [6]. Chemically the ash is composed of two main parts; the first one is the organic part includes sulfates, nitrogen-containing compounds besides unburned carbon while the other is the inorganic fraction that contains heavy metals particularly vanadium, nickel, silicon and iron in addition of aluminum, magnesium, zinc and sodium species [7]. Vanadium content in oil ashes ranges from 2.5% to 30% according to the oil origin and the ash consumption conditions. The recovery of vanadium from oil ashes has received attention since 1960 not only an economic concern (metals recovery) but also from environmental terms to humans, waters and soils [8-10].

For these purposes many techniques are applied for vanadium leaching from ashes, but the common methods are hydrometallurgy (acidic, alkaline) and pyro-hydrometallurgy (roasting) processes [11-19]. The most applied among these methods are acid leaching, alkali leaching and acid/alkali or water leaching after roasting [18]. Generally, the hydrometallurgical processes have some benefits than pyro-hydrometallurgy include; low cost requirements and air pollutions releasing [20,21]. For the hydrometallurgical processes; despite the acidic methods have many advantages than alkaline namely; higher metals recovery, simple and low cost but these methods consume high acids, require acids resistance equipment’s besides producing high wastes. In contrast with acidic methods, the alkaline solutions have many advantages which include; high selective, direct precipitation the leached metals, direct wastes discharge [22,23]. Based on the leaching medium nature and solution pH, vanadium species present as the Pourbaix diagram demonstrated FIG. 1 [24-26]. The main target of this work is optimized an environment and effective procedure to hydrothermal leaching of vanadium content from the boiler ash residue using binary carbonate-bicarbonate solution, this process is followed by vanadium precipitation from the obtained liquor solution.

![Pourbaix diagram of vanadium in the aqueous medium.](image-url)
Material and Methods

Ash sample specifications

Boiler ash is a generic term produced after burning of petroleum products in generating thermal power stations. Bottom ash refers to the ash that falls through the airflow to the bottom of the boiler and represents about 5-15 wt% from the produce ash. Consequently, bottom ash is considered as hazardous residue but less than fly ash [27]. The bottom ash properties are a friable, amorphous, greyish color after exposure to sunlight and its specific gravity is 2.3 with high carbon content.

Analytical procedures

For boiler ash quantitative analysis, the studied sample (0.25 g) is digested over hot plate in Teflon backer with mixture of concentrated acids [HNO₃ (10 ml)+HClO₄ (5 ml)+HF (10 ml)] till dryness, the residue is re-dissolved by diluted HCl (10 ml). Vanadium (V) is determined spectrophotometrically using 2-(2-pyridylazo)-5-diethylaminophenol at 595 nm [28]. Chromium as Cr (VI) was determined spectrophotometrically using 1,5-diphenyl-carbazide dye at 540 nm [29]. Titanium analysis is performed using colorimetric using Tiron (disodium-1,2-dihydroxybenzene 3,5-disulfonate) as complexing agent at 430 nm. Iron as Fe³⁺, Ca²⁺ and Mg²⁺ species are determined titrimetrically against EDTA solution using sulfosalicylic acid, Murexide and Eriochrome Black T indicators respectively. For silica and alumina oxides determination in the studied ash, a weighted sample (0.1 g) is melted with NaOH (2 g) in nickel crucible, the molten cake is re-dissolved using dilute HCl (10%) then completed with distilled water. Molybdate and Alizarine reagents are used for silica and alumina determination spectrophotometrically at 630 and 475 nm respectively [30]. The colorimetric measurements are carried out using Unicam UV2-100 spectrophotometer. The vanadium measurements beside trace ions during leaching experiments are emphasized using Unicam 969 atomic absorption spectrophotometer. The total loss of ignition (L.O.I) in the studied sample includes carbon content and the oily matrix besides water content is evaluated by thermal decomposition in muffle furnace at 750°C-850°C.

General leaching procedures

The finely grinded ash sample (74 µm) is mixed with 0.5 M Na₂CO₃-NaHCO₃ (mol/mol) mixture solution in 150 ml a round Pyrex flask equipped with a reflux condenser (Na₂CO₃-NaHCO₃ solution/ash (ml/g) ratio was 20/1) at 90°C for 2 h at 400 rpm stirring speed. The obtained solution after cooling is filtrated then subjected to metals analysis. The obtained data indicates that, iron; titanium, manganese and nickel metal ions are not leached by using Na₂CO₃-NaHCO₃ solution which improves the vanadium leaching with high purity in the precipitation step. Similar results are obtained during leaching of vanadium from ash by using NaOH to avoid iron, nickel and titanium species leaching, while small silica amount is dissolved [19]. In the present work the studied ash sample contain low silica content (1.91%) so Na₂CO₃-NaHCO₃ solution is a suitable choice for vanadium leaching from the ash sample. The other contents that consume carbonate-bicarbonate as sulphide, sulphates and organic matters are indicated in the sample. During the leaching process, the magnetic stirrer bar immersed in the solution is captured the impurities having magnetic properties especially iron which insoluble by using the alkali leaching solution.
FIG. 2. The XRD and SEM analysis of the boiler oil ash sample.

Results and Discussion
The boiler ash sample has been characterized by scanning electron microscopy (SEM) and XRD analysis FIG. 2 and the wet chemical analysis. Trace elements in the ash are determined using ICP method. The fine studied sample after grinding and sieving is subjected to alkali hydrothermal digestion process using sodium carbonate and bi-carbonate binary solution in presence of potassium permanganate as oxidant. Factors include carbonate, bi-carbonate and oxidant concentrations, sample fineness, solution/solid phase ratio; digestion time and temperature are studied and optimized.

Effect of alkali solution/ash ratio
The effect of 0.5 M Na₂CO₃-NaHCO₃/ash ratio (ml/g) on vanadium leaching from the ash sample is studied from 5/1 to 30/1 while other conditions are; 100°C reaction temperature and 150 min digestion time in presence of 0.01 M KMnO₄ as oxidant. The obtained results maintain that, the leaching efficiency of vanadium increases from 32.1 to 82.4% as the solution/ash ratio increases from 5/1 to 20/1. Increasing the solution/ash ratio to 25/1 lead to a slightly vanadium leaching increases (83.2%) as presented in FIG. 3. Increasing the liquid/ash ratio to 30/1 has a vanadium leaching decreasing (79.3%), which is believed to be due to iron sodium silicate [NaFe(SiO₃)₂] formation which makes vanadium leaching becomes worse. In this context; the study for alkali leaching of vanadium with high selectivity is carried [31]. The obtained results show selective vanadium dissolution than many species specially nickel and iron. Some studies indicate that, vanadium often present in the sample in a reduced state and carbonate form a soluble pentavalent vanadate (VO₅⁻) complexes [3]. A study for the speciation of vanadium using alkaline solutions is applied, the obtained data indicates that, vanadium species after leaching presented as VO₅⁻, VO₃(H₂O)²⁻, VO₃(H₂O)³⁻, V₂O₇⁴⁻ and VO₄³⁻ [26]. There is an additional advantage of using carbonate-bicarbonate solution is the viability of CO₃²⁻ anion to be regenerated through the reaction as equation 1 show. There is another advantage using carbonate solution than acids for vanadium leaching from samples containing, iron nickel and calcium species, the carbonate is selective for vanadium than iron and calcium species despite sodium carbonate converts the insoluble calcium-vanadate in the sample to calcium carbonate and releases water soluble sodium vanadate species as in equation 4 [32]. The proposed reactions present in equations 1-3. For maximum vanadium dissolution, the generated NaOH species in equations 1 and 2 must be removed. The produced NaOH is neutralized by NaHCO₃ addition, to prevent re-precipitation of the dissolved vanadium as equation 5 indicated.
Generally, the alkaline leaching of vanadium from ores should be applied under oxidation conditions for vanadium dissolution completely. The overall vanadium loss during the leaching process is attributed to the insufficient oxidizing of V(IV) to V(V) [25]. Vanadium oxidation is conducted using chosen oxidant as potassium permanganate, potassium chlorate, hydrogen peroxide etc. Despite oxidants efficiency in alkaline medium is less than in the acidic ones but an oxidant should be used for enhances the vanadium leaching. Potassium permanganate is a powerful oxidizing agent in alkaline or acidic solution, so in the present work we use KMnO₄ as oxidant. Several experiments are carried out at S/L of 1/20 at 90°C-95°C for 3 h digestion time to define the minimum amount of oxidant compatible with maximum vanadium leaching efficiency. Potassium permanganate amount addition is varied from 0.002 to 0.02 M. From the obtained results in FIG. 4. It is obvious that, KMnO₄ as oxidant played a critical effect in vanadium dissolution by releasing active oxygen for V(IV)/V(V) oxidation controlled as equations 6-8 indicated. The leaching efficiency of vanadium increases from 53.2 to 82.7% as KMnO₄ concentration increases from 0 to 0.005 M. As KMnO₄ concentration increases to 0.006 M, the extraction efficiency of vanadium increases to 83.4%. The same behavior is found during leaching uranium from carnotite ore in presence of KMnO₄ as oxidant using alkaline solution [33]. Complete uranium leaching is achieved in presence KMnO₄ which emphasizes the role of KMnO₄ as oxidant in the basic mediums. In this context, a study is carried out using KClO₃ as oxidant and 2% (w/w) KClO₃ is the good choice for completely vanadium leaching from oil fly ash using sodium hydroxide solution [34].

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\begin{align*}
2\text{KMnO}_4 & \rightarrow \text{K}_2\text{MnO}_4+\text{MnO}_2+\text{O}_2 \\
2\text{K}_2\text{MnO}_3 & \rightarrow 2\text{K}_2\text{MnO}_3+\text{O}_2 \\
\text{MnO}_4^-+2\text{H}_2\text{O}+3e^- & \rightarrow \text{MnO}_2+4\text{OH}^- 
\end{align*}
\]
FIG. 4. Effect of KMnO₄ oxidant on vanadium leaching from ash using Na₂CO₃-NaHCO₃ solution.

Effect of digestion temperature

The effect of digestion temperature on vanadium leaching from the studied ash sample using Na₂CO₃-NaHCO₃ binary solution is studied using leaching solution/solid ratio of 20/1 at 2.5 h digestion time in presence of 0.006 M KMnO₄ as oxidant at various temperatures ranges from 30°C to 100°C. The results in FIG. 5 show that, raising temperature in the studied range increased the recovery of vanadium which indicates that, the alkaline leaching of vanadium using Na₂CO₃-NaHCO₃ solution is an endothermic reaction, increasing the temperature lead to improve the vanadium leaching efficiency. The leaching of vanadium increases from 30.7% to 82.3% by raising the reaction temperature from 30°C to 80°C. Rising the temperature over 80°C lead to a slight vanadium leaching improvement (83.2% at 100°C). So, the optimum leaching temperature is chosen at 80°C.

FIG. 5. Effect of temperature on vanadium leaching from ash using Na₂CO₃-NaHCO₃ solution.

Effect of leaching time

To evaluate the effect of digestion time on vanadium leaching from the ash sample using Na₂CO₃-NaHCO₃ solution, series of leaching experiments is carried out at different times ranged from 5 to 150 min. The other leaching conditions are fixed at; 20/1 liquid/ash ratio in presence of 0.006 M KMnO₄ as oxidant at 100°C reaction temperature. From the results in FIG. 6.
is concluded that, as the contact time increases from 5 to 105 min the vanadium leaching efficiency increases from 8.81 to 84.23% respectively. Vanadium leaching percent steadies as the leaching time reaches 120 min then leaching efficiency has been decreased as the time increases from 120 to 135 min (82.1%). Therefore, 120 min leaching time would be adequate and could be considered as optimum time for maximum vanadium leaching.

![Graph showing vanadium leaching efficiency](image)

**FIG. 6. Effect of digestion time on vanadium leaching from ash using Na₂CO₃-NaHCO₃ solution.**

**Effect of particle grain size**

Generally; the more finely divided the solid is the faster the reactions proceed (leaching efficiency). To study the effect of ash grain size on vanadium dissolution from the treated ash using carbonate and bicarbonate mixture solution, a leaching experiment using ash grain size ranging from 105-44 µm (140 to 325 mesh size) are performed for a reaction time of 2 h at 90°C at a liquid/solid ratio of 20/1 in presence of 0.006 M KMnO₄ as oxidant. The obtained results in FIG. 7 reveals that, the leaching efficiency of vanadium is increased by decreasing the sample grain size due to increasing the surface area of grains exposed to leaching solution which facilitates the alkali attack. The recovery efficiency of vanadium is increased from 61.2% to 83.1% as the grain fineness decreased from 105 µm to 74 µm. More grinding than 44 µm resulted in only about 0.85% increases in the vanadium leaching efficiency. Therefore, 74 µm would be considered the optimum grain size.

![Graph showing effect of grain fineness](image)

**FIG. 7. Effect of grain fineness on vanadium leaching from ash using Na₂CO₃-NaHCO₃ solution.**
Vanadium precipitation procedures

The leach liquor solution after alkaline leaching using sodium carbonate and bicarbonate mixture solution under the optimized conditions is applied and the resulted solution mainly contains vanadium, silicon, sodium, magnesium and aluminum with traces of chromium species with leaching percentages.

For vanadium precipitation, the pH of the obtained leach liquor is adjusted to pH 7 using H$_2$SO$_4$ solution FIG. 8A to precipitate the leached Al species (V remained in solution). The solution pH is readjusted at 5-6, then 1 M ammonium chloride (NH$_4$Cl) is added to the solution under stirring. The reaction (under agitation at 50°C) is set to about 24 h. Vanadium is precipitated as ammonium meta-metavanadate (NH$_4$VO$_3$) as provided in FIG. 8B, the precipitation percent reaches 97% (calculated by measuring the un-precipitated vanadium amount in the leach liquor solution). The obtained red colored cake as shown in FIG. 8C after filtration and washing with distilled water is dried at 100°C for 24 h and roasted at 750°C to obtain a black colored vanadium oxide FIG. 8D. The remaining filtrates besides the obtained oxides are analyzed for metals content.

FIG. 8. Successive procedures for vanadium precipitation using ammonium chloride.

Conclusion

In the present work a modified method is investigated for leaching of vanadium from the boiler oil ash using a carbonate-bicarbonate binary mixture solution. Maximum vanadium leaching (84.3%) is obtained using leaching solution at solid/liquid phase ratio at 1/20 (g/ml) at 80°C leaching temperature for 2 h digestion time in presence of KMnO$_4$ as oxidant as optimum conditions. The obtained liquor solution is then subjected to vanadium recovery by using ammonium chloride (NH$_4$Cl) at slightly acidic solution (pH; 6) to precipitate vanadium as ammonium meta-metavanadate (NH$_4$VO$_3$). The obtained precipitate is then dried at 100°C and ignited at 750°C to obtain vanadium pentoxide (V$_2$O$_5$) residue.
REFERENCES


