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Redox Properties of Heteropolyacid Catalysts Probed by Electrochemical Analysis, UV-visible Spectroscopy, and Scanning Tunneling Microscopy

전기화학적 분석, 자외선-가시광선 분광법 및 주사 터널링 현미경을 이용한 헤테로폴리산 촉매의 산화환원 특성연구

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서울대학교 대학원
화학생물공학부
최 정호
Abstract

Redox Properties of Heteropolyacid Catalysts Probed by Electrochemical Analysis, UV-visible Spectroscopy, and Scanning Tunneling Microscopy

Jung Ho Choi
School of Chemical and Biological Engineering
The Graduate School
Seoul National University

Heteropolyacids (HPAs) are polymeric metal-oxide clusters that exhibit the diverse range of structures and compositions. Because of their robust structures and unique redox properties, HPAs have been widely employed as catalysts for several redox reactions. Catalytic oxidations over HPAs have been extensively studied over the past few decades and mixed-addenda HPAs have attracted recent attention due to the variety in stoichiometric combinations and outstanding redox natures. Physicochemical properties and catalytic activity of mixed-addenda HPAs can be easily tuned at molecular level by changing the constituent elements including counter-cation, central heteroatom, or framework addenda atom. Because a number of elements including metals, semimetals, and even non-metals could be incorporated into the HPA frameworks, a number of mixed-addenda HPAs with different structures can be
designed as a candidate for promising oxidation catalyst.

In this work, several series of transition metal-substituted HPAs were designed and synthesized. They were investigated by several experimental techniques including electrochemical analysis, UV-visible spectroscopy, and scanning tunneling microscopy (STM) in order to elucidate the effect of transition metal-substitution on redox properties and catalytic activities in oxidation catalysis. Furthermore, reliabilities of absorption edge energy and negative differential resistance (NDR) peak voltage as alternative parameters for the reducibility were also examined.

First, molybdenum-substituted $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs were prepared by “etherate method” to elucidate the effect of molybdenum-substitution on the redox properties and catalytic activity of Wells-Dawson-type tungstophosphates. Electrochemical measurements were conducted to elucidate the redox properties of HPAs. Several tungsten-based redox transitions were observed in the cyclic voltammogram of $\text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62}$. However, $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ exhibited molybdenum-based redox transitions. Interestingly, molybdenum-substituted Wells-Dawson HPAs showed an additional molybdenum-centered redox transition at more positive potential. First electron reduction potentials increased with increasing molybdenum-substitution. UV-visible spectroscopy measurements were conducted to probe the electronic structure of bulk $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs. Absorption edge energy determined from the linear fit of $[F(R_\infty) \cdot h\nu]^{1/2}$ (Tauc plot) decreased with increasing molybdenum content. Scanning tunneling microscopy (STM) measurements were performed for the
further investigation about the local surface electronic structure of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs. In STM measurements, two-dimensional self-assembled HPA arrays were observed. Tunneling spectra taken at bright corrugations showed a distinctive current-voltage responses, referred to as negative differential resistance (NDR) phenomenon. NDR peak voltage appeared at less negative voltage with increasing the molybdenum content. Gas-phase oxidative dehydrogenation of ethanol to acetaldehyde was carried out as a model reaction to probe the oxidation catalysis of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs. Yield for acetaldehyde (oxidation product) increased with increasing molybdenum content. Among the tested catalysts, $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ with the highest reduction potential showed the best catalytic performance.

In order to explore the effect of group 5 metal (V and Nb)-substitution on redox properties and catalytic activity of Wells-Dawson-type tungstophosphates. $\alpha_2$-$\text{K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ and $\alpha_2$-$\text{K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$ Wells-Dawson HPAs were synthesized via direct incorporation of transition metal into the mono-lacunary species to yield the selectively-substituted structures. $\alpha$-$\text{K}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ and $\alpha_2$-$\text{K}_6\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62}$ were also prepared for the comparison. In the electrochemical analysis, molybdenum- and vanadium-substituted Wells-Dawson tungstophosphates showed additional molybdenum- and vanadium-based redox transitions, respectively, at more positive potential. However, niobium-substituted Wells-Dawson tungstophosphates showed significantly shifted redox transitions. First electron reduction potential increased in the order of $\alpha_2$-$\text{K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$ < $\alpha$-$\text{K}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ < $\alpha_2$-$\text{K}_6\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62}$ < $\alpha_2$-$\text{K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$. Absorption edge energy determined by UV-visible
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Heteropolytungstates with different central atom, $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs were prepared to elucidate the effect of central atom on the redox properties and catalytic activity. All $\alpha$-H$_n$XW$_{12}$O$_{40}$ HPAs exhibited well-defined reversible and stepwise tungsten-centered redox transitions during the electrochemical measurements. First electron reduction potential increased in the order of $\alpha$-H$_6$CoW$_{12}$O$_{40} < \alpha$-H$_5$BW$_{12}$O$_{40} < \alpha$-H$_4$SiW$_{12}$O$_{40} < \alpha$-H$_3$PW$_{12}$O$_{40}$. Absorption edge energy determined by UV-visible spectroscopy decreased in the order of $\alpha$-H$_6$CoW$_{12}$O$_{40} > \alpha$-H$_5$BW$_{12}$O$_{40} > \alpha$-H$_4$SiW$_{12}$O$_{40} > \alpha$-H$_3$PW$_{12}$O$_{40}$. The trend of NDR peak voltage was also well consistent with that of absorption edge energy. Among the tested, PO$_4^{3-}$ anion with smaller negative charge and larger size was the most effective to enhance the reducibility.

Another series of heteropolytungstates containing AsO$_4^{3-}$ as a central unit were also examined to elucidate the effect of transition metal-substitution on the redox properties and catalytic activities of Wells-Dawson-type tungstoarsenates. A series of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) Wells-Dawson HPAs were prepared via direct incorporation of transition metal into the mono-
di-, and tri-lacunary species. In electrochemical analysis, $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ exhibited four tungsten-centered redox transitions. However, molybdenum-substituted $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=1-3$) exhibited an additional molybdenum-centered redox transition at more positive potential. First electron reduction potential increased with increasing molybdenum content. Absorption edge energy determined by UV-visible spectroscopy decreased with increasing molybdenum content. It is interesting to note that NDR peak voltage appeared at less negative voltage with increasing molybdenum content. These results are nearly same with the results in the series of H$_6$P$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0, 3, 9, 15, 18$). Gas-phase oxidative dehydrogenation of benzyl alcohol was carried out as a model reaction to track the oxidation catalysis. Yield for benzaldehyde (oxidation product) increased with increasing molybdenum content.

Furthermore, group 5 metal-substituted $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs were synthesized via direct incorporation of transition metal into the mono-lacunary species to yield the selectively-substituted structures. $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ and $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ were also prepared for the comparison. In the electrochemical analysis, molybdenum- and vanadium-substituted Wells-Dawson tungstoarsenates showed additional molybdenum- and vanadium-based redox transitions, respectively, at more positive potential. However, niobium-substituted Wells-Dawson tungstoarsenates showed significantly shifted redox transitions. First electron reduction potential increased in the order of $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ $<$ $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ $<$ $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ $<$ $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$. Absorption edge energy determined by UV-visible spectroscopy decreased in the order of $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ $>$ $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ $>$ $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ $>$ $\alpha_2$-
K\textsubscript{7}As\textsubscript{2}W\textsubscript{17}V\textsubscript{1}O\textsubscript{62}. Gas-phase oxidative dehydrogenation of benzylamine was carried out as a model reaction to track the oxidation catalysis. Yield for dibenzylimine (oxidation product) increased in the order of α\textsubscript{2}-K\textsubscript{7}As\textsubscript{2}W\textsubscript{17}Nb\textsubscript{1}O\textsubscript{62} < α-K\textsubscript{6}As\textsubscript{2}W\textsubscript{18}O\textsubscript{62} < α\textsubscript{2}-K\textsubscript{6}As\textsubscript{2}W\textsubscript{17}Mo\textsubscript{1}O\textsubscript{62} < α\textsubscript{2}-K\textsubscript{7}As\textsubscript{2}W\textsubscript{17}V\textsubscript{1}O\textsubscript{62}.

In summary, several series of transition metal-substituted HPA catalysts with different addenda atoms, central atoms, contents, and structures were prepared, characterized, and applied to the model reactions in order to elucidate the redox properties and catalytic activities in oxidation catalysis. Reliabilities of absorption edge energy and NDR peak voltage as alternative parameters for the redox properties were also examined. It was found that redox properties were easily tunable by changing the constituent elements. In addition, it could be concluded that redox properties of HPAs, which is closely related to the electronic structure, play an important role to determine the catalytic activities in the oxidation catalysis and absorption edge energy and NDR peak voltage can be utilized as alternative parameters to estimate the reducibility of HPAs.

**Keywords:** Heteropolyacids, Transition metal, Redox property, Oxidation catalysis, Electronic structure

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Chapter 1. Introduction

1.1. Heteropolyacids (HPAs)

Heteropolyacids (HPAs) are polymeric oxoanion clusters with diverse range of molecular structures and compositions that exhibit the unique acid-base and redox nature. As shown in Eq. (1), acidic elements such as W, Mo, V, and Nb generally forms the oxoanions in the aqueous solution. These oxoanions polymerize at certain pH, temperature, and concentration. Polymerization proceeds until the strong inward polarization of exterior oxygen atoms terminates further polymerizations, resulting in small discrete metal-oxygen clusters. Polyanions comprising only a kind of oxoanion are called isopolyanion. Polyanions comprising more than two kinds of oxoanions are called heteropolyanion.

\[
12\text{WO}_4^{2-} + \text{HPO}_4^{2-} + 23\text{H}^+ \rightarrow \text{PW}_{12}\text{O}_{40}^{3-} + 12\text{H}_2\text{O} \quad (1)
\]

Schematic representations of primary, secondary, and tertiary structures of Keggin-type HPAs are shown in Figure 1.1. Keggin-type heteropolyanion, which has a general formula of \( \text{XM}_{12}\text{O}_{40}^{n-} \), comprises central atom (X), addenda atom (M), and framework oxygen. A central tetrahedral \( \text{XO}_4 \) unit is surrounded by four octahedral \( \text{M}_3\text{O}_{13} \) units, resulting in soccer ball-shaped Keggin-type heteropolyanion.
Heteropolyanions are generally coordinated with charge-compensating counter-cations (e.g. protonated water dimer, H$_5$O$_2^+$) and other polar molecules to form the secondary structures. Heteropolyanions are linked each other by hydrogen bonding at the terminal or bridged oxygen. Other elements such as Na$^+$, NH$_4^+$, or Cs$^+$ are also available as a counter-cation. Tertiary structure is the structure of solid heteropoly compounds as assemble. Counter-cation plays an important role to determine the tertiary structure. The salts of small cation (e.g. Na$^+$) are generally highly water-soluble and possess low surface. On the other hand, the salts of large cation (e.g. Cs$^+$) are generally insoluble in water and possess high surface area [1].

Since Keggin-type heteropoly compounds had been discovered by Berzelius, a number of structural types of heteropoly compounds such as Wells-Dawson- [2], Finke-Droege- [3], and Preyssler-type [4] have been discovered and structurally defined. Several geometrical isomers have also been discovered. Many structural classes of HPAs with a variation of molecular sizes, compositions, and structures are now investigated in the various fields of applications. Figure 1.2 shows the several structures that can be derived from Keggin- and Wells-Dawson-type heteropolyanions.
Fig. 1.1. Schematic representations of (a) primary structure, (b) secondary structure, and (c) tertiary structure of Keggin-type HPAs.
Fig. 1.2. Several structures that can be derived from Keggin- and Wells-Dawson-type heteropolyanions.
1.2. Characteristics of HPAs

1.2.1. General properties and applications

Physicochemical characteristics of HPAs that make them well suitable for the various fields of applications are listed as followings [5].

- Metal oxide-like material
- Stable under water/air and thermally stable
- Large size of heteropolyanion
- Discrete structure
- Anionic charge
- High ionic weight
- Fully oxidizable/reducible
- Variety in oxidation numbers for the addenda atoms
  \[(E_{1/2} = +0.5 \text{ to } -1.0 \text{ V vs. SCE})\]
- Color of oxidized forms/color of reduced forms
- Photoreducible
- Arrhenius acids (pKₐ < 0)
- Can be incorporated more than 70 elements and form large number of structures
- Soluble in water and other oxygen carrying solvents (ethers, alcohols, ketones);
  also soluble or transferable into nonpolar solvents
- Hydrolysable to form deficient structures
The major applications of HPAs are found in the fields of catalysis. About 80-85% of patents and applied literatures are related to the catalysis. The remaining 15-20% of applications include coatings, analytical chemistry, membranes, sensors, dye, electrochemistry, capacitors, dopants, and clinical analysis (Table 1.2).

Most of all, excellent acid property is one of the most representative characteristics of HPAs. “Heteropolyacids”, which are acid form of heteropoly compounds, are known to considerably stronger Brønsted acid than other mineral acids and solid acids. Therefore, most of the early researches on HPAs are focused on their acid properties. Free acids of Keggin-type and Wells-Dawson-type heteropolyanions could be isolated from aqueous solution by the “etherate method” introduced by Drechsel [6]. Recently, another route for the conversion of heteropolyanion salts to the corresponding free acids using ion exchange resins was also reported [7]. Large-scale industrial processes using HPAs as acid catalysts were already developed. Several examples are hydration of isobutene to tert-butyl alcohol and polymerization of tetrahydrofuran (THF) to polyoxytetramethylene glycol (PTMEG), developed by Asahi Chemical Co. [8].

Another important characteristic of HPAs is the robust structure. Both chemical and thermal stabilities are very important in the catalytic applications. HPAs are highly stable in the most of oxidation states and exhibit inherent stability under oxidizing environments. Some HPAs are thermally stable up to ca. 600 °C and applicable in gas-phase reactions required harsh environments. It has been reported that thermal stability showed a variation with respect to counter-
cations in case of PMo$_{12}$O$_{40}^{3-}$; Ba$^{2+}$, Co$^{2+}$ (673 K, decomposition temperature) < Cu$^{2+}$, Ni$^{2+}$ (683 K) < H$^+$, Cd$^{2+}$ (693 K) < Ca$^{2+}$, Mn$^{2+}$ (700 K) < Mg$^{2+}$ (710 K) < La$^{2+}$, Ce$^{3+}$ (730 K) [9]. It was known that thermal changes in structures proceeds stepwise. Loss of water of crystallization occurs at the temperature below 473 K. At the higher temperature, acidic protons bound to the framework oxygen are released as a water (constitutional water) and HPAs finally decomposed to oxides.
<table>
<thead>
<tr>
<th>HPAs</th>
<th>Soluble</th>
<th>Hydrolytic stability</th>
<th>Thermal stability</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{H}<em>2\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}$</td>
<td>$\text{H}_2\text{O}$, organic</td>
<td>pH ≤ 6</td>
<td>≤ 600 °C</td>
</tr>
<tr>
<td>$\text{H}<em>2\text{P}<em>2\text{Mo}</em>{16}\text{O}</em>{62}$</td>
<td>$\text{H}_2\text{O}$, organic</td>
<td>-</td>
<td>375 °C</td>
</tr>
<tr>
<td>$(\text{NH}_4)<em>6\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}$, $\text{K}<em>6\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}$</td>
<td>$\text{H}_2\text{O}$, hot $\text{H}_2\text{O}$</td>
<td>pH = 3-6 for oxidized species</td>
<td>≤ 500 °C for $(\text{NH}_4)<em>6\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}$, pH = 0-13 for reduced species $\text{K}<em>6\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}$</td>
</tr>
<tr>
<td>$(\text{NH}_4)<em>6\text{As}<em>3\text{W}</em>{18}\text{O}</em>{62}$</td>
<td>for K⁺ salt</td>
<td>-</td>
<td>≤ 570 °C</td>
</tr>
<tr>
<td>$(\text{NH}_4)<em>6\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62}/\text{TiO}_2$</td>
<td>-</td>
<td>-</td>
<td>≤ 400 °C</td>
</tr>
<tr>
<td>$(\text{NH}_4)<em>6\text{As}<em>3\text{Mo}</em>{16}\text{O}</em>{62}$</td>
<td>$\text{H}_2\text{O}$</td>
<td>pH &lt; 5.3</td>
<td>-</td>
</tr>
<tr>
<td>$\text{P}<em>2\text{W}</em>{10}\text{M}(\text{H}_2\text{O})<em>2\text{O}</em>{68}^{10\text{h}}$</td>
<td>$\text{H}_2\text{O}$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$(\text{M} = \text{Co}^{2+}, \text{Cu}^{2+}, \text{Zn}^{2+})$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\text{Z}<em>2\text{P}<em>2\text{W}</em>{17}\text{O}</em>{60}(\text{M}^{2+}·\text{L})$</td>
<td>$\text{H}_2\text{O}$ for oxidized species</td>
<td>6.0 ≤ pH ≤ 7.0, for $\text{M}^{2+} = \text{Co}^{2+}$</td>
<td>-</td>
</tr>
<tr>
<td>$(\text{X} = \text{K}^+, \text{Li}^+, \text{Na}^+, \text{Bu}_4\text{N}^+)$</td>
<td>$(\text{Z} = \text{K}^+, \text{L} = \text{H}_2\text{O})$</td>
<td>5.5 ≤ pH ≤ 9.0</td>
<td>-</td>
</tr>
<tr>
<td>$(\text{Z} = \text{K}^+, \text{L} = \text{H}_2\text{O})$</td>
<td>$(\text{Z} = \text{Bu}_4\text{N}^+, \text{L} = \text{Br})$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$\text{P}<em>2\text{W}</em>{10}\text{Nb}<em>2\text{O}</em>{62}\text{X}_9$</td>
<td>$\text{H}_2\text{O}$</td>
<td>-</td>
<td>473 °C, 492 °C, 529 °C, 534 °C</td>
</tr>
<tr>
<td>$(\text{X} = \text{K}^+, \text{Li}^+, \text{Na}^+, \text{Bu}_4\text{N}^+)$</td>
<td>$(\text{K}^+, \text{Li}^+, \text{Na}^+, \text{Bu}_4\text{N}^+)$ salts</td>
<td>-</td>
<td>for $\text{Li}^+, \text{Na}^+, \text{K}^+$, $\text{Cs}^+$, respectively</td>
</tr>
<tr>
<td>$\text{Na}_{16}[\text{M}_4(\text{H}_2\text{O})<em>3(\text{P}<em>2\text{W}</em>{18}\text{O}</em>{62})_3]$</td>
<td>$\text{H}_2\text{O}$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$(\text{M} = \text{Mn}^{2+}, \text{Ni}^{2+}, \text{Co}^{2+}, \text{Cu}^{2+}, \text{Zn}^{2+})$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$[\text{n}-(\text{C}<em>9\text{H}</em>{18})_2\text{N}][(\text{C}<em>6\text{H}<em>3\text{TiP}</em>{2}\text{W}</em>{15}\text{V}<em>5\text{O}</em>{62})]$</td>
<td>Organic solvent</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Table 1.2
Other applications of HPAs [5]

<table>
<thead>
<tr>
<th>HPAs</th>
<th>Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{H}<em>3\text{P}\text{Mo}</em>{12}\text{O}_{40}$, $\text{H}<em>4\text{P}\text{Mo}</em>{10}\text{V}<em>2\text{O}</em>{40}$</td>
<td>Electrodes, capacitors</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{Mo}</em>{10}\text{O}<em>{40}$, $\text{H}<em>3\text{P}\text{W}</em>{12}\text{O}</em>{40}$</td>
<td>Electrolytic capacitors</td>
</tr>
<tr>
<td>$(\text{NH}_4)<em>3\text{H}<em>2\text{W}</em>{12}\text{O}</em>{43}$, $(\text{NH}_4)<em>3\text{P}<em>2\text{W}</em>{12}\text{O}</em>{42}$</td>
<td>Corrosion inhibitors of Al alloys</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{Mo}</em>{12}\text{O}<em>{40}$, $\text{H}<em>3\text{Si}\text{W}</em>{12}\text{O}</em>{40}$</td>
<td>Colorants for pigmenting paints, Printing inks and plastics</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{Si}\text{Mo}</em>{12}\text{O}<em>{40}$, $\text{H}<em>3\text{Si}\text{W}</em>{12}\text{O}</em>{40}$</td>
<td>Dyes for polyester and polyacrylonitrile</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{Si}\text{W}</em>{12}\text{O}_{40}$</td>
<td>Dopants of polyaniline and polypyrrole</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{W}</em>{12}\text{O}_{40}$</td>
<td>Liquid $\text{H}_2\text{O}_2$ fuel cell</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{W}</em>{12}\text{O}_{40}$</td>
<td>Photochromic coatings for copiers</td>
</tr>
<tr>
<td>$\text{K}<em>3\text{Si}\text{W}</em>{12}\text{V}<em>{40}$, $\text{Na}<em>x\text{H}</em>{5-x}\text{P}\text{Mo}</em>{13}\text{V}<em>2\text{O}</em>{40}$</td>
<td>Wood pulp bleaching</td>
</tr>
<tr>
<td>$\text{PW}<em>{11}\text{O}</em>{39}\text{Fe}^{3+}(\text{OH}_2)^+$</td>
<td>Electrode for NO determination</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{Mo}</em>\text{V}<em>2\text{O}</em>{40}$</td>
<td>Porous support for the purification of vent air</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{P}\text{Mo}</em>{10}\text{V}<em>2\text{O}</em>{40}$</td>
<td>Decontamination of mustard (HD) analogues</td>
</tr>
<tr>
<td>$\text{H}<em>3\text{Si}\text{W}</em>{12}\text{O}_{40}$ disk/$\text{SnO}_2$</td>
<td>Electrochromic display device</td>
</tr>
</tbody>
</table>
1.2.2. Redox properties

The addenda atoms in heteropolyanion frameworks are in their highest oxidation states. Therefore, HPAs are suitable for acting as an oxidizing agent. It is well known that Keggin-type molybdophosphates are easily reduced to heteropoly blue species and their reducibility can be rationalized by considering the local environments of the metal centers [11].

Metal centers in heteropolyanion frameworks occupy octahedral sites with one or two terminal oxygen atoms and represent as a mono nuclear coordination complexes, \{MOL$_5$\} or \{MO$_2$L$_4$\}. The ideal local symmetry of a metal center with mono-oxo metal centers, is C$_{4v}$. Occupation of b$_2$ by one or two electrons have no significant effect on the metal-ligand bond orders. The lowest unoccupied molecular orbital (LUMO) of Keggin-type HPAs are symmetry-adapted sets of d$_{xy}$-like metal orbitals of E symmetry. In brief, the LUMO is a non-bonding metal-centered orbital. In contrast, cis-di-oxo metal center do not have “non-bonding” d orbital, but anti-bonding with respect to the terminal M=O bonds. Therefore, stable d$^1$-complexes are not available. Some examples of reducible and irreducible heteropolyanions are summarized in Table 1.3.

In general, type I heteropolyanions are reduced easily and reversibly to form mixed-valence species (heteropoly blues). However, type II heteropolyanions are hard to be reduced. For this reason, only type I heteropoly compounds, have attracted great attention as catalysts for the redox reactions.

There were lots of efforts to elucidate the redox mechanism of HPAs. Followings are reduction mechanisms of HPAs proposed by Mizuno et al. [12].
The reduction of \( \text{H}_3\text{PM}_{12}\text{O}_{40} \) (\( \text{M} = \text{Mo, W} \)) by hydrogen proceeds in the following three steps:

\[
x\text{H}_2 \text{ (gas)} + \text{PM}_{12}\text{O}_{40}^{3-} \text{ (surface)} \leftrightarrow 2x\text{H}^+ \text{ (bulk)} + \text{PM}_{12}\text{O}_{40}^{(3+2x)} \text{ (bulk)} \quad \text{(I)}
\]
\[
2x\text{H}^+ \text{ (bulk)} + \text{PM}_{12}\text{O}_{40}^{(3+2x)} \text{ (bulk)} \leftrightarrow \text{PM}_{12}\text{O}_{40-x}^{-} \text{ (bulk)} + x\text{H}_2\text{O} \text{ (bulk)} \quad \text{(II)}
\]
\[
x\text{H}_2\text{O} \text{ (bulk)} \leftrightarrow x\text{H}_2\text{O} \text{ (gas)} \quad \text{(III)}
\]

First, a hydrogen is dissociated to protons and electrons: \( \text{H}_2 \rightarrow 2\text{H}^+ + 2\text{e}^- \). Electrons are trapped by heteropolyanions, converting \( \text{M}(\text{VI}) \) ions into \( \text{M}(\text{V}) \) ions. In the second step, protons react with framework oxygen of heteropolyanion and are evolved as water. Further reduction brings about irreversibly reduced species. However, redox behaviors of HPAs can be affected by a number of variables (structures, compositions, or nature of metal ions) and redox behaviors of several HPAs still remain question marks.
Table 1.3
Examples of type I (reducible) and type II (irreducible) heteropolyanions [11]

<table>
<thead>
<tr>
<th>Type I</th>
<th>Type II</th>
</tr>
</thead>
<tbody>
<tr>
<td>$M_6O_{19}^{6-}$</td>
<td>$Mo_7O_{22}^{6-}$</td>
</tr>
<tr>
<td>$XM_{12}O_{40}^{n-}$</td>
<td>$ThMo_{12}O_{42}^{8-}$</td>
</tr>
<tr>
<td>$P_2M_{18}O_{62}^{6-}$</td>
<td>$NiW_6O_{32}H_6^{4+}$</td>
</tr>
<tr>
<td>$NaP_3W_{30}O_{110}^{14-}$</td>
<td>$H_2W_{12}O_{42}^{10-}$</td>
</tr>
</tbody>
</table>
1.3. Oxidation catalysis over HPAs

HPAs are known to suitable materials for the catalyst design because of their several unique properties. First, acid-base and redox properties can be easily tuned by changing the constituent elements such as counter-cations, central atoms, or addenda atoms. Second, their structures are well defined than conventional metal-oxides. This indicates that catalytic properties can be designed at atomic/molecular level and theoretical cluster modeling is feasible. Catalytic oxidations over HPAs have been extensively studied over the past few decades [1]. Especially, the mixed-addenda HPAs have attracted recent attention because of their variety in stoichiometric combination and outstanding catalytic performance in oxidation reactions.

Examples of oxidation catalysis over HPAs are summarized in Table 1.5. Liquid-phase reactions include electrocatalytic oxidation of NADH [13], photocatalytic dehydrogenation of alcohols [14], and aerobic oxidation of alkyl aromatics [15]. Gas-phase reactions include oxidative dehydrogenation of alkanes [16,17] and selective oxidation of methanol [18].

It has been suggested that the aerobic oxidation of aromatics occurs via Mars-van Krevelen mechanism and the rate-determining step involves oxygen-transfer reaction from HPA to the intermediate radical cation [19]. In addition, a previous study [20] on the oxidative dehydrogenation of 2-propanol has demonstrated that the rate-determining step involves the reduction of HPA with β-hydrogen eliminated from the substrate and the intrinsic reaction rate is well correlated with the reducibility of HPA. In both cases, redox transitions of HPAs
were observed during the oxidation reactions and the rate-determining steps involved the reduction of HPAs. It is obvious that comprehensive understandings about redox properties of HPAs are of great importance in designing HPAs as oxidation catalysts.

Due to the importance of redox properties of HPAs in oxidation catalysis, comprehensive researches including quantum chemical molecular orbital study [21], hydrogen-temperature programmed reduction (TPR) measurements [22], and electrochemical analysis [23] have been conducted to examine the redox properties of HPAs. Among these, the most conventional experimental methods to probe redox properties of HPAs are electrochemical analysis and TPR measurements. Unfortunately, electrochemical analysis is strongly affected by a number of experimental conditions (pH, electrolyte, and electrode) and HPAs are not generally suitable for the dissociative adsorption of hydrogen in the TPR measurements. For this reason, only limited information on the redox properties of several HPAs is available. It is obvious that there is a need for the systematic studies about redox properties of HPAs using reliable experimental techniques.
Table 1.5
Oxidation catalysis over HPAs [10,24]

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Catalyst</th>
<th>Reaction conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toluene and nitrobenzene oxidation with H₂O₂</td>
<td>K₃P₂W₁₅Mo₂VO₆₂</td>
<td>Acetonitrile as solvent, 298 K</td>
</tr>
<tr>
<td>Aerobic oxidation of aldehydes to carboxylic acids</td>
<td>C₆H₅P₂W₁₇</td>
<td>Dichloromethane as solvent, refluxing for 30 h under O₂</td>
</tr>
<tr>
<td>Aerobic oxidation of alkyl aromatics</td>
<td>K₄H₃F₆Na₂VW₆₄O₅₆</td>
<td>Biphasic reaction media pH =5 under O₂ (5 atm), 393 K</td>
</tr>
<tr>
<td>Electrocatalytic oxidation of NADH to NAD⁺ derivatives</td>
<td>P₂Mo₃W₁₅O₆₄₁⁰⁻</td>
<td>pH = 8</td>
</tr>
<tr>
<td>Photochemical degradation of thiothers</td>
<td>(NH₄)₆P₂W₁₃O₆₂</td>
<td>298 K, acetonitrile as solvent hₐ=550-10⁵ W</td>
</tr>
</tbody>
</table>

\[
\text{CH}_2 + \text{O}_2 \rightarrow \text{CH}_2\text{O}
\]
\[
\text{CH}_3 \text{CH} = \text{CH}_2 + \text{O}_2 \rightarrow \text{CH}_3\text{CH} = \text{CH}_2 + \text{HO}
\]
\[
\text{CH}_2 = \text{CH}_2 + \text{O}_2 \rightarrow \text{CH}_2 = \text{CH}_2 + \text{H}_2\text{O}
\]
\[
\text{OH} + \text{O}_2 \rightarrow \text{O}
\]
\[
\text{Ph} + \text{O}_2 \rightarrow \text{PhCO}
\]
Chapter 2. Experimental

2.1. Mo(VI)-substituted Wells-Dawson-type tungstophosphates

2.1.1. Preparation

H₆P₂W₁₈₋ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs were prepared by “etherate method” according to the method reported in the previous literature [25] using Na₂MoO₄·2H₂O, Na₂HPO₄·2H₂O, Na₂WO₄·2H₂O, diethyl ether, H₂SO₄, and HNO₃.

In case of H₆P₂W₃Mo₁₅O₆₂, 150 g of Na₂MoO₄·2H₂O and 40 g of Na₂WO₄·2H₂O were separately dissolved in 320 ml and 50 ml of distilled water, respectively. After complete dissolution, two solutions were mixed. Equivalent Na₂HPO₄·2H₂O aqueous solution were subsequently added to the mixed solution containing molybdenum and tungsten precursors. The mixed solution was treated with 80 ml of H₂SO₄ and it was refluxed for 8 h to form the heteropolyanion frameworks. After cooling the solution, it was extracted with diethyl ether. The “etherate” was dried in a convection oven to obtain a crude solid product. The yellow-colored solid was recrystallized from boiling water (small amounts of HNO₃ was added to avoid the formation of reduced species) to obtain the H₆P₂W₃Mo₁₅O₆₂. All the catalysts were thermally treated at 300 °C in a stream of nitrogen prior to characterization and catalytic reaction.
2.1.2. Characterization

2.1.2.1. Formation of HPA structure

Successful formation of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs frameworks was confirmed by diffuse reflectance infrared Fourier transform (DRIFT) spectroscopy using a Nicolet 6700 (Nicolet) spectrometer. Chemical compositions were confirmed by inductively coupled plasma-atomic emission spectroscopy (ICP-AES) analyses using an ICPS-1000IV (Shimadzu) instrumentation.

2.1.2.2. Electrochemical analysis

Cyclic voltammograms of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs were obtained by an electrochemical method in solution. Electrochemical measurements were performed using a conventional three-electrode system (Autolab 302N, Eco Chemie). Glassy carbon with diameter of 3.0 mm was used as a working electrode. Platinum rod and saturated calomel electrode (KCl saturated) were used as a counter electrode and a reference electrode, respectively. 0.5 M Na$_2$SO$_4$ aqueous solution was used as electrolyte. Each HPA sample (1 mM) was dissolved in electrolyte and cyclic voltammograms were acquired at a scan rate of 50 mV/s. Samples were purged for 20 min under He flow prior to electrochemical measurements.
2.1.2.3. UV-visible spectroscopy

UV-visible spectra of H₆P₂W₁₈₋ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs were obtained using a Lambda-35 (Perkin-Elmer) spectrometer. 1 mM of sample solution was prepared for UV-visible spectroscopy measurement. The Kubelka-Munk function [26] was used to convert reflectance into equivalent absorption spectrum using BaSO₄ as a standard. Absorption edge energy was directly obtained from the [F(R∞)·hν]¹/² curve.

2.1.2.4. Scanning tunneling microscopy

Scanning tunneling microscopy (STM) and tunneling spectroscopy measurements were carried out in air using an Autoprobe CP (Thermomicroscope) instrument with mechanically formed Pt/Ir (90/10) tip. Samples for STM studies were prepared as followings. A certain amount of each HPA was dissolved in water to prepare 0.01 M aqueous HPA solution. A drop of sample solution was then deposited on freshly cleaved highly oriented pyrolytic graphite (HOPG) surface and it was allowed to dry in atmosphere for 1 h.

STM image was acquired in the constant current mode at a tunneling current of 1-2 nA and a sample bias of 100 mV. Tip was calibrated by imaging bare HOPG and measuring periodicity of bare HOPG. The image presented in this work was not filtered, and the reported periodicity and included angle are average values determined by performing 2-dimensional fast Fourier transform (2D FFT) analyses. For tunneling spectroscopy measurement, STM probe was
positioned above the HPA molecule of interest and the tunneling current was monitored while the bias voltage was ramped from -2.0 to +2.0 V. The voltage axis in the tunneling spectrum represents the potential applied to the sample relative to that of tip.
2.1.3. Catalytic test (oxidative dehydrogenation of ethanol)

Gas-phase oxidative dehydrogenation of ethanol over $H_{6}P_{2}W_{18-x}Mo_{x}O_{62}$ ($x=0, 3, 6, 9, 15, 18$) Wells-Dawson HPA catalysts was carried out in a continuous flow fixed-bed reactor at atmospheric pressure. Each catalyst (0.5 g) was charged into a tubular quartz reactor, and pretreated with a mixed stream of nitrogen (10 ml/min) and oxygen (10 ml/min) at 260 °C for 1 h. Ethanol ($3.43 \times 10^{-3}$ mol/h) was sufficiently vaporized by passing through the pre-heating zone and continuously fed into the reactor together with oxygen and nitrogen carrier. The feed composition (molar ratio) was fixed at ethanol (0.2): oxygen (1.0): nitrogen (1.0). The catalytic reaction was carried out at 240 °C for 5 h under steady-state conditions. The products were periodically sampled and analyzed with a gas chromatograph (HP 5890II), equipped with a capillary column (Supelco, VOCOL, 60 m × 1.5 μm × 0.25 mm). The conversion of ethanol and yield for acetaldehyde were calculated based on carbon balance.
2.2. V(V) and Nb(V)-substituted Wells-Dawson-type tungstophosphates

2.2.1. Preparation

α₂-K₇P₂W₁₇M₁O₆₂ (M=V and Nb) Wells-Dawson HPAs were prepared according to the method reported in the literature [27]. For the comparison, α-K₆P₂W₁₈O₆₂ and α₂-K₆P₂W₁₇Mo₁O₆₂ Wells-Dawson HPAs were also prepared. The catalysts were prepared by direct incorporation of transition metal into the framework of mono-lacunary heteropolyanions, α₂-P₂W₁₇O₆₁₀⁻. This method is proper than “etherate method” to prepare the structurally well-defined heteropolyanion because only uniform site (cap site) is available for metal-substitution in the mono-lacunary heteropolyanions.

2.2.1.1. Preparation of tungstophosphate (α-K₆P₂W₁₈O₆₂)

α-K₆P₂W₁₈O₆₂ Wells-Dawson HPA was prepared according to the method reported in the literature [28]. 300 g of Na₂WO₄·2H₂O dissolved in 350 ml deionized water was acidified by addition of 50 ml of 4 M HCl aqueous solution. After complete dissolution, 250 ml of 4 M H₃PO₄ aqueous solution was slowly added. The pale yellow and limpid solution was refluxed for 24 h. The resulting solution was cooled and treated with excess amounts of KCl. The precipitate was filtered and air-dried to obtain the crude product. The crude product was dissolved in 650 ml deionized water and it was heated to 80 °C. After heating for
72 h, the resulting solution was cooled and placed in refrigerator at 4 °C overnight. After few days, yellow crystals of α-K₆P₂W₁₈O₆₂ Wells-Dawson HPA were collected and air-dried.

2.2.1.2. Preparation of mono-lacunary tungstophosphate (α₂-K₁₀P₂W₁₇O₆₁)

Wells-Dawson-type mono-lacunary tungstophosphate (α₂-K₁₀P₂W₁₇O₆₁) was prepared according to the reported method [29]. 70 g of α-K₆P₂W₁₈O₆₂·14H₂O was dissolved in 250 ml of deionized water. 1 M aqueous solution of KHCO₃ was added dropwise into α-K₆P₂W₁₈O₆₂ solution, and pH of the solution adjusted to 7.5. After 30 minutes stirring, white precipitate was collected and dried. The crude white crystal was then recrystallized from boiling water. The white crystal was washed with deionized water, ethanol, and diethyl ether. The resulting solid was dried at 50 °C overnight, and α₂-K₁₀P₂W₁₇O₆₁·15H₂O was obtained.

2.2.1.3. Preparation of α₂-K₇P₂W₁₇M₁O₆₂ (M=V and Nb)

Typical procedures for the preparation of α₂-K₇P₂W₁₇M₁O₆₂ (M=V and Nb) Wells-Dawson HPAs are similar. For example, α₂-K₇P₂W₁₇Nb₁O₆₂ was prepared as followings. 0.360 g of NbCl₅ was dissolved in 150 ml of oxalic acid aqueous solution. 5 g of α₂-K₁₀P₂W₁₇O₆₁ was dissolved in 100 ml of boiling water. After complete dissolution, two solutions were then mixed. 35% HCl was added to the mixed solution until pH of the solution becomes <1.0. The mixed solution
was refluxed for 4 h. The resulting solution was cooled down to room temperature and treated with excess amounts of KCl. The white precipitates were collected and washed with deionized water several times to obtain the crude product. The crude product were recrystallized from boiling water to obtain $\alpha_2$-$K_2P_2W_{17}NbO_{62}$. $\alpha_2$-$K_2P_2W_{17}V_1O_{62}$ Wells-Dawson HPA was prepared by similar method using NaVO$_3$ as a vanadium source. For the comparison, $\alpha_2$-$K_6P_2W_{17}MoO_{62}$ Wells-Dawson HPA was prepared similar method using Na$_2$MoO$_4$·2H$_2$O as a molybdenum source. All the catalysts were thermally treated at 300 °C in a stream of nitrogen prior to characterization and catalytic reaction.
2.2.2. Characterization

2.2.2.1. Formation of HPA structure

Successful formation of $\alpha_2$-$K_7P_2W_{17}M_{1}O_{62}$ (M=V and Nb) Wells-Dawson HPAs were confirmed by diffuse reflectance infrared Fourier transform (DRIFT) spectroscopy using a Nicolet 6700 (Nicolet) spectrometer. Chemical compositions were confirmed by inductively coupled plasma-atomic emission spectroscopy (ICP-AES) analyses using an ICPS-1000IV (Shimadzu) instrumentation. $^{31}$P NMR measurement (AVANCE 600, Bruker) was also performed using D$_2$O as a solvent, and 85% aqueous H$_3$PO$_4$ was used as an external standard.

2.2.2.2. Electrochemical analysis

Reduction potentials of $\alpha_2$-$K_7P_2W_{17}M_{1}O_{62}$ (M=V and Nb) Wells-Dawson HPAs were measured by an electrochemical method in solution. 1 mM sample solutions were prepared by dissolving each catalyst in 0.5 M Na$_2$SO$_4$ solution. The sample solutions were then purged with He for 20 min, prior to the cyclic voltammetry measurements. Electrochemical experiments were performed using a Potentiostat/Galvanostat (Eco Chemie, Autolab 302N) instrument with a computer-controlled cyclic voltammetry system. The working electrode consisted of glassy carbon with an electrode diameter of 3.0 mm. A Pt rod and saturated calomel electrode (KCl saturated) were used as a counter electrode and
a reference electrode, respectively. The cyclic voltammograms were obtained at a scan rate of 25, 50, 100, 150, and 200 mV/s.

2.2.2.3. UV-visible spectroscopy

UV-visible spectroscopy of $\alpha_2$-$K_7P_2W_{17}M_1O_{62}$ (M=V and Nb) Wells-Dawson HPAs was performed at room temperature using a Lambda-35 spectrometer (Perkin-Elmer). 1 mM aqueous solution of each catalyst was used for UV-visible spectroscopy measurement. The Kubelka-Munk function ($F(R_{\infty})$) [26] was used to convert reflectance measurements into equivalent absorption spectra using the reflectance of BaSO$_4$ as a reference. Absorption edge energies was directly obtained from the $[F(R_{\infty}) \cdot h\nu]^{1/2}$ curves.

2.2.2.4. Scanning tunneling microscopy

0.01 M aqueous solutions of $\alpha_2$-$K_7P_2W_{17}M_1O_{62}$ (M=V and Nb) Wells-Dawson HPAs were prepared. A drop of solution was deposited on HOPG surface and allowed to dry in air for 1 h. STM and tunneling spectroscopy were carried out in air using an Autoprobe CP (Thermomicroscope) instrument. Mechanically formed Pt/Ir (90/10) tips were used as a probe. STM tips were calibrated by imaging bare HOPG and confirming its standard periodicity (2.46 Å). Scanning was done in the constant current mode at a tunneling current of 1-2 nA and a sample bias of +100 mV. Lattice constants of unit cell represented average values determined by performing two-dimensional fast Fourier transform (2D FFT)
analyses.

To acquire the tunneling spectrum, sample bias voltage was ramped from -2.0 to +2.0 V with respect to the tip and the tunneling current was monitored. The voltage axis in the tunneling spectrum represented the potential applied to the sample relative to that of the tip. Current-voltage spectra were measured several times with three different tips to obtain more accurate and reproducible results, and to provide a basis for statistical analyses.
2.2.3. Catalytic test (oxidative dehydrogenation of benzylamine)

Gas-phase oxidative dehydrogenation of benzylamine over $\alpha_2$-K$_7$P$_2$W$_{17}$M$_1$O$_{62}$ (M=V and Nb) was carried out in a fixed-bed reactor under atmospheric pressure. The reaction was also carried out over $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$ and $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$ Wells-Dawson HPAs. 0.4 g of each HPA was charged into a tubular pyrex reactor and pretreated with a mixed stream of nitrogen (30 ml/min) and oxygen (10 ml/min) at 300 °C for 1 h. Benzylamine (0.5 ml/h) was sufficiently vaporized by passing through a preheating zone and it was continuously fed into the reactor together with nitrogen (30 ml/min) and oxygen (10 ml/min). Catalytic reaction was carried out at 300 °C for 5 h. Reaction products were periodically sampled and analyzed using a gas chromatograph (YL6100 GC, Younglin) equipped with a flame ionization detector. DB-5 (Agilent, 60 m × 0.32 mm) capillary column was used for product separation. The conversion of benzylamine and yield for products were calculated based on carbon balance.
2.3. Keggin-type heteropolytungstates with different central atom

2.3.1. Preparation

2.3.1.1. Preparation of $\alpha$-H$_3$PW$_{12}$O$_{40}$ and $\alpha$-H$_4$SiW$_{12}$O$_{40}$

Commercially available $\alpha$-H$_3$PW$_{12}$O$_{40}$ (Kanto Chem.) and $\alpha$-H$_4$SiW$_{12}$O$_{40}$ (Fluka) were purchased. The catalysts were thermally treated at 300 °C in a stream of nitrogen prior to characterization and catalytic reaction.

2.3.1.2. Preparation of $\alpha$-H$_5$BW$_{12}$O$_{40}$

$\alpha$-K$_5$BW$_{12}$O$_{40}$ was prepared by according to the method as described in the literature [30]. 100 g of Na$_2$WO$_4$·2H$_2$O was dissolved in the 100 ml of deionized water. 5 g of boric acid and 60 ml of 6 M HCl aqueous solution was added to the solution containing tungsten precursor. After complete dissolution, the solution was refluxed for 4 h and allowed to cool down. The solution was filtered and the filtrate was acidified by 6 M HCl aqueous solution until pH becomes 2.0. The solution was refluxed for 0.5 h and treated by 20 g of KCl. The white precipitate of $\alpha$-K$_5$BW$_{12}$O$_{40}$ was obtained after the filtration, washing with ether, and recrystallization. Potassium salt was converted to acid form by etherate method or ion exchange process using ion exchange resin (Dowex HCR-W2 hydrogen form, Sigma-Aldrich). The catalyst was thermally treated at 300 °C in a stream
of nitrogen prior to characterization and catalytic reaction.

2.3.1.3. Preparation of α-H₆CoW₁₂O₄₀

α-H₆CoW₁₂O₄₀ was prepared to the methods as described in the literature [31]. 99 g of Na₂WO₄·2H₂O was dissolved in the 200 ml of deionized water. The solution was treated by 20 ml of acetic acid (The pH of solution was adjusted to 6.5~7.5.). 12.35 g of Co(CH₃CO₂)₂·4H₂O was dissolved in 65 ml of deionized water. The solution containing Co precursor was slowly added to the solution containing W precursor. The mixture solution was gently heated for 10 min and insoluble impurities were filtered. The filtrate was treated by excess amounts of KCl aqueous solution (85 g KCl in 150 ml deionized water) and stirred for an hour. The resulting solution was allowed to settle down for 24 h. The dark green precipitate (the mixture of dark blue and green crystals) was collected and treated by 200 ml of 2 M H₂SO₄. The solution turned into dark blue. The solution was concentrated by evaporation and green needles was filtered off. The solution was cooled in ice-bath. The dark blue crystals were collected after 24 h and allowed to dry in air. Potassium salt was converted to acid form by etherate method or ion exchange process using ion exchange resin (Dowex HCR-W2 hydrogen form, Sigma-Aldrich). The catalyst was thermally treated at 300 °C in a stream of nitrogen prior to characterization and catalytic reaction. The catalyst was thermally treated at 300 °C in a stream of nitrogen prior to characterization and catalytic reaction.
2.3.2. Characterization

2.3.2.1. Formation of HPA structure

Successful formation of $\alpha$-XW$_{12}$O$_{40}$\(^{n-}\) (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) heteropolyanion frameworks was confirmed by diffuse reflectance infrared Fourier transform (DRIFT) spectroscopy using a Nicolet 6700 (Nicolet) spectrometer.

2.3.2.2. Electrochemical analysis

Cyclic voltammograms of $\alpha$-XW$_{12}$O$_{40}$\(^{n-}\) (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) were measured by a conventional three-electrode system. Electrochemical measurements were carried out using a Potentiostat/Galvanostat (Eco Chemie, Autolab 302N) instrument. Glassy carbon electrode with diameter of 3.0 mm and platinum rod were used as a working electrode and a counter electrode, respectively. Saturated calomel electrode (KCl saturated) was used as a reference. Each HPA catalyst (1 mM) was dissolved in 0.5 M Na$_2$SO$_4$ aqueous solution and purged for 20 min under He flow (20 ml/min) prior to electrochemical measurement. Cyclic voltammograms were acquired at a scan rate of 25, 50, 100, 150, and 200 mV/s.
2.3.2.3. UV-visible spectroscopy

UV-visible spectra of α-XW_{12}O_{40}^{n−} (X=Co^{2+}, B^{3+}, Si^{4+}, and P^{5+}) were obtained using a Lambda-35 (Perkin-Elmer) spectrometer. 1 mM of sample solutions were prepared for UV-visible spectroscopy measurements. The Kubelka-Munk function (F(R∞)) [26] was used to convert reflectance measurement into equivalent absorption spectrum.

2.3.2.4. Scanning tunneling microscopy

Scanning tunneling microscopy and tunneling spectroscopy studies for α-XW_{12}O_{40}^{n−} (X=Co^{2+}, B^{3+}, Si^{4+}, and P^{5+}) were previously reported [32]. Data were taken from the previous literature [32].
2.4. Mo(VI)-substituted Wells-Dawson-type tungstoarsenates

2.4.1. Preparation

Wells-Dawson tungstoarsenate ($\alpha$-$K_{6}As_{2}W_{18}O_{62}$) was prepared according to the modified method in the previous literature [33]. Molybdenum-substituted Wells-Dawson tungstoarsenates ($\alpha$-$K_{6}As_{2}W_{18-x}Mo_{x}O_{62}$ ($x=1\text{-}3$)) were prepared by direct incorporation of molybdenum into mono-, di-, and tri-vacant lacunary species according to the methods described in the previous literature [34].

2.4.1.1. Preparation of $\alpha$-$K_{6}As_{2}W_{18}O_{62}$

66 g of $\text{Na}_2\text{WO}_4\cdot2\text{H}_2\text{O}$ and excess amount of $\text{Na}_2\text{HAsO}_4\cdot7\text{H}_2\text{O}$ were separately dissolved in 90 ml of deionized water. 30 ml of 4 M HCl was added into the sodium tungstate solution. Sodium hydrogen arsenate solution was then introduced into the sodium tungstate solution. The pH of the mixed solution was adjusted to ca. 1.3 using HCl. The solution was subsequently refluxed at 80 °C for 4 h with vigorous stirring. The solution was then cooled down to room temperature and treated with 40 g of KCl. The resulting precipitate was filtered, washed with saturated KCl solution, and dried under atmospheric conditions.
2.4.1.2. Preparation of mono-lacunary tungstoarsenate (α-\(K_{10}As_2W_{17}O_{61}\))

118.5 g (24 mmol) of α-\(K_6As_2W_{18}O_{62}\) was dissolved in 360 ml of boiling water. 360 ml of 1 M KHCO\(_3\) aqueous solution was added to the solution. The resulting solution was stirred for 40 min and filtered. The crude solid was recrystallized from 500 ml of boiling water to obtain α-\(K_{10}As_2W_{17}O_{61}\).

2.4.1.3. Preparation of di-lacunary tungstoarsenate (α-\(K_{11}HAs_2W_{16}O_{59}\))

39.5 g (8 mmol) of α-\(K_6As_2W_{18}O_{62}\) was dissolved in 150 ml of water. 100 ml of 2 M tris(hydroxymethyl)aminomethane was quickly added to the solution. After stirring for 5 min, the resulting solution was treated with 150 ml of saturated KCl solution. The solution was further stirred for 10 min. The white precipitates were collected by filtration, washing with 50 ml of ethanol, and drying.

2.4.1.4. Preparation of tri-lacunary tungstoarsenate (α-\(Na_{12}As_2W_{15}O_{56}\))

99 g (20 mmol) of α-\(K_6As_2W_{18}O_{62}\) was dissolved in 250 ml water. 89 g of NaClO\(_4\) was added to the solution. After 1 h, the solution was cooled down to 10 °C. The white precipitates (KClO\(_4\)) were removed by filtration. The pH of filtrate was adjusted to 9.0 using 1 M Na\(_2\)CO\(_3\) aqueous solution and it was maintained for 1 h. The resulting precipitates were collected by filtration, washing with 100 ml ethanol, and drying.
2.4.1.5. Preparation of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=1-3)

$\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=1-3) were prepared by similar methods via direct incorporation of molybdenum into mono-, di-, and tri-vacant lacunary species according to the modified methods described in the previous literature [34]. 5 g of mono-, di-, or tri-lacunary tungstoarsenates were dissolved in 20 ml of boiling water. Equivalent Na$_2$MoO$_4$·2H$_2$O was added to the solution. After complete dissolution, the pH of solution was adjusted to < 1.0 using 35% HCl aqueous solution. After stirring for 1 h, the resulting solution was treated with 4 g of KCl. The yellow precipitates were collected by filtration and recrystallized in boiling water (pH < 2.0).
2.4.2. Characterization

2.4.2.1. Formation of HPA structure

Successful formation of $\alpha$-As$_2$W$_{18-x}$Mo$_x$O$_{62}^{6-}$ ($x=0-3$) heteropolyanion frameworks was confirmed by diffuse reflectance infrared Fourier transform (DRIFT) spectroscopy using a Nicolet 6700 (Nicolet) spectrometer. Chemical compositions of the catalysts were confirmed by inductively coupled plasma-atomic emission spectroscopy (ICP-AES) analyses using an ICPS-1000IV (Shimadzu) instrumentation.

2.4.2.2. Electrochemical analysis

Cyclic voltammograms of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$) were obtained by an electrochemical method in solution. Electrochemical measurements were performed using a conventional three-electrode system (Autolab 302N, Eco Chemie). Glassy carbon with diameter of 3.0 mm was used as a working electrode. Platinum rod and saturated calomel electrode (KCl saturated) were used as a counter electrode and a reference electrode, respectively. 0.5 M Na$_2$SO$_4$ aqueous solution was used as electrolyte. Each HPA sample (1 mM) was dissolved in electrolyte and cyclic voltammograms were acquired at a scan rate of 10, 25, 50, 100, 150, and 200 mV/s. Samples were purged for 20 min under He flow (20 ml/min) prior to electrochemical measurements.
2.4.2.3. UV-visible spectroscopy

UV-visible spectra of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0\text{-}3$) were obtained using a Lambda-35 (Perkin-Elmer) spectrometer. 1 mM of sample solution was prepared for UV-visible spectroscopy measurement. The Kubelka-Munk function [26] was used to convert reflectance into equivalent absorption spectrum using BaSO$_4$ as a standard. Absorption edge energy was directly obtained from the $[F(R_\infty)\cdot h\nu]^{1/2}$ curve.

2.4.2.4. Scanning tunneling microscopy

Each $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0\text{-}3$) catalyst was dissolved in deionized water (1 mM). Each HPA solution was physically deposited on the freshly cleaved HOPG. The sample was allowed to dry under atmosphere. STM investigations were performed using a XE-100E instrument (Park Systems) equipped with mechanically formed Pt/Ir (90/10) tip. The tip was calibrated by imaging bare HOPG surface and confirming its standard periodicity (2.46 Å) prior to imaging process. STM images of nano-structured HPAs were acquired in the constant-current mode. STM image presented in this work was not filtered, and dimension of unit cell was determined from two-dimensional fast Fourier transform (2-D FFT) analysis. Tunneling spectrum of each bright corrugation in the STM image was also measured. Current-voltage response was monitored as ramping the bias from -2.0 to +2.0 V. The voltage axis in the tunneling spectra represented the potential applied to the sample relative to that of the tip.
Tunneling spectra were measured several times with three different tips to provide more reproducible and accurate results, and to provide a basis for statistical analyses.
2.4.3. Catalytic test (oxidative dehydrogenation of benzyl alcohol)

Gas-phase oxidative dehydrogenation of benzyl alcohol was carried out over \( \alpha-K_6As_2W_{18-x}Mo_xO_{62} \) (\( x=0-3 \)) catalysts. 0.5 g of HPA was charged into the tubular glass reactor and pretreated with a mixed stream of oxygen (10 ml/min) and nitrogen (20 ml/min) at 300 °C for 1 h. Vaporized benzyl alcohol (0.15 ml/h) was fed into the reactor together with a mixed stream of oxygen (10 ml/min) and nitrogen (20 ml/min). Catalytic reaction was performed at 300 °C for 5 h. Reaction products were periodically sampled and analyzed using a gas chromatograph (YL6100 GC, Younglin) equipped with a flame ionization detector. DB-5 (Agilent, 60 m × 0.32 mm) capillary column was used for product separation.
2.5. V(V) and Nb(V)-substituted Wells-Dawson-type tungstoarsenates

2.5.1. Preparation

2.5.1.1. Preparation of $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$

$\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ was prepared by similar methods via direct incorporation of vanadium into mono-lacunary species according to the modified method described in the previous literature [34]. 5 g of $\alpha_2$-K$_{10}$As$_2$W$_{17}$O$_{61}$ was dissolved in 25 ml of boiling water. Equivalent NaVO$_3$ was added to the solution. After complete dissolution, the pH of solution was adjusted to < 1.0 using 35% HCl aqueous solution. After stirring for 1 h, the resulting solution was treated with 5 g of KCl. The orange precipitates were collected by filtration and recrystallized in boiling water. For the comparison, $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ and $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ were also prepared.

2.5.1.2. Preparation of $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$

$\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ was prepared by direct incorporation of niobium into mono-lacunary species. 0.110 g of NbCl$_3$ was dissolved in 30 ml of oxalic acid aqueous solution. 1.7 g of $\alpha_2$-K$_{10}$P$_2$W$_{17}$O$_{61}$ was dissolved in 20 ml of boiling water. After complete dissolution, two solutions were then mixed. 35% HCl was added to mixed solution until pH of the solution becomes <1.0. The mixed
solution was refluxed for 4 h and it was cooled down to room temperature. The resulting solution was cooled and transfer to 300 ml of methanol. Then mixed solution was treated with excess amounts of KCl. The white precipitates were collected and washed with methanol to obtain $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$. 
2.5.2. Characterization

2.5.2.1. Formation of HPA structure

Successful formation of $\alpha_2$-K$_2$As$_2$W$_{17}$M$_1$O$_{62}$ (M=V and Nb) heteropolyanion frameworks was confirmed by diffuse reflectance infrared Fourier transform (DRIFT) spectroscopy using a Nicolet 6700 (Nicolet) spectrometer. Chemical compositions of the catalysts were confirmed by inductively coupled plasma-atomic emission spectroscopy (ICP-AES) analyses using an ICPS-1000IV (Shimadzu) instrumentation.

2.5.2.2. Electrochemical analysis

Cyclic voltammograms of $\alpha_2$-K$_2$As$_2$W$_{17}$M$_1$O$_{62}$ (M=V and Nb) were obtained by an electrochemical method in solution. Electrochemical measurements were performed using a conventional three-electrode system (Autolab 302N, Eco Chemie). Glassy carbon with diameter of 3.0 mm was used as a working electrode. Platinum rod and saturated calomel electrode (KCl saturated) were used as a counter electrode and a reference electrode, respectively. 0.5 M Na$_2$SO$_4$ aqueous solution was used as electrolyte. Each HPA sample (1 mM) was dissolved in electrolyte and cyclic voltammograms were acquired at a scan rate of 10, 25, 50, 100, 150, and 200 mV/s. Samples were purged for 20 min under He flow (20 ml/min) prior to electrochemical measurements.
2.5.2.3. UV-visible spectroscopy

UV-visible spectra of $\alpha_2$-K$_2$As$_2$W$_17$M$_1$O$_{62}$ (M=V and Nb) were obtained using a Lambda-35 (Perkin-Elmer) spectrometer. 1 mM of sample solution was prepared for UV-visible spectroscopy measurement. The Kubelka-Munk function [26] was used to convert reflectance into equivalent absorption spectrum using BaSO$_4$ as a standard. Absorption edge energy was directly obtained from the $[F(R_c)\cdot h\nu]^{1/2}$ curve.
2.5.3. Catalytic test (oxidative dehydrogenation of benzylamine)

Gas-phase oxidative dehydrogenation of benzyl amine was carried out over $\alpha_2$-K$_7$As$_2$W$_{17}$M$_1$O$_{62}$ (M=V and Nb) catalysts. The reaction was also carried out over $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ and $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ Wells-Dawson HPAs. 0.4 g of each HPA was charged into a tubular pyrex reactor and pretreated with a mixed stream of nitrogen (30 ml/min) and oxygen (10 ml/min) at 300 °C for 1 h. Benzylamine (0.5 ml/h) was sufficiently vaporized by passing through a preheating zone and it was continuously fed into the reactor together with nitrogen (30 ml/min) and oxygen (10 ml/min). Catalytic reaction was carried out at 300 °C for 5 h. Reaction products were periodically sampled and analyzed using a gas chromatograph (YL6100 GC, Younglin) equipped with a flame ionization detector. DB-5 (Agilent, 60 m x 0.32 mm) capillary column was used for product separation. The conversion of benzylamine and yield for products were calculated based on carbon balance.
Chapter 3. Results and Discussion

3.1. Mo(VI)-substituted Wells-Dawson-type tungstophosphates

3.1.1. Formation of HPA structure

Figure 3.1 shows the schematic diagram for the preparation procedures of \( H_6P_2W_{18-x}Mo_xO_{62} \) (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. Wells-Dawson heteropolyanion, \([X_2M_{18}O_{62}]^6\) (X=central atom, M=addenda atom), is a polymeric oxoanion formed by condensation of more than two different o xoanions. The molecular structure of Wells-Dawson heteropolyanion could be considered as a combination of two defected Keggin units, \([XMoO_{34}]^6\); Two defected units were linked through six metal-oxygen-metal bonds. In this work, \( H_6P_2W_{18-x}Mo_xO_{62} \) with x=0, 3, 9, 15, and 18 were denoted as \( P_2W_{18} \), \( P_2W_{15}Mo_3 \), \( P_2W_9Mo_9 \), \( P_2W_3Mo_{15} \), and \( P_2Mo_{18} \), respectively.

Figure 3.2 shows the DRIFT spectra of \( H_6P_2W_{18-x}Mo_xO_{62} \) (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. Four characteristic bands attributed to P-O, M=O, and M-O-M (M=W or Mo) were clearly observed in the range of 700 - 1200 cm\(^{-1}\). Detailed assignment of each band of \( H_6P_2W_{18-x}Mo_xO_{62} \) (x=0, 3, 9, 15, 18) Wells-Dawson HPAs is summarized in Table 3.1. A single band for P-O stretching vibration was observed at 1067 - 1081 cm\(^{-1}\). Three bands for M=O\(_t\) (O\(_t\): terminal oxygen), M-O\(_c\)-M (O\(_c\): corner-shared oxygen), and M-O\(_e\)-M (O\(_e\): edge-shared oxygen) stretching vibrations were observed at 959 - 984 cm\(^{-1}\), 871 - 889 cm\(^{-1}\),
and 789 - 808 cm$^{-1}$, respectively. The characteristic bands of molybdenum-
substituted Wells-Dawson tungstophosphates slightly shifted to lower
wavenumber with increasing molybdenum content. Band positions were in good
agreement with the previous results [25].

Chemical compositions of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-
Dawson HPAs was further confirmed by ICP-AES analyses. Chemical
compositions of phosphorous, tungsten, and molybdenum in the $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs determined by ICP-AES
analyses are summarized in Table 3.2. The measured P:W:Mo ratios in the
$\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ were in good agreement with the theoretical values, indicating
successful formation of heteropolyanion framework.
Fig. 3.1. Schematic diagram for the preparation procedures of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs.
Fig. 3.2. DRIFT spectra of H$_6$P$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x$=0, 3, 9, 15, 18) Wells-Dawson HPAs.
Table 3.1
Detailed assignment of each band of H₆P₂W₁₈₋ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wavenumber (cm⁻¹)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P-O</td>
<td>M=Oₜ</td>
<td>M-Oₓ-M</td>
<td>M-Oₑ-M</td>
</tr>
<tr>
<td>P₂Mo₁₈</td>
<td>1067</td>
<td>959</td>
<td>871</td>
<td>789</td>
</tr>
<tr>
<td>P₂W₃Mo₁₅</td>
<td>1068</td>
<td>963</td>
<td>873</td>
<td>792</td>
</tr>
<tr>
<td>P₂W₉Mo₉</td>
<td>1075</td>
<td>973</td>
<td>880</td>
<td>795</td>
</tr>
<tr>
<td>P₂W₁₅Mo₃</td>
<td>1078</td>
<td>982</td>
<td>886</td>
<td>804</td>
</tr>
<tr>
<td>P₂W₁₈</td>
<td>1081</td>
<td>984</td>
<td>889</td>
<td>808</td>
</tr>
</tbody>
</table>

*Oₜ: terminal oxygen, Oₓ: corner-shared oxygen, Oₑ: edge-shared oxygen
Table 3.2
Chemical compositions of phosphorous, tungsten, and molybdenum in the H₆P₂W₁₈₋ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Ratio of P:W:Mo</th>
<th>Theoretical value</th>
<th>Measured value</th>
</tr>
</thead>
<tbody>
<tr>
<td>P₂Mo₁₈</td>
<td>2.0 : 0.0: 18.0</td>
<td>1.6 : 0.0 : 18.0</td>
<td></td>
</tr>
<tr>
<td>P₂W₃Mo₁₅</td>
<td>2.0 : 3.0: 15.0</td>
<td>1.6 : 2.8 : 15.2</td>
<td></td>
</tr>
<tr>
<td>P₂W₉Mo₉</td>
<td>2.0 : 9.0: 9.0</td>
<td>1.6 : 9.4 : 8.6</td>
<td></td>
</tr>
<tr>
<td>P₂W₁₅Mo₃</td>
<td>2.0 : 15.0: 3.0</td>
<td>2.4 : 14.9 : 3.1</td>
<td></td>
</tr>
<tr>
<td>P₂W₁₈</td>
<td>2.0 : 18.0: 0.0</td>
<td>1.8 : 18.0 : 0.0</td>
<td></td>
</tr>
</tbody>
</table>
3.1.2. Characterization

3.1.2.1. Cyclic voltammetry

Electrochemical measurement is one of the most conventional techniques to measure the redox properties of bulk HPAs. Figure 3.3 shows the cyclic voltammograms of \( \text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62} \) and \( \text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62} \) Wells-Dawson HPAs. They exhibited well-defined reversible and stepwise redox transitions during the electrochemical measurements. Overall shapes of cyclic voltammograms were retained at a scan rate up to 200 mV/s (maximum scan rate in this work, not shown here) and the peak currents increased monotonically. Each peak current was proportional to the square root of scan rate, indicating diffusion-controlled process which can be expressed by the Randles-Sevcik equation:

\[
I_p = 2.69 \times 10^5 \cdot n^{3/2} \cdot D_0^{1/2} \cdot \nu^{1/2} \cdot C_o \cdot A
\]  

(2)

where \( I_p \), \( n \), \( D_0 \), \( \nu \), \( C_o \), and \( A \) denote peak current, number of electron involved in the redox event, diffusion coefficient, scan rate, concentrations, and electrode surface area, respectively.

Several tungsten-based redox transitions were observed in the cyclic voltammogram of \( \text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62} \), while \( \text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62} \) exhibited molybdenum-based redox transitions. Interestingly, molybdenum-substituted Wells-Dawson HPAs showed an additional molybdenum-based redox transition. This indicates that molybdenum centers were more electrochemically accessible and reducible.
than tungsten centers in the heteropolyanion framework. Because the highest potential represented the reducibility of HPAs, the first electron reduction potential was taken as the representative reduction potential of HPAs [35]. First electron reduction potentials are summarized in Table 3.3. Reduction potential increased with increasing molybdenum-substitution.

The variation of reduction potential of HPAs could be understood by taking into account of their enthalpy, entropy, and electronic properties. However, enthalpy and entropy contributions on the redox properties could be neglected in the series of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs because of their identical molecular size. Therefore, it is inferred that the difference in redox properties of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs was mainly due to the difference in electronic structure caused by molybdenum-substitution.
Fig. 3.3. Cyclic voltammograms of (a) $\text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ and (b) $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ Wells-Dawson HPAs.
3.1.2.2. UV-visible spectroscopy

UV-visible spectroscopy measurements were conducted to probe the electronic structure of H₆P₂W₁₈ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. It has been demonstrated that absorption edge energy determined by UV-visible spectroscopy was well correlated with catalytic activity and reducibility in the series of supported metal oxide clusters and metatungstate [36]. In the metal oxide, it is known that absorption edge energy is generally sensitive to average size of metal oxide domains. However, HPAs in solution are molecularly discrete clusters with nearly identical size and have an advantage for applying UV-visible spectroscopy. Therefore, UV-visible spectroscopy were performed as a simple diagnostic method for energy state and reducibility of HPAs.

Absorption edge energy (energy gap) was calculated from the following Tauc equation using Kubelka-Munk function (F(R∞)) to convert the reflectance to corresponding absorption spectrum.

\[(\alpha h\nu)^{1/2} = A (h\nu - E_g)\]  \hspace{1cm} (3)

where \(\alpha\), h, \(\nu\), A, and \(E_g\) denote absorption coefficient, Plank constant, scan rate, constant, and optical energy gap, respectively. Figure 3.4 shows the \([F(R_{\infty}) \cdot h\nu]^{1/2}\) curves of H₆P₂W₁₈ₓMoₓO₆₂ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. Absorption edge energies are summarized in Table 3.3. Absorption edge energy decreased with increasing molybdenum content. The trend of absorption edge energy is well consistent with that of reduction potential. Absorption edge energy represents the
energy required for electron promotion from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO) [37]. By UV-visible irradiation, electrons are promoted from 2p-orbitals on the oxygens (the HOMO) to the low-lying high-energy states which comprise a combination of d-orbitals on the framework metal centers and 2p-orbitals on the framework oxygens (the LUMO). Thus, the energy gap between the HOMO and the LUMO decreased with increasing molybdenum content in the series of $H_{6}P_{2}W_{18-x}Mo_{x}O_{62}$. It is obvious that the HOMO is little affected by molybdenum-substitution because the HOMO mostly comprises 2p-orbitals on the framework oxygens. This indicates that the smaller absorption edge energy represents the lower energy level of the LUMO.

A previous density functional theory (DFT) study [38] about vanadium-substituted Wells-Dawson tungstophosphate showed that redox properties were closely related to energy state and composition of the LUMO, comprising an combination of d-orbitals on the metal centers and p-orbitals on the framework oxygens. In case of Wells-Dawson tungstophosphate, added electrons are generally delocalized over twelve tungsten centers at equatorial sites. In case of vanadium-substituted heteropolyanion, however, the energy state of metal centers at cap sites was stabilized by more reducible vanadium and the electrochemical reduction of vanadium centers at cap sites became available. The effect of molybdenum-substitution might be understood in a similar manner by considering the fact that molybdenum center is more reducible and contribution of molybdenum to the LUMO increases with increasing molybdenum content.
Fig. 3.4. $[F(R_\infty)\cdot h\nu]^{1/2}$ curves of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs
3.1.2.3. Scanning tunneling microscopy

Scanning tunneling microscopy measurements were performed for the further investigation about the local electronic structure of H$_6$P$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. In the STM measurements, STM image represents convolution of geometric and electronic structures rather than purely geometric structure of the specimen [39]. For adsorbate-substrate system, STM imaging would be more complex due to the presence of interaction between adsorbate and substrate. This indicates that STM image of adsorbate-covered surface may contain the electronic contributions of substrate and may not reflect the actual topography of surface. In this work, therefore, chemically inert graphite was employed as a substrate for HPA deposition and HPAs were physically deposited by solvent evaporation in order to minimize the interaction.

Figure 3.5 shows the STM images of H$_6$P$_2$W$_{18}$O$_{62}$ and H$_6$P$_2$Mo$_{18}$O$_{62}$ Wells-Dawson HPAs. Two-dimensional self-assembled HPA array was observed in the STM measurement. Each bright corrugation in the STM image represents individual heteropolyanion. The periodicity of H$_6$P$_2$W$_{18}$O$_{62}$ unit cell constructed on the basis of lattice constants determined from two-dimensional fast Fourier transform analysis (2-D FFT) was 11.2 Å × 14.4 Å with included angle of 57.2 °. The periodicity of H$_6$P$_2$Mo$_{18}$O$_{62}$ unit cell was found to 14.4 Å × 10.5 Å with included angle of 85.8 °. The dimensions of unit cells determined from STM images were in good agreement with the lattice constant determined by X-ray crystallography [2].

Figure 3.6 shows the typical current-voltage responses of H$_6$P$_2$W$_{18}$O$_{62}$ and
H$_6$P$_2$Mo$_{18}$O$_{62}$ Wells-Dawson HPAs taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.5. Current-voltage response at bare graphite was also measured for the comparison. It is noteworthy that each current-voltage response taken at interstitial space between bright corrugations was almost identical to that of bare graphite, indicating that two-dimensional HPA arrays shown in Figure 3.5 were monolayer. On the other hand, tunneling spectrum taken at bright corrugation showed a distinctive current-voltage response, referred to as negative differential resistance (NDR) phenomenon.

NDR phenomenon often explained by resonant tunneling model through a double barrier quantum well structure; tunneling electrons can pass through the molecular well without any attenuation when energy state of incident electron is in resonance with a virtual empty state in the molecular well (Figure 3.7) [40,41]. This resonance tunneling contains sequential electron transfer from negative potential to positive potential via framework oxygen and metal ions of heteropolyanion [42]. It is inferred that HPA molecule might serve as a quantum well, and resonant tunneling occurs when electrons pass through the frontier orbitals of HPA molecule [32].

The average values of NDR peak voltage were summarized in Table 3.3. It is interesting that NDR peak voltage appeared at less negative voltage with increasing the molybdenum content. This results is well consistent with the results of UV-visible spectroscopy, by considering the fact that absorption edge energy (energy gap between the HOMO and the LUMO) decreased with increasing the molybdenum content and NDR peak voltage reflects the frontier
orbital (the LUMO) of the HPAs.

Figure 3.8 shows the absorption edge energy and NDR peak voltage plotted as a function of reduction potential. It is inferred that the most stabilized energy level of the LUMO in the $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ (due to the contribution of reducible molybdenum metal centers) provided the electrochemically accessible metal centers and resulted in the highest reduction potential. The correlation also suggested that absorption edge energy and NDR peak voltage could be utilized as alternative correlating parameters for reduction potential of bulk HPAs in the series of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs.
Fig. 3.5. STM images of (a) $\text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ and (b) $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ Wells-Dawson HPAs
Fig. 3.6. Typical current-voltage responses of (a) H$_6$P$_2$W$_{18}$O$_{62}$ and (b) H$_6$P$_2$Mo$_{18}$O$_{62}$ Wells-Dawson HPAs taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.5.
Fig. 3.7. Resonant tunneling model through a double barrier quantum well structure: (a) schematic representations and (b) corresponding spectra [41].
Table 3.3.
First electron reduction potentials, absorption edge energies, and NDR peak voltages of $H_6P_2W_{18-x}Mo_xO_{62}$ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>$1^{st}$ electron reduction potential (volts vs. SCE)</th>
<th>UV-visible absorption edge energy (eV)</th>
<th>NDR peak voltage (volts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P_2Mo_{18}$</td>
<td>+0.310</td>
<td>2.58</td>
<td>-0.85</td>
</tr>
<tr>
<td>$P_2W_3Mo_{15}$</td>
<td>+0.278</td>
<td>2.82</td>
<td>-0.94</td>
</tr>
<tr>
<td>$P_2W_9Mo_9$</td>
<td>+0.216</td>
<td>2.90</td>
<td>-1.00</td>
</tr>
<tr>
<td>$P_2W_{15}Mo_3$</td>
<td>+0.209</td>
<td>3.05</td>
<td>-1.08</td>
</tr>
<tr>
<td>$P_2W_{18}$</td>
<td>+0.030</td>
<td>3.31</td>
<td>-1.13</td>
</tr>
</tbody>
</table>
Fig. 3.8. Absorption edge energy and NDR peak voltage plotted as a function of reduction potential.
3.1.3. Catalytic test

Gas-phase oxidative dehydrogenation of ethanol to acetaldehyde was carried out as a model reaction to probe oxidation catalysis of H$_6$P$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0, 3, 9, 15, 18) Wells-Dawson HPAs. Figure 3.9 shows the catalytic performance of H$_6$P$_2$Mo$_{18}$O$_{62}$ during a 6 h-reaction and yield for acetaldehyde plotted as a function of molybdenum content. The catalysts exhibited a stable catalytic performance during a 5 h-reaction. Acetaldehyde was mainly formed as an oxidation product, and ethylene/diethyl ether was produced as acid-catalyzed products. It is noteworthy that yield for acetaldehyde (oxidation product) increased with increasing molybdenum content. Among the tested catalysts, H$_6$P$_2$Mo$_{18}$O$_{62}$ which exhibited the highest reduction potential showed the best catalytic performance.

The previous extended Hückel calculation for HPA [21] suggested that hydrogen was initially transferred to reducible metal center and rate-determining step involved direct interaction between hydrogen and reducible metal center during the oxidative dehydrogenation reaction. Furthermore, the adjacent metal center has a effect on the activation energy of reaction and may affect the pre-exponential factor of rate constant by contributing to the density of accessible electronic states (unoccupied orbitals). This result suggests that reducibility and composition of the metal centers in the HPA framework play an important role in the oxidative dehydrogenation reaction.

Similarly, a previous study [20] on the oxidative dehydrogenation of 2-propanol showed that the rate-determining step involved reduction of HPA with
β-hydrogen eliminated from the substrate and the intrinsic reaction rate was well correlated with the reducibility of HPA. When considering the fact that molybdenum center is easier to be reduced in the electrochemical measurement, it is inferred that molybdenum center provides efficient active sites for oxidative dehydrogenation by contributing to the density of accessible electronic states (unoccupied orbitals).
Fig. 3.9. (a) Catalytic performance of H₆P₂Mo₁₈O₆₂ during a 5 h-reaction and (b) yield for acetaldehyde after a 5 h-reaction plotted as a function of molybdenum content.
3.2. V(V) and Nb(V)-substituted Wells-Dawson-type tungstophosphates

3.2.1. Formation of HPA structure

Figure 3.10 shows the schematic diagram for the preparation procedures of $\alpha$-$K_6P_2W_{18}O_{62}$, $\alpha_2$-$K_6P_2W_{17}MoO_{62}$, $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs. $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$ and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ were prepared to elucidate the effect of group 5-metal-substitution on the redox properties and the electronic structure. $\alpha$-$K_6P_2W_{18}O_{62}$ and $\alpha_2$-$K_6P_2W_{17}MoO_{62}$ were also prepared for the comparison. In this work, mono-transition metal-substituted Wells-Dawson HPAs were prepared by direct incorporation of transition metal into the mono-lacunary $\alpha_2$-$K_{10}P_2W_{17}O_{61}$\textsuperscript{10}. Removal of a cap WO$_6$ unit in the $\alpha$-$K_6P_2W_{18}O_{62}$ by pH control yields mono-lacunary heteropolyanion, $\alpha_2$-$P_2W_{17}O_{61}$\textsuperscript{10}. Mono-lacunary heteropolyanion generally acts as a tetradentate ligand for lanthanides and actinides, and as a pentadentate ligand for transition metals [43]. As a results, a tungsten center could be selectively replaced by a transition metal. This preparation method is very useful to yield the uniformly-substituted Wells-Dawson structure, rather than “etherate method (randomly-substituted structure)”.  

Figure 3.11 shows the DRIFT spectra of $\alpha$-$K_6P_2W_{18}O_{62}$, $\alpha_2$-$K_6P_2W_{17}MoO_{62}$, $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs. Successful formations of the heteropolyanions can be confirmed by four characteristic bands in the range of 1200-700 cm$^{-1}$ which represent P-O, M=O$\delta$, inter-octahedral M-O$_c$-M, and intra-octahedral M-O$_c$-M bonds (M=metal).
Detailed assignment of each characteristic band was summarized in Table 3.4. Four characteristic bands were clearly observed in DRIFT spectra, indicating the successful formation of the heteropolyanions.

The HPAs were also confirmed by $^{31}$P NMR analyses. Figure 3.12 shows the $^{31}$P NMR spectra of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs dissolved in D$_2$O. In $^{31}$P NMR spectra, $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$ HPAs showed a single peak at -12.6 ppm which arise from two identical central phosphorous heteroatoms. However, other HPAs showed two different peaks. Two different peaks arise from two inequivalent central phosphorous heteroatoms due to the metal-substitution. In niobium-substituted Wells-Dawson tungstophosphate, for example, a peak at -10.8 ppm corresponds to a phosphorous atom near the niobium center. Another peak at -13.1 ppm corresponds to a phosphorous atom far from the niobium center. Any other noticeable peaks were not observed except small peaks of impurity in the $^{31}$P NMR spectra.

Chemical compositions of phosphorous, tungsten, and substituted-metal in the HPAs were further confirmed by ICP-AES analyses. Chemical compositions of phosphorous, tungsten, and substituted-metal in the $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs were summarized in table 3.5. The molar ratios of P:W:M were in good agreement with the theoretical values, suggesting the successful formation of HPA structure.
Fig. 3.10. Schematic diagram for the preparation procedures of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.
Fig. 3.11. DRIFT spectra of $\alpha$-K$_8$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_8$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.
Table 3.4.
Detailed assignment of each characteristic band

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<th>Catalyst</th>
<th>P-O</th>
<th>M=O_t</th>
<th>M-O_c-M</th>
<th>M-O_e-M</th>
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<td>960</td>
<td>913</td>
<td>785</td>
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</table>

*O_t: terminal oxygen, O_c: corner-shared oxygen, O_e: edge-shared oxygen
Fig. 3.12. $^{31}$P NMR spectra of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs dissolved in D$_2$O.
Table 3.5.
Chemical compositions of phosphorous, tungsten, and substituted-metal (M) in the $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs

<table>
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<th>Measured Value</th>
</tr>
</thead>
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</tr>
<tr>
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<td>1.9 : 17.0 : 1.0</td>
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</tr>
<tr>
<td>KP$<em>2$W$</em>{17}$V$_1$</td>
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<td>1.9 : 17.0 : 1.1</td>
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</tr>
<tr>
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<td>2.2 : 17.0 : 1.1</td>
<td></td>
</tr>
</tbody>
</table>
3.2.2. Characterization

3.2.2.1. Cyclic voltammetry

Figure 3.13 shows the cyclic voltammograms of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. Four tungsten-based redox transitions were observed in the cyclic voltammogram of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$. In case of molybdenum- and vanadium-substituted Wells-Dawson tungstophosphates showed an additional molybdenum- and vanadium-based redox transitions, respectively. However, niobium-substituted Wells-Dawson tungstophosphates showed significantly shifted redox couple. This indicates vanadium or molybdenum center in the heteropolyanion framework was more electrochemically accessible and easier to be reduced than tungsten centers. Otherwise, the niobium-substitution has negative effect to the reducibility of Wells-Dawson tungstophosphates. The first electron reduction potentials are summarized in Table 3.6. Reduction potential increased in the order of $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62} < \alpha$-K$_6$P$_2$W$_{18}$O$_{62} < \alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62} < \alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$.
Fig. 3.13. Cyclic voltammograms of (a) $\alpha$-$K_6P_2W_{18}O_{62}$, (b) $\alpha_2$-$K_6P_2W_{17}Mo_1O_{62}$, (c) $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and (d) $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs.
UV-visible spectroscopy measurements were conducted to probe the bulk electronic structures of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. Figure 3.14 shows the $[F(R_\infty) \cdot h\nu]^{1/2}$ curves of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. UV-visible absorption edge energies are summarized in Table 3.6. Absorption edge energy decreased in the order of $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ > $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$ > $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$ > $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$. As mentioned above, absorption edge energy determined from the linear fit of $[F(R_\infty) \cdot h\nu]^{1/2}$ represents the energy required for electron promotion from the HOMO to the LUMO [37]. Thus, the energy gap between the HOMO and the LUMO decreased in the order of $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ > $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$ > $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$ > $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$. It could be inferred that $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ with the smallest absorption edge energy has the lowest energy level of the LUMO, because the HOMO is little perturbed by metal-substitution in the series of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.

A previous density functional theory (DFT) study [44] about the tri-substituted Keggin-type tungstosilicates showed the similar results. Theoretically calculated energy gap between the HOMO and the LUMO decreased when molybdenum/vanadium is substituted but increased when niobium is substituted in the Keggin-type tungstosilicates. They also reported that the relative energy and composition of the LUMO was well correlated with the electron affinity of
metal center. The niobium center shows the greater predisposition than vanadium center to lose its electrons (hard to be reduced). Otherwise, vanadium-center largely contributes and stabilizes the energy level of the LUMO and an additional electron is delocalized over the three vanadium centers in the mono-reduced vanadium-substituted Keggin-type tungstosilicate. The differences in reduction potentials and absorption edge energies could be understood in a similar manner. For Wells-Dawson tungstophosphate, the energy of metal centers at cap sites was stabilized by more reducible vanadium and the electrochemical reduction of vanadium centers at cap sites became available. Therefore, electrochemically added electrons are preferentially localized on the vanadium center. In case of $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$, it could be inferred that niobium center could not stabilize but elevate the energy level of metal centers. Therefore, reduction of metal centers at cap site was not available and showed negatively-shifted reduction potential. The difference in absorption edge energy could be understood by considering the differences in electron affinity of metal center, contribution of metal center to the LUMO, and symmetry of HPA structure (the symmetries of HPA structures are almost identical as a $\alpha_2$-type isomer). It is noteworthy that $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ with the smallest band gap showed the highest reduction potential.
Fig. 3.14. \( [F(R\infty)\cdot h\nu]^{1/2} \) curves of \( \alpha-K_6P_2W_{18}O_{62} \), \( \alpha_2-K_6P_2W_{17}Mo_1O_{62} \), \( \alpha_2-K_7P_2W_{17}V_1O_{62} \), and \( \alpha_2-K_7P_2W_{17}Nb_1O_{62} \) Wells-Dawson HPAs.
3.2.2.3. Scanning tunneling microscopy

Figure 3.15 shows the STM images of $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs deposited on graphite surface. Each bright corrugation represents individual heteropolyanion. STM images clearly showed the formation of self-assembled and well-ordered HPA arrays on HOPG surface. The periodicity and included angle of unit cell constructed on the basis of lattice constants determined from two-dimensional fast Fourier transformation (2D FFT) are also shown in Figure 3.15. The periodicity of $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ unit cell was found to 0.85 Å $\times$ 13.1 Å with included angle of 83.2 °. The periodicity of $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ unit cell was found to 11.1 Å $\times$ 11.5 Å with included angle of 61.0 °. Periodicity of unit cell was well consistent with lattice constants of Wells-Dawson HPAs determined by X-ray crystallography [2].

Tunneling spectroscopy was conducted in order to elucidate the local surface electronic structures of $\alpha$-K$_8$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_8$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. Figure 3.16 shows the typical tunneling spectra of $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs catalyst taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.15. The tunneling spectra taken at bright corrugation showed a distinctive current-voltage behavior referred to as negative differential resistance (NDR), which was known to be a characteristic of HPAs. On the other hands, tunneling spectra taken at interstitial space between bright corrugations showed almost the same current-voltage behavior with that of bare HOPG. This indicates that the self-assembled and well-ordered HPA arrays shown in Figure 3.15 are a monolayer.
The average value of NDR peak voltage was taken as the representative NDR peak voltage. The average values of NDR peak voltage were summarized in Table 3.6. It is interesting that the trend of NDR peak voltage is well consistent with that of UV-visible spectroscopy. As mentioned, it can be inferred that less negative NDR voltage of HPA catalysts corresponds to lower energy state of the LUMO. Because energy gap only depends on energy state of LUMO in the series of $\alpha$-$K_6P_2W_{18}O_{62}$, $\alpha_2$-$K_6P_2W_{17}Mo_1O_{62}$, $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs, smaller absorption edge energy corresponds to lower energy state of the LUMO, leading to less negative NDR peak voltage.

Figure 3.17 shows the absorption edge energy and NDR peak voltage plotted as a function of reduction potential. The correlations suggested that reduction potential could be estimated from the electronic structure and absorption edge energy/NDR peak voltage could be utilized as alternative correlating parameters for reduction potential of bulk HPAs in the series of $\alpha$-$K_6P_2W_{18}O_{62}$, $\alpha_2$-$K_6P_2W_{17}Mo_1O_{62}$, $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs. Among the tested, vanadium center is the most effective to enhance the reducibility of heteropolyanion by the stabilization of energy state of the LUMO.
Fig. 3.15. STM images of (a) $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$ and (b) $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.
Fig. 3.16. Typical tunneling spectra of (a) $\alpha_2-K_7P_2W_{17}V_1O_{62}$ and (b) $\alpha_2-K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs catalyst taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.15.
Table. 3.6.

First electron reduction potentials, absorption edge energies, and NDR peak voltages of $\alpha$-K$_6$P$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$P$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>1$^{st}$ electron reduction potential (volts vs. SCE)</th>
<th>UV-visible absorption edge energy (eV)</th>
<th>NDR peak voltage (volts)</th>
</tr>
</thead>
<tbody>
<tr>
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<td>KP$<em>2$W$</em>{17}$Nb$_1$</td>
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<td>3.06</td>
<td>-1.02</td>
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</table>
Fig. 3.17. Absorption edge energy and NDR peak voltage plotted as a function of reduction potential.
3.2.3. Catalytic test

Gas-phase oxidative dehydrogenation of benzylamine was carried out over $\alpha$-$K_6P_2W_{18}O_{62}$, $\alpha_2$-$K_6P_2W_{17}Mo_1O_{62}$, $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs. The catalysts exhibited a stable catalytic performance during a 5 h-reaction. It was reported that dibenzylimine, benzonitrile, and benzaldehyde could be formed from the dehydrogenated intermediates, phenylmethanimine as shown in Figure 3.18 [45]. Unstable phenylmethanimine reacts with another molecule benzylamine to form aminal and then release an NH$_3$ to form the dibenzylimine. Benzonitrile is formed by further oxidative dehydrogenation of phynylmethanimine. Benzaldehyde is formed by hydrolysis of phenylmethanimine.

In this work, dibenzylimine was mainly formed and trace amounts of benzonitrile and benzaldehyde were detected. It is noteworthy that none of them was detected without oxygen, supporting that dibenzylimine, benzonitrile, benzaldehyde is formed via oxidative dehydrogenation reaction. It was also found that benzylamine readily reacted with benzaldehyde to form dibenzylimine even at room temperature, indicating that as-produced benzaldehyde could react with benzylamine to form dibenzylimine. Figure 3.19 shows the catalytic performance of $\alpha_2$-$K_7P_2W_{17}V_1O_{62}$ during the 5 h-reaction and yield for dibenzylimine after a 5 h-reaction. Interestingly, yield for dibenzylimine increased in the order of $\alpha_2$-$K_7P_2W_{17}Nb_1O_{62} < \alpha$-$K_6P_2W_{18}O_{62} < \alpha_2$-$K_6P_2W_{17}Mo_1O_{62} < \alpha_2$-$K_7P_2W_{17}V_1O_{62}$.

When considering the fact that vanadium center is easier to be reduced
than other metal centers in the electrochemical measurement, it is inferred that vanadium-substitution provided efficient active sites for oxidative dehydrogenation by contributing to the density of accessible electronic states (unoccupied orbitals). This result also suggests that reducibility of mono-transition metal-substituted HPAs plays an important role in the oxidative dehydrogenation of benzylamine.
Fig. 3.18. Plausible reaction mechanism for the oxidative dehydrogenation of benzylamine [45].
Fig. 3.19. (a) Catalytic performance of $\alpha_2$-K$_2$P$_2$W$_{17}$V$_1$O$_{62}$ during the 5 h-reaction and (b) yield for dibenzylimine after a 5 h-reaction.
3.3. **Keggin-type heteropolytungstates with different central atom**

3.3.1. **Formation of HPA structure**

Figure 3.20 shows the DRIFT spectra of $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs. The spectrum of $\alpha$-H$_6$CoW$_{12}$O$_{40}$ catalyst exhibited characteristic bands at 960, 889, and 761 cm$^{-1}$, which were assigned to W=O$_t$, W-O$_e$-W, and W-O$_e$-W stretching vibrations, respectively. The characteristic band for Co-O stretching vibration (445 cm$^{-1}$) was not presented in this figure. The spectrum of $\alpha$-H$_5$BW$_{12}$O$_{40}$ catalyst exhibited characteristic bands at 955, 904, 814, and 753 cm$^{-1}$, which were assigned to W=O$_t$, W-O, W-O$_e$-W, and W-O$_e$-W stretching vibrations, respectively. The characteristic band at 1001 cm$^{-1}$ corresponded to the maximum of W-O-B stretching mode for $\alpha$-H$_5$BW$_{12}$O$_{40}$. The characteristic band for B-O stretching vibration (1410 cm$^{-1}$) was not presented in this figure. The spectrum of $\alpha$-H$_4$SiW$_{12}$O$_{40}$ catalyst exhibited characteristic bands at 979, 924, 885, and 787 cm$^{-1}$, which were assigned to W=O$_t$, Si-O, W-O$_e$-W, and W-O$_e$-W stretching vibrations, respectively. The spectrum of $\alpha$-H$_3$PW$_{12}$O$_{40}$ catalyst exhibited characteristic bands at 1080, 984, 888, and 810 cm$^{-1}$, which were assigned to P-O, W=O$_t$, W-O$_e$-W, and W-O$_e$-W stretching vibrations, respectively. X-O bands of $\alpha$-H$_n$XW$_{12}$O$_{40}$ HPA catalysts did not show any band splitting, indicating the existence of T$_d$ local symmetry of a central XO$_4$ unit that is a characteristic feature of $\alpha$-Keggin structure. All positions of characteristic IR bands of $\alpha$-H$_n$XW$_{12}$O$_{40}$ HPA catalysts were in good agreement with the results of
previous works [1,46,47], indicating successful formation of heteropolyanion frameworks.

Elemental compositions of $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs were calculated from the results of ICP-AES measurements. Chemical compositions of central atom and tungsten in the $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs were summarized in Table 3.7.
Fig. 3.20. DRIFT spectra of $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs
Table. 3.7.

Chemical compositions of central atom (X) and tungsten in the $\alpha$-H$_8$XW$_{12}$O$_{40}$

$(X=\text{Co}^{2+}, \text{B}^{3+}, \text{Si}^{4+}, \text{and P}^{5+})$ Keggin HPAs

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<tr>
<th>Catalyst</th>
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</table>
3.3.2. Characterization

3.3.2.1. Cyclic voltammetry

Figure 3.21 shows the cyclic voltammograms of α-H₄XW₁₂O₄₀ (X=Co²⁺, B³⁺, Si⁴⁺, and P⁵⁺) Keggin HPAs. Electrochemical measurements were performed under consistent experimental conditions to provide the comparable results. All α-H₄XW₁₂O₄₀ HPA catalysts exhibited well-defined reversible and stepwise tungsten-centered redox transitions during the electrochemical measurements. Five couples of one-electron redox waves were observed for α-H₃PW₁₂O₄₀ catalyst. Three couples of redox waves were observed for α-H₄SiW₁₂O₄₀ catalyst with electron ratios of 1:1:2. Two couples of redox waves with electron ratios of 2:1 were observed for α-H₅BW₁₂O₄₀ catalyst. Three couples of one-electron redox waves were observed for α-H₆CoW₁₂O₄₀ catalyst.

First electron reduction potential was increased in the order of α-H₆CoW₁₂O₄₀ < α-H₅BW₁₂O₄₀ < α-H₄SiW₁₂O₄₀ < α-H₃PW₁₂O₄₀ catalysts. First electron reduction potential was summarized in Table 3.8. Among the tested HPA catalysts, α-H₃PW₁₂O₄₀ catalyst showed the most positive reduction potential. It is well known that added electrons to α-type Keggin heteropolytungstates were delocalized over tungsten sites. This indicates that each redox transition during the electrochemical measurements corresponded to electron transfer related to tungsten sites and nature of central atom would not directly affect the reduction potential, but indirectly [48].

Previous density functional theory (DFT) calculations [49,50] has been
reported the effect of heteroatom charge and size on electronic structure using a clathrate model, in which tetrahedral $\text{XO}_4^{n-}$ units are encapsulated within neutral $\text{W}_{12}\text{O}_{40}$ cage ($\text{XO}_4^{n-}@\text{W}_{12}\text{O}_{40}$). It has been demonstrated that Keggin-type heteropolyanion with larger negative charge and smaller size resulted in poor reducibility due to the smaller capacity to accept electrons, in good agreement with the results of this work.
Fig. 3.21. Cyclic voltammograms of $\alpha$-H$_6$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs.
3.3.2.2. UV-visible spectroscopy

Fig. 3.22 shows the \([F(R_\infty)\cdot h\nu]^{1/2}\) curves of \(\alpha\text{-H}_n\text{XW}_{12}\text{O}_{40}\) (X=Co\(^{2+}\), B\(^{3+}\), Si\(^{4+}\), and P\(^{5+}\)) Keggin HPAs. Because absorption edge energy of HPA catalysts are typically sensitive to number of crystalline water molecules, UV-visible spectroscopy measurements were carried out in solution to be fully solvated by water, giving rise to consistent number of crystalline water. Absorption edge energy of \(\alpha\text{-H}_n\text{XW}_{12}\text{O}_{40}\) HPA catalysts determined by UV-visible spectroscopy decreased in the order of \(\alpha\text{-H}_6\text{CoW}_{12}\text{O}_{40} > \alpha\text{-H}_3\text{BW}_{12}\text{O}_{40} > \alpha\text{-H}_4\text{SiW}_{12}\text{O}_{40} > \alpha\text{-H}_3\text{PW}_{12}\text{O}_{40}\) catalysts. Absorption edge energies were summarized in Table 3.8. The trend of absorption edge energy is well consistent with that of reduction potential.

Unlike the case of metal-substitution in the site of addenda atom, the central atom within neutral \(\text{W}_{12}\text{O}_{40}\) cage (\(\text{XO}_n^{\text{=}n}\text{}@\text{W}_{12}\text{O}_{40}\)) significantly affect the both the HOMO and the LUMO [49]. Therefore, it is very hard to figure out the relative energy level of the LUMO. In the series of \(\alpha\text{-H}_n\text{XW}_{12}\text{O}_{40}\) (X=Co\(^{2+}\), B\(^{3+}\), Si\(^{4+}\), and P\(^{5+}\)) Keggin HPAs, the only information estimated from the absorption edge energy was that the heteropolyanion with larger negative charge and smaller size resulted in large energy gap.
Fig. 3.22. $[F(R_{\infty}) \cdot h\nu]^{1/2}$ curves of $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs.
3.3.2.3. Scanning tunneling microscopy

NDR peak voltages of α-HₙXW₁₂O₄₀ (X=Co²⁺, B³⁺, Si⁴⁺, and P⁵⁺) Keggin HPAs determined by tunneling spectroscopy were taken from the previous literature [32]. The average values of NDR peak voltage were summarized in Table 3.8. The trend of NDR peak voltage is well consistent with that of absorption edge energy. As mentioned, it is very hard to figure out the relative energy level of the LUMO from absorption edge energy because the central atom within neutral W₁₂O₄₀ cage (XO₄ⁿ⁻@W₁₂O₄₀) significantly affect the both the HOMO and the LUMO. However, it can be inferred from the trend of NDR peak voltages that energy level of the LUMO decreased in the order of α-H₆CoW₁₂O₄₀ > α-H₃BW₁₂O₄₀ > α-H₄SiW₁₂O₄₀ > α-H₃PW₁₂O₄₀.

Figure 3.23 shows the absorption edge energy and NDR peak voltage plotted as a function of reduction potential. The correlations suggested that reduction potential could be estimated from the electronic structure in the Keggin-type heteropolytungstates with different central atom. Among the tested, P⁵⁺ is the most effective to enhance the reducibility of heteropolyanion framework. The central PO₄³⁻ unit provided electrochemically accessible tungsten centers which is easier to be reduced (than other central XO₄ⁿ⁻ units), resulting in the highest reduction potential. The correlations also suggested that absorption edge energy and NDR peak voltage could be utilized as alternative correlating parameters for reduction potential in the series of α-HₙXW₁₂O₄₀ (X=Co²⁺, B³⁺, Si⁴⁺, and P⁵⁺) Keggin HPAs.
Table 3.8.

First electron reduction potentials, absorption edge energies, and NDR peak voltages of $\alpha$-H$_n$XW$_{12}$O$_{40}$ (X=Co$^{2+}$, B$^{3+}$, Si$^{4+}$, and P$^{5+}$) Keggin HPAs.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>1$^{st}$ electron reduction potential (volts vs. SCE)</th>
<th>UV-visible absorption edge energy (eV)</th>
<th>NDR peak voltage (volts)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PW$_{12}$</td>
<td>-0.020</td>
<td>3.35</td>
<td>-1.14</td>
</tr>
<tr>
<td>SiW$_{12}$</td>
<td>-0.227</td>
<td>3.37</td>
<td>-1.19</td>
</tr>
<tr>
<td>BW$_{12}$</td>
<td>-0.511</td>
<td>3.47</td>
<td>-1.28</td>
</tr>
<tr>
<td>CoW$_{12}$</td>
<td>-0.554</td>
<td>3.48</td>
<td>-1.36</td>
</tr>
</tbody>
</table>

*Data were taken from the previous literature [32].
Fig. 3.23. Absorption edge energy and NDR peak voltage plotted as a function of reduction potential.
3.4. Mo(VI)-substituted Wells-Dawson-type tungstoarsenates

3.4.1. Formation of HPA structure

Figure 3.24 shows the schematic diagram for the preparation procedures of $\alpha$-$\text{K}_6\text{As}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0-3$) Wells-Dawson HPAs. Removal of cap WO$_6$ units in the $\alpha$-$\text{K}_6\text{As}_2\text{W}_{18}\text{O}_{62}$ by pH control yields mono-, di-, and tri-lacunary heteropolyanion. In this work, molybdenum-substituted Wells-Dawson tungstoarsenates were prepared by direct incorporation of molybdenum ions into mono-, di-, and tri-lacunary tungstoarsenates, derived from $\alpha$-$\text{K}_6\text{As}_2\text{W}_{18}\text{O}_{62}$ via selective degradation. Therefore, molybdenum was selectively substituted on the cap sites of Wells-Dawson structure, giving the uniform molecular structure.

Successful formation of heteropolyanion frameworks was confirmed by DRIFT spectroscopy. Figure 3.25 shows the DRIFT spectra of $\alpha$-$\text{K}_6\text{As}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0-3$) Wells-Dawson HPAs. Typical characteristic bands of Wells-Dawson heteropolyanion were observed in the range of 700-1000 cm$^{-1}$. For $\alpha$-$\text{K}_6\text{As}_2\text{W}_{18}\text{O}_{62}$, the characteristic bands attributed to W-O-W asymmetric stretching vibrations were observed at 894, 868, 833, 770 cm$^{-1}$. The band attributed to W=O asymmetric stretching was observed at 974 cm$^{-1}$. The single band attributed to As-O vibration appeared at the wavenumber (974 cm$^{-1}$) near W-O-W bands. Molybdenum-substituted tungstoarsenates ($\alpha$-$\text{K}_6\text{As}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=1-3$)) exhibited a little shifted bands with no significant difference compared to $\text{K}_6\text{As}_2\text{W}_{18}\text{O}_{62}$. Although the local $T_d$ symmetry of central AsO$_4$ units might be
lowered by molybdenum-substitution, the bands attributed to vibration mode of As-O appeared as a single weak shoulder with no band splitting. Therefore, it is inferred that the bands corresponding to central AsO₄ were hidden under the broad and intense M-O-M bands, resulting in a single band for central AsO₄. This result was well consistent with the result of previous work [34], indicating that a series of molybdenum-substituted Wells-Dawson tungstoarsenates were successfully prepared. Successful preparation of the catalysts were further confirmed by ICP-AES analyses. The chemical compositions of arsenic, tungsten, and molybdenum in the \( \alpha-K_6\text{As}_2W_{18-x}\text{Mo}_x\text{O}_{62} \) (x=0-3) Wells-Dawson HPAs determined by ICP-AES measurements were summarized in Table 3.7. Chemical compositions were in good agreement with the designed values.
Fig. 3.24. Schematic diagram for the preparation procedures of $\alpha$-$K_6As_2W_{18-x}Mo_xO_{62}$ ($x=0-3$) Wells-Dawson HPAs.
Fig. 3.25. DRIFT spectra of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) Wells-Dawson HPAs.
### Table 3.9.

Detailed assignment of each characteristic band

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wavenumber (cm&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>As-O</th>
<th>M=O&lt;sub&gt;t&lt;/sub&gt;</th>
<th>M-O-M</th>
</tr>
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<tbody>
<tr>
<td>KAs&lt;sub&gt;2&lt;/sub&gt;W&lt;sub&gt;18&lt;/sub&gt;</td>
<td>974, 928, 894, 868, 833, 770</td>
<td>974</td>
<td>928</td>
<td>894, 868, 833, 770</td>
</tr>
<tr>
<td>KAs&lt;sub&gt;2&lt;/sub&gt;W&lt;sub&gt;17&lt;/sub&gt;Mo&lt;sub&gt;1&lt;/sub&gt;</td>
<td>970, 924, 891, 865, 837, 769</td>
<td>970</td>
<td>924</td>
<td>891, 865, 837, 769</td>
</tr>
<tr>
<td>KAs&lt;sub&gt;2&lt;/sub&gt;W&lt;sub&gt;16&lt;/sub&gt;Mo&lt;sub&gt;2&lt;/sub&gt;</td>
<td>968, 923, 890, 862, 829, 770</td>
<td>968</td>
<td>923</td>
<td>890, 862, 829, 770</td>
</tr>
<tr>
<td>KAs&lt;sub&gt;2&lt;/sub&gt;W&lt;sub&gt;15&lt;/sub&gt;Mo&lt;sub&gt;3&lt;/sub&gt;</td>
<td>968, 925, 893, 861, 831, 772</td>
<td>968</td>
<td>925</td>
<td>893, 861, 831, 772</td>
</tr>
</tbody>
</table>

*O<sub>t</sub>: terminal oxygen
Table 3.10.

Chemical compositions of arsenic, tungsten, and molybdenum in the $\alpha$-$K_{6}As_{2}W_{18-x}Mo_{x}O_{62}$ (x=0-3) Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Ratio of As:W:Mo</th>
<th>Theoretical value</th>
<th>Measured value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$KAs_{2}W_{18}$</td>
<td>2.0 : 18.0 : 0</td>
<td>2.0 : 18.0 : 0</td>
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</tr>
<tr>
<td>$KAs_{2}W_{17}Mo_{1}$</td>
<td>2.0 : 17.0 : 1.0</td>
<td>1.9 : 17.0 : 0.9</td>
<td></td>
</tr>
<tr>
<td>$KAs_{2}W_{16}Mo_{2}$</td>
<td>2.0 : 16.0 : 2.0</td>
<td>1.9 : 16.0 : 1.9</td>
<td></td>
</tr>
<tr>
<td>$KAs_{2}W_{15}Mo_{3}$</td>
<td>2.0 : 15.0 : 3.0</td>
<td>1.9 : 15.0 : 2.9</td>
<td></td>
</tr>
</tbody>
</table>
3.4.2. Characterization

3.4.2.1. Cyclic voltammetry

Electrochemical measurements were carried out to examine the electrochemical redox transitions of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$) Wells-Dawson HPAs. Figure 3.26 shows the cyclic voltammograms of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$) Wells-Dawson HPAs. $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$) showed reversible and stepwise redox transitions during the electrochemical measurements. Overall shapes of redox waves were maintained at the measured scan rates in this work (up to 200 mV/s). For all $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$), the linear relationship between peak current and square root of scan rate was observed, suggesting that these redox transitions were diffusion-controlled process expressed by the Randles-Sevcik equation. $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ exhibited four tungsten-centered redox transitions in this scan range. However, molybdenum-substituted $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=1-3$) exhibited an additional molybdenum-centered redox transition at more positive potential. This indicates that molybdenum center is electrochemically more accessible and added electrons are more easily localized on molybdenum sites than tungsten sites.

First electron reduction potentials determined by electrochemical analysis are summarized in Table 3.11. First electron reduction potential increased with increasing molybdenum content in the series of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0-3$) Wells-Dawson HPAs.
Fig. 3.26. Cyclic voltammograms of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) Wells-Dawson HPAs.
3.4.2.2. UV-visible spectroscopy

Figure 3.27 shows the [(F(R∞)·hv)]^{1/2} curves of α-K₆As₂W₁₈₋ₓMoₓO₆₂ (x=0-3). Absorption edge energies were directly obtained from the linear fit of [(F(R∞)·hv)]^{1/2} curve and summarized in Table 3.11. Absorption edge energy (energy gap between the HOMO and the LUMO) determined by UV-visible spectroscopy decreased with increasing molybdenum content. The trend of absorption edge energy is well consistent with that of reduction potential. Like the case of transition metal-substituted Wells-Dawson tungstophosphates, it could be understood by considering the fact that molybdenum-substitution stabilized the energy level of metal centers at cap sites and reduction of metal centers at cap sites was available. Difference in energy gap of α-K₆As₂W₁₈₋ₓMoₓO₆₂ (x=1-3) can also be understood by considering the contribution of molybdenum to the LUMO. Because the HOMO mostly comprises 2p-orbitals on the framework oxygens, as mentioned, it is nearly unchanged by molybdenum-substitution and the energy gap only depends on the LUMO. This indicates that absorption edge energy can be utilized as an alternative diagnostic to estimate the energy state and reduction potential in the series of molybdenum-substituted Wells-Dawson tungstoarsenates.
Fig. 3.27. $[F(R,\nu)\cdot h\nu]^{1/2}$ curves of $\alpha$-K₆As₂W₁₈₋ₓMoₓO₆₂ (x=0-3) Wells-Dawson HPAs.
3.4.2.3. Scanning tunneling microscopy

Figure 3.28 shows the STM images of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$. Two-dimensional self-assembled HPA array was observed in the STM measurement. Each bright corrugation in the STM images represents individual heteropolyanion. The periodicity of unit cell constructed on the basis of lattice constants determined from two-dimensional fast Fourier transform analysis (2D FFT) was 20.9 Å × 21.9 Å with included angle of 60.2°. The dimension of unit cell determined from STM image was matched with the lattice constant determined by X-ray crystallography [2]. Figure 3.29 shows the typical current-voltage responses of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.28. Current-voltage response at bare graphite was also measured for comparison purpose. It is noteworthy that current-voltage response taken at interstitial space between bright corrugations was almost identical to that of bare graphite, indicating that two-dimensional HPA array shown in Figure 3.28 was monolayer. On the other hand, tunneling spectrum taken at bright corrugation showed a distinctive current-voltage response, referred to as negative differential resistance (NDR) phenomenon. The statistical average value of NDR peak voltage of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ was found to be -1.01 V. The statistical NDR peak voltages of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) are summarized in Table 3.11. It is interesting to note that NDR peak voltage of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) shifted less negative voltage with increasing molybdenum content.

Figure 3.30 shows the absorption edge energy and NDR peak voltage plotted as a function of reduction potential. The correlation suggested that
reduction potential could be estimated from the electronic structure. As a results, molybdenum center is more effective to enhance the reducibility of heteropolyanion framework than tungsten center in the series of $\alpha$-$K_6\text{As}_2W_{18-x}\text{Mo}_xO_{62}$ ($x=0-3$) Wells-Dawson HPAs. The molybdenum-substitution provided electrochemically accessible molybdenum centers which is easier to be reduced, resulting in the higher reduction potential. The correlations suggested that absorption edge energy and NDR peak voltage could be utilized as alternative correlating parameters for reduction potential of bulk HPAs in the series of $\alpha$-$K_6\text{As}_2W_{18-x}\text{Mo}_xO_{62}$ ($x=0-3$) Wells-Dawson HPAs.
Fig. 3.28. STM images of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ Wells-Dawson HPA: (a) 20 x 20 nm and (b) 10 x 10 nm.
Fig. 3.29. Typical tunneling spectra of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ Wells-Dawson HPA taken at two different sites (bright corrugation and interstitial space between bright corrugations) in Figure 3.28.
Table 3.11.
First electron reduction potentials, absorption edge energies, and NDR peak voltages of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ (x=0-3) Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>1st electron reduction potential (volts vs. SCE)</th>
<th>UV-visible absorption edge energy (eV)</th>
<th>NDR peak voltage (volts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KAs$<em>2$W$</em>{18}$</td>
<td>+0.049</td>
<td>2.97</td>
<td>-1.01</td>
</tr>
<tr>
<td>KAs$<em>2$W$</em>{17}$Mo$_1$</td>
<td>+0.243</td>
<td>2.90</td>
<td>-0.99</td>
</tr>
<tr>
<td>KAs$<em>2$W$</em>{16}$Mo$_2$</td>
<td>+0.256</td>
<td>2.84</td>
<td>-0.98</td>
</tr>
<tr>
<td>KAs$<em>2$W$</em>{15}$Mo$_3$</td>
<td>+0.270</td>
<td>2.79</td>
<td>-0.95</td>
</tr>
</tbody>
</table>
Fig. 3.30. Absorption edge energy and NDR peak voltage plotted as a function of reduction potential.
3.4.3. Catalytic test

Gas-phase oxidative dehydrogenation of benzyl alcohol was carried out over $\alpha$-$K_6As_2W_{18-x}Mo_xO_{62}$ ($x=0-3$) Wells-Dawson HPAs. The catalysts exhibited a stable catalytic performance during a 5 h-reaction. Benzaldehyde was mainly formed as an oxidation product, and small amounts of toluene and benzyl ether were also detected [51]. Yield for benzaldehyde (oxidation product) increased with increasing molybdenum content. This supports that HPAs with reducible metal centers shows the excellent catalytic acidity in the oxidative dehydrogenation reaction. Similar results have been reported in the oxidative dehydrogenation of 2-propanol [20]. They showed that the rate-determining step involved reduction of HPA with $\beta$-hydrogen eliminated from the substrate and the intrinsic reaction rate was well correlated with reducibility of HPA.

When considering the fact that molybdenum center is easier to be reduced than other tungsten centers in the electrochemical measurement, it is inferred that molybdenum-substitution provided efficient active sites for oxidative dehydrogenation by contributing the accessible electronic state (unoccupied orbital). This result suggests that reducibility of the metal centers in the HPA framework plays an important role in the oxidative dehydrogenation reaction.
Fig. 3.31. (a) Catalytic performance of $\alpha$-$K_6As_2W_{18}O_{62}$ during a 5 h-reaction and (b) yield for acetaldehyde after a 5 h-reaction plotted as a function of molybdenum content.
3.5. V(V) and Nb(V)-substituted Wells-Dawson-type tungstoarsenates

3.5.1. Formation of HPA structure

α₂-K₇As₂W₁₇V₁O₆₂ and α₂-K₇As₂W₁₇Nb₁O₆₂ were prepared to elucidate the effect of group 5-metal-substitution on the redox properties and electronic structure in the Wells-Dawson-type tungstoarsenate frameworks. α-K₆As₂W₁₈O₆₂ and α₂-K₆As₂W₁₇Mo₁O₆₂ were also prepared for the comparison. Figure 3.32 shows the schematic diagram for the preparation procedures of α-K₆As₂W₁₈O₆₂, α₂-K₆As₂W₁₇Mo₁O₆₂, α₂-K₇As₂W₁₇V₁O₆₂, and α₂-K₇As₂W₁₇Nb₁O₆₂ Wells-Dawson HPAs. Mono-transition metal-substituted Wells-Dawson tungstoarsenates were prepared by direct incorporation of transition metal into the mono-lacunary α₂-K₁₀As₂W₁₇O₆₁₀. As a results, a tungsten center at cap site was selectively replaced by a transition metal.

Figure 3.33 shows the DRIFT spectra of α-K₆As₂W₁₈O₆₂, α₂-K₆As₂W₁₇Mo₁O₆₂, α₂-K₇As₂W₁₇V₁O₆₂, and α₂-K₇As₂W₁₇Nb₁O₆₂ Wells-Dawson HPAs. Successful formations of the heteropolyanions were confirmed by four characteristic bands in the range of 1000-700 cm⁻¹, which represents As-O, M=O, inter-octahedral M-Oₖ-M, and intra-octahedral M-Oₖ-M bonds (M=metal). Detailed assignment of each characteristic band was summarized in Table 3.12. Unfortunately, α-K₆As₂W₁₈O₆₂, α₂-K₆As₂W₁₇Mo₁O₆₂, α₂-K₇As₂W₁₇V₁O₆₂, and α₂-K₇As₂W₁₇Nb₁O₆₂ Wells-Dawson HPAs showed no peaks in ⁷⁵As NMR analyses.
Chemical compositions of phosphorous, tungsten, and substituted-metal of the HPAs were further confirmed by ICP-AES analyses. Chemical compositions of arsenic, tungsten, and substituted-metal in the $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs were summarized in table 3.13. The molar ratios of P:W:M were in good agreement with the theoretical values, suggesting the successful formation of HPA structures.
Fig. 3.32. Schematic diagram for the preparation procedures of $\alpha$-$K_6As_2W_{18}O_{62}$, $\alpha_2$-$K_6As_2W_{17}Mo_1O_{62}$, $\alpha_2$-$K_7As_2W_{17}V_1O_{62}$, and $\alpha_2$-$K_7As_2W_{17}Nb_1O_{62}$ Wells-Dawson HPAs.
Fig. 3.33. DRIFT spectra of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.
Table 3.12.

Detailed assignment of each characteristic band

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wavenumber (cm(^{-1}))</th>
<th>As-O</th>
<th>M=O(_t)</th>
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<td>KAs(<em>2) W(</em>{18})</td>
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<td>974</td>
<td>928</td>
<td>894, 868, 833, 770</td>
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<tr>
<td>KAs(<em>2) W(</em>{17})M(_{1})</td>
<td></td>
<td>970</td>
<td>924</td>
<td>891, 865, 837, 769</td>
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<tr>
<td>KAs(<em>2) W(</em>{17})V(_{1})</td>
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<td>968</td>
<td>-</td>
<td>892, 865, 830, 775</td>
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<tr>
<td>KAs(<em>2) W(</em>{17})Nb(_{1})</td>
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<td>968</td>
<td>926</td>
<td>899, 864, 825, 776</td>
</tr>
</tbody>
</table>

\(\*O_t\): terminal oxygen
Table 3.13.
Chemical compositions of arsenic, tungsten, and substituted-metal (M) in the $\alpha$-$K_6\text{As}_2\text{W}_{18}\text{O}_{62}$, $\alpha_2$-$K_6\text{As}_2\text{W}_{17}\text{Mo}_1\text{O}_{62}$, $\alpha_2$-$K_7\text{As}_2\text{W}_{17}\text{V}_1\text{O}_{62}$, and $\alpha_2$-$K_7\text{As}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$ Wells-Dawson HPAs

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Ratio of As:W:M</th>
<th>Theoretical value</th>
<th>Measured value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K\text{As}<em>2\text{W}</em>{18}$</td>
<td>2.0 : 18.0 : 0</td>
<td>2.0 : 18.0 : 0</td>
<td></td>
</tr>
<tr>
<td>$K\text{As}<em>2\text{W}</em>{17}\text{Mo}_1$</td>
<td>2.0 : 17.0 : 1.0</td>
<td>1.9 : 17.0 : 0.9</td>
<td></td>
</tr>
<tr>
<td>$K\text{As}<em>2\text{W}</em>{17}\text{V}_1$</td>
<td>2.0 : 17.0 : 1.0</td>
<td>2.0 : 17.0 : 1.0</td>
<td></td>
</tr>
<tr>
<td>$K\text{As}<em>2\text{W}</em>{17}\text{Nb}_1$</td>
<td>2.0 : 17.0 : 1.0</td>
<td>2.0 : 17.0 : 1.0</td>
<td></td>
</tr>
</tbody>
</table>
3.5.2. Characterization

3.5.2.1. Cyclic voltammetry

Figure 3.34 shows the cyclic voltammograms of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$PAs$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. Four tungsten-based redox transitions were observed in the cyclic voltammogram of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$. In case of molybdenum- and vanadium-substituted Wells-Dawson tungstoarsenates showed an additional molybdenum- and vanadium-based redox transitions, respectively. The results are very similar with the case of Wells-Dawson tungstophosphates. However, niobium-substituted Wells-Dawson tungstoarsenates showed significantly shifted redox couples. This indicates vanadium or molybdenum center in the heteropolyanion framework was more electrochemically accessible and reducible than tungsten center in the Wells-Dawson tungstoarsenate frameworks. Otherwise, substituted-niobium has negative effect to the reducibility of Wells-Dawson tungstoarsenates. First electron reduction potentials are summarized in Table 3.14. Like, first electron reduction potential increased in the order of $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ < $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ < $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$ < $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$. 
Fig. 3.34. Cyclic voltammograms of \( \alpha-K_6As_2W_{18}O_{62} \), \( \alpha_2-K_6As_2W_{17}Mo_1O_{62} \), \( \alpha_2-K_7As_2W_{17}V_1O_{62} \), and \( \alpha_2-K_7As_2W_{17}Nb_1O_{62} \) Wells-Dawson HPAs.
3.5.2.2. UV-visible spectroscopy

UV-visible spectroscopy measurements were conducted to probe the bulk electronic structures. Figure 3.35 shows the \[\sqrt{F(R_\infty) \cdot h\nu}\] curves of \(\alpha\)-K\(_6\)As\(_2\)W\(_{18}\)O\(_{62}\), \(\alpha_2\)-K\(_6\)As\(_2\)W\(_{17}\)Mo\(_1\)O\(_{62}\), \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)V\(_1\)O\(_{62}\), and \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)Nb\(_1\)O\(_{62}\) Wells-Dawson HPAs. Absorption edge energies are summarized in Table 3.14. Absorption edge energy (energy gap between the HOMO and the LUMO) decreased in the order of \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)Nb\(_1\)O\(_{62}\) > \(\alpha\)-K\(_6\)As\(_2\)W\(_{18}\)O\(_{62}\) > \(\alpha_2\)-K\(_6\)As\(_2\)W\(_{17}\)Mo\(_1\)O\(_{62}\) > \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)V\(_1\)O\(_{62}\). Because the HOMO is little perturbed by metal-substitution, it could be inferred that \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)V\(_1\)O\(_{62}\) with the smallest absorption edge energy has the lowest energy level of the LUMO.

The difference in absorption edge energy could be understood by considering the differences in electron affinity of metal center, contribution of metal center to the LUMO, and symmetry of HPA structures. As mentioned above, niobium centers shows the greater predisposition than vanadium center to lose its electrons (hard to be reduced). Otherwise, vanadium-center largely contributes/stabilizes the LUMO. Similar with the case of Wells-Dawson tungstophosphates, the energy of metal centers at cap sites were stabilized by vanadium center and the electrochemical reduction of vanadium centers at cap sites became available. It is inferred that added electrons are preferentially delocalized over the vanadium centers. In case of \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)Nb\(_1\)O\(_{62}\), it could be inferred that niobium center could not stabilize but elevate the energy level of metal centers. Therefore, reduction of metal centers at cap site was not available and showed negatively-shifted reduction potential. Among the tested, \(\alpha_2\)-
K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ with the smallest band gap showed the highest reduction potential.

Figure 3.36 shows the absorption edge energy plotted as a function of reduction potential. Reduction potential of the catalysts increased with decreasing absorption edge energy. This suggests that the electronic properties plays an important role to determine the reducibility of HPAs and absorption edge energy can be utilized as an alternative diagnostic to estimate the electronic structure and reduction potential in the series of $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$, $\alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs.
Fig. 3.35. \([F(R_\infty)\cdot h\nu]^{1/2}\) curves of \(\alpha\)-K\(_6\)As\(_2\)W\(_{18}\)O\(_{62}\), \(\alpha_2\)-K\(_6\)As\(_2\)W\(_{17}\)Mo\(_1\)O\(_{62}\), \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)V\(_1\)O\(_{62}\), and \(\alpha_2\)-K\(_7\)As\(_2\)W\(_{17}\)Nb\(_1\)O\(_{62}\) Wells-Dawson HPAs.
Table 3.14.
First electron reduction potentials and absorption edge energies of α-K\textsubscript{6}As\textsubscript{2}W\textsubscript{18}O\textsubscript{62}, α\textsubscript{2}-K\textsubscript{6}As\textsubscript{2}W\textsubscript{17}Mo\textsubscript{1}O\textsubscript{62}, α\textsubscript{2}-K\textsubscript{7}As\textsubscript{2}W\textsubscript{17}V\textsubscript{1}O\textsubscript{62}, and α\textsubscript{2}-K\textsubscript{7}As\textsubscript{2}W\textsubscript{17}Nb\textsubscript{1}O\textsubscript{62} Wells-Dawson HPAs.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>1st electron reduction potential (volts vs. SCE)</th>
<th>UV-visible absorption edge energy (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KAs\textsubscript{2}W\textsubscript{18}</td>
<td>+0.049</td>
<td>2.97</td>
</tr>
<tr>
<td>KAs\textsubscript{2}W\textsubscript{17}Mo\textsubscript{1}</td>
<td>+0.243</td>
<td>2.90</td>
</tr>
<tr>
<td>KAs\textsubscript{2}W\textsubscript{17}V\textsubscript{1}</td>
<td>+0.378</td>
<td>2.36</td>
</tr>
<tr>
<td>KAs\textsubscript{2}W\textsubscript{17}Nb\textsubscript{1}</td>
<td>-0.162</td>
<td>3.01</td>
</tr>
</tbody>
</table>
Fig. 3.36. Absorption edge energy plotted as a function of reduction potential.
3.5.3. Catalytic test

Gas-phase oxidative dehydrogenation of benzylamine was carried out over $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$, $\alpha$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62}$, $\alpha$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$, and $\alpha$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs. The catalysts exhibited a stable catalytic performance during a 5 h-reaction. In this work, dibenzylimine was mainly formed and trace amounts of benzonitrile and benzaldehyde were detected. None of them was detected without oxygen, indicating that dibenzylimine, benzonitrile, benzaldehyde is formed via oxidative dehydrogenation reaction. Figure 3.37 shows the catalytic performance of $\alpha$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ during the 5 h-reaction and yield for dibenzylimine after a 5 h-reaction. Yield for dibenzylimine increased in the order of $\alpha$-K$_7$P$_2$W$_{17}$Nb$_1$O$_{62} < \alpha$-K$_6$P$_2$W$_{18}$O$_{62} < K_6P_2W_{17}Mo_1O_{62} < \alpha$-K$_7$P$_2$W$_{17}$V$_1$O$_{62}$.

When considering the fact that vanadium center is easier to be reduced than other metal centers in the electrochemical measurement, it is inferred that vanadium-substitution provided efficient active sites for oxidative dehydrogenation by contributing to the density of accessible electronic states (unoccupied orbitals) in the Wells-Dawson tungstoarsenate frameworks. This result suggests that reducibility of the metal centers in the Wells-Dawson tungstoarsenate frameworks plays an important role in the oxidative dehydrogenation reaction of benzylamine. Interestingly, metal-substitution (as addenda atom) showed the similar effect on the redox properties and the catalytic activity, even in a different series of HPAs.
Fig. 3.37. (a) Catalytic performance of $\alpha_2$-$K_7\text{As}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ during the 5 h-reaction and (b) yield for dibenzylimine after a 5 h-reaction.
Chapter 4. Conclusions

In this work, several series of transition metal-substituted HPA catalysts with different addenda atoms, central atoms, contents, and structures were designed and synthesized in order to elucidate the catalytic redox properties and catalytic activities in oxidation catalysis. They were investigated by several experimental techniques including electrochemical analysis, UV-visible spectroscopy, and scanning tunneling microscopy (STM) in order to elucidate the relationship between electronic structure and redox properties. Furthermore, the reliabilities of absorption edge energy and negative differential resistance (NDR) peak voltage determined by UV-visible spectroscopy and scanning tunneling microscopy, respectively, were also examined as alternative parameters for the redox property.

In molybdenum-substituted $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs, molybdenum-substituted Wells-Dawson HPAs showed an additional molybdenum-based redox transition at more positive potential. First electron reduction potentials increase with increasing molybdenum substitution. Absorption edge energy determined from the linear fit of $[F(R_{\infty}) \cdot h\nu]^{1/2}$ decreased with increasing molybdenum content. In STM measurements, two-dimensional self-assembled HPA arrays were observed. Tunneling spectra taken at bright corrugations showed a distinctive current-voltage responses, referred to as negative differential resistance (NDR) phenomenon. NDR peak voltage appeared at less negative voltage with increasing the molybdenum content. In case of Wells-Dawson tungstophosphate, added electrons are generally delocalized over
twelve tungsten centers at equatorial sites. In case of molybdenum-substituted heteropolyanion, however, the energy of metal centers at cap sites is stabilized by more reducible molybdenum and the electrochemical reduction of molybdenum centers at cap sites would be available. Among the investigated, $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ showed the highest reduction potential and the smallest energy gap. It is inferred that the most stabilized energy level of the LUMO in the $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$, due to the contribution of reducible molybdenum metal centers, provided the electrochemically accessible metal centers and resulted in the highest reduction potential. Gas-phase oxidative dehydrogenation of ethanol to acetaldehyde was carried out as a model reaction to probe oxidation catalysis of $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) Wells-Dawson HPAs. Yield for acetaldehyde (oxidation product) increased with increasing molybdenum content. Among the tested catalysts, $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ with the highest reduction potential showed the best catalytic performance. Molybdenum-substitution provided efficient active sites for oxidative dehydrogenation by contributing to the density of accessible electronic states (unoccupied orbitals). The results suggests that reducibility of the metal centers in the Wells-Dawson tungstophosphate frameworks plays an important role in the oxidative dehydrogenation of ethanol.

Group 5 metal (V and Nb)-substituted Wells-Dawson tungstophosphates were synthesized via direct incorporation of transition metal into the monolacunary species to yield the selectively-substituted structures. In the electrochemical analysis, vanadium-substituted Wells-Dawson tungstophosphates showed an additional vanadium-based redox transitions at more positive potential. However, niobium-substituted Wells-Dawson tungstophosphates showed significantly shifted redox transitions. First electron
reduction potential increased in the order of \( \alpha_2-K_7P_2W_{17}Nb_1O_{62} < \alpha-K_6P_2W_{18}O_{62} < \alpha_2-K_6P_2W_{17}Mo_1O_{62} < \alpha_2-K_2P_2W_{17}V_1O_{62} \). Absorption edge energy determined by UV-visible spectroscopy decreased in the order of \( \alpha_2-K_7P_2W_{17}Nb_1O_{62} > \alpha-K_6P_2W_{18}O_{62} > \alpha_2-K_6P_2W_{17}Mo_1O_{62} > \alpha_2-K_2P_2W_{17}V_1O_{62} \). The trend of NDR peak voltage was also well consistent with that of absorption edge energy. It is inferred that the energy level of metal centers at cap sites was stabilized by more reducible vanadium which largely contributes and stabilizes the LUMO and the electrochemical reduction of vanadium centers at cap sites would be available. Therefore, electrochemically added electrons are preferentially localized on the vanadium center. Otherwise, niobium center shows the greater predisposition than vanadium center to lose its electrons (hard to be reduced). In case of \( \alpha_2-K_7P_2W_{17}Nb_1O_{62} \), niobium center could not stabilize but elevate the energy level of metal centers. Gas-phase oxidative dehydrogenation of benzylamine was carried out as a model reaction to probe oxidation catalysis. Yield for dibenzylimine (oxidation product) increased in the order of \( \alpha_2-K_7P_2W_{17}Nb_1O_{62} < \alpha-K_6P_2W_{18}O_{62} < \alpha_2-K_6P_2W_{17}Mo_1O_{62} < \alpha_2-K_2P_2W_{17}V_1O_{62} \).

Heteropolytungstates with different central atom, \( \alpha-H_nXW_{12}O_{40} \) (\( X=\text{Co}^{2+}, \text{B}^{3+}, \text{Si}^{4+}, \text{and} \text{P}^{5+} \)) Keggin HPAs were prepared to elucidate the effect of central atom on the redox properties and catalytic activity. All \( \alpha-H_nXW_{12}O_{40} \) HPA catalysts exhibited well-defined reversible and stepwise tungsten-centered redox transitions during the electrochemical measurements. First electron reduction potential increased in the order of \( \alpha-H_6\text{CoW}_{12}O_{40} < \alpha-H_5\text{BW}_{12}O_{40} < \alpha-H_4\text{SiW}_{12}O_{40} < \alpha-H_3\text{PW}_{12}O_{40} \) catalysts. Absorption edge energy determined by UV-visible spectroscopy decreased in the order of \( \alpha-H_6\text{CoW}_{12}O_{40} > \alpha-H_5\text{BW}_{12}O_{40} > \alpha-H_4\text{SiW}_{12}O_{40} > \alpha-H_3\text{PW}_{12}O_{40} \) catalysts. The trend of NDR peak
Voltage was also well consistent with that of absorption edge energy. Keggin-type heteropolyanion with larger negative charge and smaller size resulted in poor reducibility of heteropolyanion due to the smaller capacity to accept electrons. Among the tested, PO$_4^{3-}$ anion with smaller negative charge and larger size is the most effective to enhance the reducibility.

Another heteropolytungstates containing AsO$_4^{3-}$ as a central unit were also examined. The series of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0$-3) Wells-Dawson HPAs were prepared via direct incorporation of transition metal into the mono-, di-, and tri-lacunary species. In electrochemical analysis, $\alpha$-K$_6$As$_2$W$_{18}$O$_{62}$ exhibited four tungsten-centered redox transitions. However, molybdenum-substituted $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=1$-3) exhibited an additional molybdenum-centered redox transition at more positive potential. First electron reduction potential increased with increasing molybdenum content. Absorption edge energy decreased with increasing molybdenum content. NDR peak voltage appeared at less negative voltage with increasing molybdenum content. These results are well consistent with the results in the series of $\alpha$-K$_6$As$_2$W$_{18-x}$Mo$_x$O$_{62}$ ($x=0$, 3, 9, 15, 18). Gas-phase oxidative dehydrogenation of benzyl alcohol was carried out as a model reaction to track the oxidation catalysis. Yield for benzaldehyde (oxidation product) increased with increasing molybdenum content.

Furthermore, group 5 metal-substituted $\alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ and $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62}$ Wells-Dawson HPAs were examined. In the electrochemical analysis, vanadium-substituted Wells-Dawson tungstoarsenates showed additional vanadium-centered redox transitions at more positive potential. However, niobium-substituted Wells-Dawson tungstoarsenates showed significantly shifted redox transitions. First electron reduction potential increased
in the order of $\alpha_2$-$K_7$As$_2$W$_{17}$Nb$_1$O$_{62} < \alpha$-$K_6$As$_2$W$_{18}$O$_{62} < \alpha_2$-$K_6$As$_2$W$_{17}$Mo$_1$O$_{62} < \alpha_2$-$K_7$As$_2$W$_{17}$V$_1$O$_{62}$. Absorption edge energy determined by UV-visible spectroscopy decreased in the order of $\alpha_2$-$K_7$As$_2$W$_{17}$Nb$_1$O$_{62} > \alpha$-$K_6$As$_2$W$_{18}$O$_{62} > \alpha_2$-$K_6$As$_2$W$_{17}$Mo$_1$O$_{62} > \alpha_2$-$K_7$As$_2$W$_{17}$V$_1$O$_{62}$. Gas-phase oxidative dehydrogenation of benzylamine was carried out as a model reaction to track the oxidation catalyst. Yield for dibenzyylimine (oxidation product) increased in the order of $\alpha_2$-$K_7$As$_2$W$_{17}$Nb$_1$O$_{62} < \alpha$-$K_6$As$_2$W$_{18}$O$_{62} < \alpha_2$-$K_6$As$_2$W$_{17}$Mo$_1$O$_{62} < \alpha_2$-$K_7$As$_2$W$_{17}$V$_1$O$_{62}$.

From the results, it was found that redox properties of HPAs could be altered by changing the constituent elements, compositions, or structures. It could be concluded that redox properties of HPAs, which are closely related to the electronic structure, play an important role to determine the catalytic activities in the oxidative dehydrogenation reactions. Furthermore, it was successfully demonstrated that absorption edge energy and NDR peak voltage can be utilized as alternative parameters to estimate the reducibility of HPAs.
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“Rationalization and improvement of the syntheses of two octadecatungstoarsenates: the novel $\alpha$-K$_7$[H$_4$AsW$_{18}$O$_{62}$]$\cdot$18H$_2$O and the well known symmetrical $\alpha$-K$_6$[As$_2$W$_{18}$O$_{62}$]$\cdot$14H$_2$O”, *Inorg. Chim. Acta* 342 (2003) 219.


요 약(국문초록)

헤테로폴리산은 다양한 구조와 조성을 지닌 금속산화물 클러스터이며 고유의 산화환원 특성과 구조 안정성으로 인해 산화반응 및 광촉매적 환원 반응에서 촉매로 널리 이용되고 있다. 헤테로폴리산을 이용한 산화반응은 지속적으로 연구되어 왔으며, 최근에는 배위원소가 치환된 (Mixed-addenda) 헤테로폴리산이 산화촉매로서 주목받고 있다. 헤테로폴리산의 촉매적 특성과 촉매활성은 촉매의 구성원소를 달리함으로써 분자 수준에서 조절할 수 있는 것으로 알려져 있다. 금속, 반금속 또는 비금속 원소들이 구성원소로서 헤테로폴리산 구조 내에 포함될 수 있으며 이에 따라 이론적으로 조합 가능한 헤테로폴리산은 무수히 많다. 이는 헤테로폴리산의 물성분포가 매우 광범위한 특성을 의미하며, 요구에 따라 다양한 물성을 지닌 헤테로폴리산을 설계할 수 있음을 나타낸다.

본 연구에서는 다양한 전이금속이 치환된 헤테로폴리산을 제조하여 조성과 구조가 산화환원 특성 및 산화반응에서의 촉매활성에 미치는 영향을 확인하고자 하였다. 이를 위해 다양한 구조의 헤테로폴리산 촉매를 제조하고 전기화학적 분석, 자외선-가시광선 분광법 및 주사 터널링 현미경을 이용해 분석하였다. 특히, 자외선-가시광선 분광법으로 측정되는 흡수단 에너지 (Absorption edge energy) 및 주사 터널링 현미경으로 측정되는 NDR (Negative differential
전압과 산화환원 특성 간의 상관관계를 분석하였다. 먼저, 서로 다른 조성으로 물리브테니아 치환된 Wells-Dawson형 $\text{H}_6\text{P}_2\text{W}_{18-x}\text{Mo}_x\text{O}_{62}$ ($x=0, 3, 9, 15, 18$) 헤테로폴리산을 “Etherate 제조법”으로 제조하고 물리브테니 치환 Wells-Dawson 형 탕스토인산의 산화환원 특성과 촉매활성에 미치는 영향을 확인하고자 하였다. 순환전압전류법을 통해 $\text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ 헤테로폴리산 촉매가 탕스텐 기반의 산화환원쌍을 보임을 확인하였다. 반면 $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ 헤테로폴리산 촉매는 물리브테니 기반의 산화환원쌍을 나타내었다. 황미롭게도 물리브테니가 일부 치환된 다른 촉매들에서는 $\text{H}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ 헤테로폴리산과 비교해 물리브테니 기반의 산화환원쌍이 추가적으로 관찰되었다. 환원전위는 헤테로폴리산 내 물리브테니 함량이 증가함에 따라 증가하였다. 헤테로폴리산의 전자구조를 확인하기 위해 자외선-가시광선 분광법을 수행하였으며 이로부터 측정된 흡수단 에너지는 물리브테니 함량이 증가함에 따라 감소하였다. 추가적인 표면의 국부적 전자구조 분석을 위해 주사 터널링 현미경을 이용하였다. 주사 터널링 현미경을 통해 얻어진 이미지를 통해 단순한 물리적 증착을 통해서도 헤테로폴리산이 그래파이트 상에서 정렬된 2차원 구조를 형성함을 확인하였다. 이미지의 밝은 부분에서 측정한 투과 스펙트럼에서는 NDR 현상이 관찰되었으며, NDR 전압은 물리브테니 함량이 증가함에 따라 증가하였다. 최종적으로 산화반응에서의 촉매활성을 확인하기 위해 에탄올의 산화적 탈수소화 반응을 수행하였다. 아세트알데히드 수율은 물리브테니 함량이
증가함에 따라 증가하였으며 모든 촉매 중에서 가장 높은 환원전위값을 보인 $\text{H}_6\text{P}_2\text{Mo}_{18}\text{O}_{62}$ 헤테로폴리산 촉매가 가장 높은 알세트알데히드 수율을 나타내었다.

추가적으로 5족 금속이 치환된 Wells-Dawson형 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ 및 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$ 헤테로폴리산을 제조하고 5족 금속 치환이 Wells-Dawson형 팀스토인산의 산화환원 특성과 촉매활성에 미치는 영향을 확인하고자 하였다. 해당 촉매는 금속의 선택적 치환을 위해 Lacunary 화합물을 이용하여 제조하였다. $\alpha\text{-K}_6\text{P}_2\text{W}_{18}\text{O}_{62}$ 및 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62}$ 헤테로폴리산 또한 대조군으로 제조하였다. 순환전압전류법을 통해 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ 및 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62}$ 헤테로폴리산에서는 $\alpha\text{-K}_6\text{P}_2\text{W}_{18}\text{O}_6$ 헤테로폴리산과 비교해 추가적인 바나듐 및 물리브테데 기반의 산화환원쌍이 나타남을 확인하였다. 반면, $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62}$ 에서는 추가적인 산화환원쌍이 나타나지 않고 팀스턴 기반의 산화환원쌍이 음전압폭으로 크게 이동함을 확인하였다. 전기화학적 분석을 통해 측정된 환원전위는 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62} < \alpha\text{-K}_6\text{P}_2\text{W}_{18}\text{O}_{62} < \alpha_2\text{-K}_8\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62} < \alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ 순으로 증가하였다. 자외선-가시광선 분광법으로 측정된 흡수단 에너지는 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62} > \alpha\text{-K}_6\text{P}_2\text{W}_{18}\text{O}_{62} > \alpha_2\text{-K}_8\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62} > \alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ 순으로 감소하였다. 주사 터널링 현미경을 통해 얻어진 이미지를 통해 해당 촉매들이 그래파이드 표면에서 형성된 2차원 구조를 형성함을 확인하였다. 측정된 NDR 전압은 $\alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{Nb}_1\text{O}_{62} < \alpha\text{-K}_6\text{P}_2\text{W}_{18}\text{O}_{62} < \alpha_2\text{-K}_8\text{P}_2\text{W}_{17}\text{Mo}_1\text{O}_{62} < \alpha_2\text{-K}_7\text{P}_2\text{W}_{17}\text{V}_1\text{O}_{62}$ 순서로 증가하였다.
해당 촉매의 산화반응에서의 촉매활성을 확인하기 위해 벤질아민의 산화적 탈수소화 반응을 수행하였으며, 디벤질아민의 수율은 \( \alpha_2K_7P_2W_{17}Nb_{1}O_{62} < \alpha_K_6P_2W_{18}O_{62} < \alpha_2K_6P_2W_{17}Mo_{1}O_{62} < \alpha_2K_7P_2W_{17}V_{1}O_{62} \) 순서로 증가하였다.

서로 다른 중심원소가 산화환원 특성에 미치는 영향을 확인하기 위해 Keggin형 \( \alpha-H_nXW_{12}O_{40} \) \( (X=Co^{2+}, B^{3+}, Si^{4+}, \text{and} P^{5+}) \) 헤테로폴리산을 제조하였다. 모든 촉매의 순환전류전압 곡선에서 가역적·순차적인 텅스텐 기반의 산화환원쌍이 관찰되었다. 환원전위는 \( \alpha-H_6CoW_{12}O_{40} < \alpha-H_5BW_{12}O_{40} < \alpha-H_4SiW_{12}O_{40} < \alpha-H_3PW_{12}O_{40} \) 순으로 증가하였다. 외전-가시광선 분광법으로 측정된 흡수단 에너지는 \( \alpha-H_6CoW_{12}O_{40} > \alpha-H_5BW_{12}O_{40} > \alpha-H_4SiW_{12}O_{40} > \alpha-H_3PW_{12}O_{40} \) 순으로 감소 하였다. 이러한 경향은 주사 터널링 현미경 분석을 통해 얻어진 NDR 전압에서도 유사하게 관찰되었다. 상대적으로 큰 사이즈와 높은 전하를 지닌 \( PO_4^{3-} \) 음이온이 촉매의 환원력을 높이는데 효과적인 것으로 나타났다.

또 다른 3가의 \( AsO_4^{3-} \) 음이온이 중심 금속으로 포함된 Wells-Dawson형 텅스토비소산에서 전이금속의 치환이 산화환원 특성과 촉매활성에 미치는 영향을 확인하고자 하였다. 이를 위해 서로 다른 조성으로 몰리브데넘이 치환된 Wells-Dawson형 \( \alpha-K_6As_2W_{18-x}Mo_{x}O_{62} \) \( (x=0-3) \) 헤테로폴리산을 제조하였다. 순환전류전압 곡선에서 \( \alpha-K_6As_2W_{18}O_{62} \)가 텅스텐 기반의 산화환원쌍을 나타내는 반면, 몰리브데넘이 치환된 Wells-Dawson형 텅스토비소산에서는 추가적인 몰리브데넘 기반의 산화환원쌍이 관찰되었다. Wells-Dawson형
텅스토인산의 결과와 매우 유사하게, 환원전위는 헤테로폴리산 내 물리브테넘 함량이 증가함에 따라 증가하였다. 자외선-가시광선 분광법으로 측정된 홍수단 에너지는 물리브테넘 함량이 증가함에 따라 감소하였다. NDR 전압은 물리브테넘 함량이 증가함에 따라 증가하였다. 해당 촉매의 산화반응에서의 촉매활성을 확인하기 위해 벤질알코올의 산화적 탈수소화 반응을 수행하였으며, 벤زال데히드 수용은 물리브테넘 함량이 증가함에 따라 증가하였다.

마지막으로 5족 금속인 Wells-Dawson형 텅스토비소산의 산화환원 특성과 촉매활성에 미치는 영향을 확인하고자 \( \alpha_2-K_7As_2W_{17}V_1O_{62} \) 및 \( \alpha_2-K_7As_2W_{17}Nb_1O_{62} \) 헤테로폴리산을 제조하였다. \( \alpha-K_6As_2W_{18}O_{62} \) 및 \( \alpha_2-K_6As_2W_{17}Mo_1O_{62} \) 헤테로폴리산 또한 대조군으로 제조하였다. 순환전압전류법을 통해 \( \alpha_2-K_7As_2W_{17}V_1O_{62} \) 및 \( \alpha_2-K_7As_2W_{17}Mo_1O_{62} \) 헤테로폴리산에서는 \( \alpha-K_6As_2W_{18}O_{62} \) 헤테로폴리산과 비교해 추가적인 바나듐 및 물리브테넘 기반의 산화환원쌍이 나타남을 확인하였다. 반면, \( \alpha_2-K_7As_2W_{17}Nb_1O_{62} \)에서는 추가적인 산화환원쌍이 관찰되지 않고 텅스텔 기반의 산화환원쌍이 음전압 영역으로 크게 이동함을 확인하였다. 환원전위는 \( \alpha_2-K_7As_2W_{17}Nb_1O_{62} < \alpha-K_6As_2W_{18}O_{62} < \alpha_2-K_6As_2W_{17}Mo_1O_{62} < \alpha_2-K_7As_2W_{17}V_1O_{62} \) 순으로 증가하였다. 자외선-가시광선 분광법으로 측정된 홍수단 에너지는 \( \alpha_2-K_7As_2W_{17}Nb_1O_{62} > \alpha-K_6As_2W_{18}O_{62} > \alpha_2-K_6As_2W_{17}Mo_1O_{62} > \alpha_2-K_7As_2W_{17}V_1O_{62} \) 순으로 감소하였다. 해당 촉매의 산화반응에서의 촉매활성을 확인하기 위해 벤질아민의 산화적 탈수소화 반응을 수행하였으며,
디벤질이민의 수율은 $\alpha_2$-K$_7$As$_2$W$_{17}$Nb$_1$O$_{62} < \alpha$-K$_6$As$_2$W$_{18}$O$_{62} < \alpha_2$-K$_6$As$_2$W$_{17}$Mo$_1$O$_{62} < \alpha_2$-K$_7$As$_2$W$_{17}$V$_1$O$_{62}$ 순서로 증가하였다.

본 연구에서는 다양한 배위원소, 중심원소, 조성 및 구조를 지니는 헤테로폴리산의 제조, 분석 및 모델 산화반응으로의 적용을 통해 산화환원 특성과 산화촉매 반응을 이해하고자 하였다. 본 연구를 통해 헤테로폴리산 촉매의 산화환원 특성은 분자 내 구성성분을 달리함에 따라 쉽게 변화하는 것으로 나타났다. 특히, 헤테로폴리산 촉매의 산화환원 특성은 전자구조와 밀접한 연관성을 가지고 있는 것으로 판단할 수 있었으며, 산화반응에서의 촉매활성을 결정하는 중요한 특성임을 확인하였다. 또한 흡수단 애너지와 NDR 전압은 촉매의 환원력을 측정하기 위한 상호보완적 수치로서 이용될 수 있음을 확인하였다.

주요어: 헤테로폴리산, 전이 금속, 산화환원 특성, 산화반응, 전자구조

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