Low temperature oxidation of VOCs in air by catalytic ozonation

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By

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ABSTRACT

Alumina supported manganese oxides were used in the gas phase oxidation of toluene by ozone. Catalyst activity and characterization, the promotional effect of noble metals (Pt and Pd) on the activity of manganese oxides, and the kinetics and mechanism of the reaction were investigated in this PhD thesis. It was shown that MnO₂ and Mn₃O₄ were the active sites of the catalyst capable of oxidizing toluene to CO and CO₂ below 100 °C. Catalysts were deactivated at room temperature due to the accumulation of carbonaceous species on their surface. At least 65 °C was required for the stable operation of the catalysts. X-ray absorption spectroscopy was used to study the structure and electronic properties of the mono metallic and bimetallic catalysts. It was found that the catalysts with higher Mn loading resulted in higher oxidation states of Mn which were less favorable for the oxidation of toluene. The addition of Pt to the Mn containing catalyst increased the reaction rate by transferring electrons from Pt to Mn. On the other hand, no promotional effect was observed by the addition of Pd to Mn. The Oxidation state of Mn atoms was one of the most important parameters, controlling the rate of toluene oxidation. Lower oxidation states of Mn were able to easily transfer electrons to ozone, accelerating the rate of toluene oxidation. A reaction mechanism was proposed for the catalytic oxidation of toluene over manganese oxides. In this mechanism, the oxidation of toluene was carried out by the abstraction of hydrogen atoms followed by the oxidation of toluene carbon skeleton. A rate equation was derived based on this mechanism, determining the reaction orders of -1 and 2 for toluene and ozone, respectively. It was concluded that catalytic ozonation is an effective method for the low temperature oxidation of volatile organic compounds (VOCs) in air. The significance of this method is related to energy saving in air purifying systems by reducing the required temperature to oxidize VOCs. Catalytic ozonation can be used in indoor and outdoor applications for removal of VOCs from enclosed environments or polluted industrial streams.
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<thead>
<tr>
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<th>Description</th>
</tr>
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<tbody>
<tr>
<td>BE</td>
<td>binding energy (eV)</td>
</tr>
<tr>
<td>BET</td>
<td>Brunauer–Emmett–Teller</td>
</tr>
<tr>
<td>CLS</td>
<td>Canadian Light Source</td>
</tr>
<tr>
<td>CN</td>
<td>coordination number</td>
</tr>
<tr>
<td>$d_p$</td>
<td>particle size (μm)</td>
</tr>
<tr>
<td>$E$</td>
<td>activation energy (kJ mol$^{-1}$)</td>
</tr>
<tr>
<td>EPA</td>
<td>Environmental Protection Agency</td>
</tr>
<tr>
<td>EXAFS</td>
<td>extended X-ray absorption fine structure</td>
</tr>
<tr>
<td>$\Delta E_o$</td>
<td>difference between theory and data in EXAFS energy scale (eV)</td>
</tr>
<tr>
<td>$F_c$</td>
<td>F value in Fisher’s F-test</td>
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<tr>
<td>$F_{tol.}$</td>
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</tr>
<tr>
<td>FT-IR</td>
<td>Fourier transform infra red</td>
</tr>
<tr>
<td>GC</td>
<td>gas chromatograph</td>
</tr>
<tr>
<td>HVAC</td>
<td>heating, ventilating and air conditioning</td>
</tr>
<tr>
<td>$\Delta H$</td>
<td>enthalpy of adsorption (kJ mol$^{-1}$)</td>
</tr>
<tr>
<td>IAQ</td>
<td>indoor air quality</td>
</tr>
<tr>
<td>ICP-MS</td>
<td>inductively coupled plasma mass spectrometry</td>
</tr>
<tr>
<td>$k$</td>
<td>kinetic rate constant (atm$^{-1}$, mol$<em>{tol.}$ kg$</em>{cat.}$$^{-1}$ s$^{-1}$ atm$^{-2}$, mol$<em>{tol.}$ kg$</em>{cat.}$$^{-1}$ s$^{-1}$ atm$^{-1}$)</td>
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<td>$k_0$</td>
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<td>LH</td>
<td>Langmuir-Hinshelwood</td>
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<tr>
<td>---------</td>
<td>---------------------------------------------</td>
</tr>
<tr>
<td>Mn1</td>
<td>first Mn site in Mn$_2$O$_3$ structure</td>
</tr>
<tr>
<td>Mn2</td>
<td>second Mn site in Mn$_2$O$_3$ structure</td>
</tr>
<tr>
<td>MnO$_x$</td>
<td>manganese oxides</td>
</tr>
<tr>
<td>MS</td>
<td>mass spectrometer</td>
</tr>
<tr>
<td>NTP</td>
<td>non-thermal plasma</td>
</tr>
<tr>
<td>p</td>
<td>number of fitting parameter</td>
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<tr>
<td>P</td>
<td>pressure (atm)</td>
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<td>PCO</td>
<td>photo-catalytic oxidation</td>
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<tr>
<td>PDF</td>
<td>powder diffraction file</td>
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<tr>
<td>r</td>
<td>reaction rate (mol$<em>{\text{mol}}$, kg$</em>{\text{cat}}^{-1}$ s$^{-1}$)</td>
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<td>$R_f$</td>
<td>absolute misfit between theory and data in EXAFS fitting</td>
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<td>interatomic distance between cation i and anion j (Å)</td>
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<tr>
<td>$\Delta R$</td>
<td>change in the bond length between atoms (Å)</td>
</tr>
<tr>
<td>S</td>
<td>reaction selectivity</td>
</tr>
<tr>
<td>SBS</td>
<td>sick building syndrome</td>
</tr>
<tr>
<td>$S_0^2$</td>
<td>amplitude reduction factor (-)</td>
</tr>
<tr>
<td>$S_{ij}$</td>
<td>valence of the bond between cation i and anion j (-)</td>
</tr>
<tr>
<td>$\Delta S$</td>
<td>entropy of adsorption (J mol$^{-1}$ K$^{-1}$)</td>
</tr>
<tr>
<td>T</td>
<td>temperature (°C or K)</td>
</tr>
<tr>
<td>TIC</td>
<td>total ion chromatograph</td>
</tr>
<tr>
<td>TPR</td>
<td>temperature programmed reduction</td>
</tr>
<tr>
<td>UV</td>
<td>ultra violet</td>
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</table>
VOCs  volatile organic compounds
$V_i$  formal oxidation state of cation $i$
WHO  World Health Organization
$W_{\text{cat.}}$  catalyst weight (kg)
$X_{\text{tol.}}$  toluene conversion (mol$_{\text{tol.}}$ mol$_{\text{tol. in}}^{-1}$)
XANES  X-ray absorption near edge structure
XAS  X-ray absorption spectroscopy
XPS  X-ray photoelectron spectroscopy
XRD  X-ray diffraction

Greek letters

$\theta$  fraction of occupied or vacant catalyst active sites
$\nu$  theoretical stoichiometric coefficient
$\sigma^2$  mean-square displacement of the bond length ($\text{Å}^2$)
$\chi^2_\nu$  reduced chi-square

Subscripts

cal.  calculated
cat.  catalyst
eff.  effective
exp.  experimental
in  inlet
O  oxygen
ozo.  ozone

tol.  toluene

**Superscripts**

app.  apparent

o  standard state of gases

*  catalyst active site
CHAPTER 1
INTRODUCTION

1.1. **Research background and motivation**

There has been an increasing concern over air pollution control in recent years due to public and environmental health and safety issues. People are now dealing with the problem of air pollution more than in the past because of living in metropolitans and spending more time indoors. Outdoor air pollutants such as gaseous molecules (organic and inorganic) and particulate matters (suspended liquid or solid fine particles) can cause damage to people and the environment [1, 2]. There have been several reports on the effect of air pollutants on humans’ health. "Headache, dizziness, nausea, eye and skin irritations, respiratory system damages, lung and heart diseases, neurological effects, and cancer” are among the short and long term consequences of air pollution on people [1, 2].

The issue of air pollution is not limited to outdoor air. People inside enclosed environments such as residential buildings and work offices are exposed to air pollution problems. Sick building syndrome (SBS) is believed to be responsible for the development of diseases and allergies as the result of long term exposure of people to air pollutants inside buildings [3]. World Health Organization (WHO) has indicated that SBS can be identified in almost 30% of new houses. 6 million people of the population of Canada and half a million of people working in offices in Canada suffer from SBS [4]. Asbestos, CO, formaldehyde, NOx, SO2, radon, and volatile organic compounds (VOCs) are the most frequently detected pollutants in indoor air [5]. WHO statistics show that human deaths as the result of indoor air pollution in large cities are 14 times higher than those caused by outdoor air pollution. The annual death rate related to indoor air pollution is about 2.8 million which is equal to 5.5% of the total deaths.
occurring around the world each year [3]. WHO has considered the smoke emitted from burning solid fuels as one of the major risk factors for human health inside buildings [6]. These are all showing the importance of indoor air quality (IAQ) control which is the main motivation for the current research work.

VOCs are one of the most commonly found pollutants in indoor air. The concentration of indoor VOCs can be reduced by increasing the exchange rate of make-up air in heating, ventilating and air conditioning (HVAC) systems. However, homes nowadays have lower ventilation rates compared to the past because of the energy saving in HVAC systems, causing the accumulation of VOCs inside buildings [7]. The issue of energy saving is more serious in countries such as Canada with cold weather in most seasons of a year.

Several technologies have been developed for the removal of indoor VOCs to assist reducing energy consumption of HVAC systems. Sorption filtration, photo-catalytic oxidation (PCO), biological treatment systems, ozonation, non-thermal plasma (NTP), and combustion reactors are among the methods used for removal of indoor air VOCs [8-12]. Sorption filtration seems to be the most efficient method for the removal of VOCs from indoor air [10]. However, pollutants are not eliminated by adsorption. They are only transferred from a gas phase to a solid phase, making frequent regeneration of adsorbents necessary. Combustion of VOCs can also be considered to remove VOCs from air. In spite of the effectiveness of this method, combustion processes have poor energy efficiency. Performance of combustion chambers can be enhanced by using heterogeneous catalysts. It has been shown that VOCs such as benzene, toluene and xylene can be completely oxidized at around 200 °C using appropriate catalysts [13-16].

More decrease in the oxidation temperature of VOCs can be achieved if one replaces oxygen with ozone and uses appropriate catalysts. Transition metal oxides are effective catalysts
for ozonation of VOCs. Considering the high price of noble metal catalysts used for the oxidation of VOCs by oxygen, transition metal oxides have an economic advantage over noble metals due to their substantially lower cost of synthesis.

Catalytic ozonation has been explored for the oxidation of pollutants in liquid or gas phase over the past decades [17]. Application of this method in the gas phase has opened a new catalytic approach to low temperature oxidation of VOCs at trace concentration levels. Ethanol, carbon monoxide, acetone, cyclohexane, benzene, chlorobenzene, toluene, and formaldehyde are among the pollutants which have been oxidized by ozone over various catalysts [18-27]. The results show that the apparent activation energy of the oxidation of VOCs can be greatly reduced by ozone in comparison to that of oxygen, reducing the temperature of total oxidation of VOCs to less than 100 °C.

Manganese oxides (MnO₅) are among the most active transition metal oxides for low temperature decomposition of ozone to active oxygen species necessary for the oxidation of VOCs [28]. It is believed that the exceptional catalytic activity of MnO₅ is related to the capability of these oxides to have different oxidation states during a catalytic cycle, making them the most effective catalyst for the oxidation of VOCs by ozone [29].

Therefore, based on the needs for the development of novel air purifying systems, this dissertation is focusing on the oxidation of VOCs by catalytic ozonation at low temperatures. Toluene has been chosen as the model compound for this work since it has been frequently reported as one of indoor air pollutants. The catalyst of interest is manganese oxides (Mn₂O₃ and MnO₂) supported on γ-alumina because of the high activity of manganese oxides in the oxidation of VOCs by ozone.
1.2. **Structure of the thesis**

The framework of this thesis has been set to elucidate existing knowledge gaps in catalytic oxidation of toluene by ozone. The general performance of toluene oxidation by ozone, the catalyst activity of different manganese loadings, the effect of the addition of promoters to manganese oxides and the kinetics and mechanism of the reaction are among the topics investigated in this work. Technology development and field applications of the current method are not within the scope of this thesis which can be explored in the future works. It has to be noted that the outcomes of this work are not limited to indoor air and they can be extended to the removal of VOCs from outdoor air as well. A literature review on indoor air quality control and the oxidation of VOCs by catalytic ozonation is given in Chapter 2. Experimental details will be explained in Chapter 3 while the general performance of the catalytic oxidation of toluene is presented in Chapter 4. Chapter 5 contains the characterization and activities of different loadings of manganese oxides dispersed on alumina. The promotional effect of noble metals on manganese oxides will be presented in Chapter 6. The kinetics and mechanism of the reaction will be discussed in Chapter 7. An overall conclusion along with the recommendations for the future works are given in Chapter 8.

It has to be mentioned that the major content of this dissertation is already published in 4 peer reviewed journal articles which are quoted in this thesis. The articles are as follows:


The publisher of the papers is Elsevier which allows using the published papers in the dissertation, provided that the publisher is acknowledged. Therefore, by acknowledging the publisher in this section, no further citing will be made to the above mentioned papers throughout the rest of this thesis.

The author of this thesis, Ebrahim Rezaei Geshnizgani, was the major contributor of the above published papers. Literature review, ordering and purchasing the experimental supplies and equipment, design and construction of the experimental set-up, development of the analytical methods for GC-MS, planning and conducting the experiments, interpretation of the results, and writing and revising the manuscripts were done by Ebrahim Rezaei Geshnizgani. Dr Jafar Soltan was in charge of the supervision of the thesis, providing feedback on the manuscripts, and handling the submission process of the papers. Dr Ning Chen and Jinru Lin were included in the third paper for collecting the EXAFS data of the bimetallic catalysts in Chapter 6. Dr Ning Chen was also included in the second paper because of his technical assistance to Ebrahim Rezaei Geshnizgani throughout running the samples in HXMA beamline at CLS. This data was included in Chapter 5 on the study of atomic structure and the oxidation state of different loadings of MnOₓ/γ-alumina. Analysis of all the characterization techniques such as XRD, BET, H₂-TPR, FT-IR, laser Raman spectroscopy, CHNS, and X-ray absorption spectroscopy were performed by Ebrahim Rezaei Geshnizgani.
This chapter presents a review on indoor air quality (IAQ) and the currently available methods for the removal of VOCs from indoor air. It provides a brief introduction about air cleaner systems along with their advantages and disadvantages. It continues with a detail literature survey on the catalytic oxidation of VOCs by ozone in order to outline the current progress made in this area. By summarizing the work reported by different research groups, the knowledge gaps are identified and the research objectives are established.

2.1. Indoor air quality (IAQ)

It has been reported that the ancient humans were the first people affected by SBS due to exposure to the smoke from fire in caves [30]. It was in the middle of 1950s that outdoor and indoor air pollutants were clearly identified and measured [4]. However, the initial concern was only on CO$_2$, odors, micro-organisms and dust as indoor air pollutants [4]. It was accepted in the mid 1960s that several health problems and diseases of people inside buildings are related to indoor air pollution [4]. Since then, there has been the establishment of several acts and protocols to reduce the concentration of indoor air pollutants [7]. Some of the major steps in addressing indoor air quality are “clean air act (1963), establishment of US EPA (1970), ban of asbestos for structural insulation (1973), limiting the lead content of residential paints (1978), report of National Research Council (NRC) on indoor air pollutants (1981), restricting the maximum emission level of formaldehyde plywood used in homes by Department of Housing and Urban Development (HUD) (1985), establishment of indoor air quality office by EPA (1985), identification of high radon content areas by EPA (1988), smoking restriction in workplace in
California (1994), and issuing rules to limit the emission of VOCs from consumer products and architectural coatings by EPA (1998)”.

It can be seen that the presence of several contaminants have been restricted by different organizations over the past decades. Indoor air pollutants can be generally categorized as followings [5]:

- “Inorganic contaminants such as asbestos, radon, lead, CO, SO\textsubscript{2}, NO\textsubscript{x}.
- Organic contaminants such as VOCs.
- Biological contaminants such as viruses, bacteria, and mold.”

VOCs are one of the most commonly found pollutants inside enclosed environments. VOCs are molecules with boiling points between 50 and 260 °C [9]. They contain “aliphatic, cyclic hydrocarbons, aromatic hydrocarbons, aldehydes, terpenes, alcohols, esters, halocarbons, ketones, alkanes, organic acids, and ethers” [9]. WHO has identified 320 VOCs, in which 261 and 61 of them can be found in outdoor and indoor environments, respectively [4, 31]. Most of the indoor VOCs come from sources inside buildings. Some examples are chemicals used in building materials, furniture, cleaning agents, electronic appliances, and equipment [7]. High concentrations of many VOCs may result in neurological disorders [32]. They are also responsible for irritation of eyes, skin and lungs. In addition, they can cause “headache, drowsiness, fatigue, and confusion” [32]. Some VOCs such as benzene are also suspect to have carcinogenic effects. Among different VOCs, acetaldehyde, benzene, toluene, tetrachloroethylene, and trichloroethylene have been identified as some of the major pollutants by WHO, EPA, and French indoor air quality observatory (OQAI) [11]. The presence of VOCs in indoor air has made it necessary to develop new technologies for purifications of air in
enclosed environments. The next section provides a brief overview on different methods used to purify indoor air from VOCs.

2.2. Removal of VOCs from indoor air

There are five major methods for the removal of indoor VOCs. They are sorption filtration, photo-catalytic oxidation (PCO), non-thermal plasma (NTP), and ozonation. A summary on the principle and performance of these methods is provided below.

2.2.1. Sorption filtration

Gaseous VOCs can be effectively captured on the surface of adsorbents. Excellent adsorbents usually have high surface area within their porous structure. “Activated carbon (AC), zeolites, activated alumina, porous clay minerals, silica gel, and other molecular sieves” are among the commonly used adsorbents [5]. However, AC is often used for indoor air purification. The surface area of activated carbon is around 1000 m$^2$/g. Only a few grams of AC can adsorb most of the indoor air pollutants. One gram of activated carbon can adsorb around 1.0, 0.7, and 0.4 grams of 1,2-dibromomethane, carbon tetrachloride, and 1,1-dimethyl hydrazine, respectively [5]. AC is also an efficient adsorbent under humid conditions since it has a non-polar surface, resulting in the adsorption of hydrocarbons rather than water vapor [5].

The main problem with the sorption filtration is that adsorbents become deactivated over time and they frequently need to be regenerated. Another deficiency of adsorbents is that they cannot remove light molecules such as formaldehyde and acetaldehyde. An active catalyst can be impregnated on the surface of adsorbents in order to increase the removal efficiency of the
adsorption process by oxidizing light hydrocarbons. Nevertheless, it can be said that the most efficient method for the removal of indoor VOCs is adsorption [10].

2.2.2. Photo-catalytic oxidation (PCO)

Photo-catalytic oxidation (PCO) works under the principles of excitation of a semiconductor under ultra violet (UV) light. An electron can be ejected from the valence band of a semiconductor when it absorbs the energy of UV light, provided that UV light provides enough energy for the excitation of the electron [10]. This phenomenon creates a hole in the valence band. The holes and excited electrons are considered as active species in converting water vapor molecules to hydroxyl radicals, oxidizing organic pollutants [33]. PCO activity depends on the presence of water vapor and the activity of PCO reactors decreases in the absence of humidity. On the other hand, high concentrations of water vapor suppress the PCO reaction due to the adsorption of water molecules on PCO catalysts. In addition, it has been reported that there is an optimum point for inlet concentration of pollutants, giving the maximum oxidation rate of VOCs [8, 33].

PCO catalysts consist of semiconductors such as oxides of titanium, tin, zinc, iron, manganese, and tungsten [8]. TiO₂ is one of the most effective photo-catalysts in the oxidation of VOCs under UV light. Performance of PCO is related to several factors such as water vapor concentration, the source of UV light, and the inlet concentration of pollutants. Different types of PCO reactors also have different removal efficiencies, making the design of the reactors to be one of the major tasks in PCO based processes [33]. TiO₂ catalysts doped with transition metals have also shown to be capable of operating under visible light to oxidize VOCs, replacing UV light with natural sources such as sunlight [33].
PCO is a safe air cleaner at ambient conditions. It is effective for removal of different types of VOCs because of the high oxidation potential of OH radicals. However, there are a few drawbacks attributed to PCO. The first problem is related to the partial oxidation of VOCs and production of harmful by-products [33]. Photo-catalysts might also become deactivated by the accumulation of reaction by-products [33]. The energy efficiency of PCO is also low [10, 33]. TiO₂ has a polar structure which is not suitable for the removal of non-polar compounds. Overall, the mentioned limitations have restricted the widespread application of PCO [10, 11, 33].

2.2.3. Non-thermal plasma (NTP)

Highly energetic electrons are generated in atmospheric plasma in a way that the air matrix remains at room temperature. Gas phase molecules (nitrogen, oxygen, and water vapor) can be ionized with the produced electrons, generating reactive species such as atomic oxygen, hydroxyl radicals, and ozone [34]. These species are highly reactive and are able to oxidize organic pollutants. However, harmful by-products such as ozone, CO, and NOₓ are generated by NTP with low CO₂ selectivity of 0.3-0.7 [8, 35-37]. NTP reactors do not work steadily and their energy consumption is high [8, 35, 37].

In order to enhance the degradation efficiency and the energy consumption of non-thermal plasma reactors, it is possible to couple NTP with heterogeneous catalysts. Two different configurations can be used in order to incorporate catalysts in NTP [35]. The first option is to put catalysts inside the plasma region in a configuration called in plasma catalyst (IPC). The second choice is to utilize catalysts after the NTP reactors called post plasma catalyst (PPC) configuration.
ICP allows production of new active species such as peroxide or hydroxyl radicals, oxidizing VOCs on the surface of the catalysts. On the other hand, if catalysts are used after NTP (PPC), it is possible to use the generated ozone inside NTP to further oxidize undesired reaction by-products.

Manganese oxides dispersed on alumina was used as a PPC reactor, resulting in total oxidation of benzene, toluene and xylene to CO₂ [37]. A semiconductor such as TiO₂ can also be incorporated inside NTP reactors, combining non-thermal plasma and photo-catalytic oxidation techniques. This approach was used in the oxidation of toluene using TiO₂ as IPC and Pd, Cu, and Mn containing catalysts as PPC [34, 38]. Even though the combination of NTP and PCO has proven to be effective, it has been observed that the degradation of VOCs decreases by increasing the relative humidity of the inlet gas stream.

In general, the sole NTP method is not an effective technique for the removal of VOCs due to high energy consumption and low CO₂ selectivity, producing harmful by-products. The addition of catalysts inside or after the plasma region can increase the efficiency of the NTP method, but further investigations are needed before it is introduced into the market as a commercial product.

2.2.4. Ozone generators

Gaseous ozone can be considered as a potential oxidant for the removal of odor and VOCs from indoor air. Ozone generators have been used widely worldwide. Although it has been shown that ozone can be used for the elimination of odor, it reacts very slowly with organic pollutants in the gas phase. The half life of homogeneous reaction of ozone with VOCs such as formaldehyde, cyclohexane, toluene, and xylene ranges from 0.9 to 4400 years [39]. Formaldehyde is one of the most refractory molecules to react with ozone (half life of 4400
years) and toluene is one of the reactive species toward ozone with half life of 0.9 year. Due to the low reaction rate of ozone with VOCs, it is necessary to use high concentrations of ozone to completely oxidize organic compounds.

An ozone generator was used to study the performance of gaseous ozone in the removal of phenycyclohexene, styrene, and 4-ethynylcyclohexene [39]. These VOCs were emitted from a carpet located in an experimental chamber. Even though ozone was effective to reduce the concentration of these compounds, several harmful by-products such as formaldehyde and organic acids were also generated. As a conclusion, it can be said that the use of ozone generators as indoor air cleaners is not recommended due to the low reaction rate and the production of harmful by-products [5, 10, 39].

In an extended study, the performance of the above-mentioned methods except NTP, was tested using different indoor air cleaners, including sorption filtration, photo-catalytic oxidation and ozone generators [10]. The air cleaners were tested on several model compounds consisting of alkanes, aromatics, chlorocarbons, aldehydes, ketones, and alcohols. In comparison of different methods, it was concluded that sorption methods were efficient except for the removal of light hydrocarbons such as formaldehyde, acetaldehyde, and dichloromethane. PCO showed promising results in eliminating formaldehyde and acetaldehyde. The most important deficiencies of PCO were related to the high energy consumption and the efficient design of PCO reactors. It was suggested that further studies were needed to design high performance PCO air cleaners [10]. Authors did not recommend use of ozone generators for indoor air purifications [10].
2.3. Review on catalysts used for oxidation of VOCs by ozone

As mentioned in section 2.2, currently, there is not a single promising method for the elimination of VOCs from indoor air except the sorption filtration. It seems that further studies are needed on PCO and NTP methods before they can be commercialized for indoor air applications. The homogeneous reaction of ozone and VOCs is also slow and does not lead to the mineralization of organic pollutants. These limitations have encouraged researchers to study alternative approaches such as catalytic ozonation for indoor air quality control. Transition metal oxides such as (Fe, Co, Cu, Mn, Ni, and Cr) have been used for the oxidation of various VOCs by ozone. The results show that depending on the reaction condition and the type of catalysts used, VOCs can be oxidized in the temperature range of 22-100 ℃. The low temperature activity of catalytic ozonation makes this method suitable for indoor air treatment applications. This section describes the recent literature of catalytic oxidation of VOCs by ozone which helps to determine the knowledge gaps that exist in this research area.

Ozone is a reactive gas which was first discovered in 1839. But it was later in 1865 that the molecular structure of ozone was determined [17]. Ozone consists of three oxygen atoms forming an angle of 116°49’ between terminal atoms. The molecule represents four resonance structures shown in Fig. 2.1. The high reactivity of ozone is attributed to the electron configurations of the molecule. Lack or excess of electrons in different resonance forms of ozone determines the electrophilic or nucleophilic characteristics of ozone, respectively [40]. These properties make ozone highly reactive and also thermodynamically unstable leading to the decomposition of the molecule to oxygen by the reaction of $2O_3 \rightarrow 3O_2$. 

13
Ozone has an oxidation potential of 2.07 eV which is only less than fluorine, atomic oxygen, OH radical, and a few other oxidants [17]. Some other properties of ozone are given in Table 2.1 [17, 40].

<table>
<thead>
<tr>
<th>Properties</th>
<th>Value</th>
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<tbody>
<tr>
<td>Standard heat of formation ($\Delta H_{298}^0$)</td>
<td>-142 kJ mol$^{-1}$</td>
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<tr>
<td>Standard free energy of formation ($\Delta G_{298}^0$)</td>
<td>-163 kJ mol$^{-1}$</td>
</tr>
<tr>
<td>Standard entropy ($S_{298}$)</td>
<td>239 J K$^{-1}$ mol$^{-1}$</td>
</tr>
<tr>
<td>Melting point</td>
<td>-251 °C</td>
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<tr>
<td>Boiling point</td>
<td>-112 °C</td>
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</tbody>
</table>

There are three main research groups worked systematically on different aspects of the catalytic ozonation in the past two decades. The first group belongs to Naydenov and Mehandjiev and co-workers which started working on the oxidation of benzene and CO over transition metal oxides since 90s [19, 41-45]. MnO$_2$, NiMnO$_3$, NiMn$_2$O$_4$, NiO$_x$/alumina, CoO$_x$/alumina, Cu-Cr/alumina, and Co-Cr/alumina are among the transition metal oxides
investigated by them. Their results show that transition metal oxides are appropriate catalysts for low temperature decomposition of ozone and oxidation of benzene [41]. They have also indicated that the activation energy of benzene oxidation is greatly reduced by using ozone instead of oxygen. It was reported that the application of ozone could reduce the oxidation temperature of benzene from 150-200°C to 20-80°C [43]. It has been suggested that the formation of atomic oxygen on the surface of the catalyst as the result of ozone decomposition is responsible for the oxidation of benzene. Alumina supported Cu-Cr and Co-Cr oxides were studied in the oxidation of benzene and CO using oxygen or ozone [44]. Ozone greatly decreased the reaction temperature in the presence of Co-Cr/Alumina catalyst. Complete oxidation of benzene was achieved at around 100 °C by ozone.

The second research group in this field is Oyama and co-workers [18, 20, 28, 29, 46-51]. They also started their work in 90s, focusing on ethanol oxidation and the mechanism of ozone decomposition reaction. One of their first conclusions was that manganese oxides were the most active transition metal oxides in the decomposition of ozone. Oxides of Mn, Co, Ni, Cr, Ag, Cu, Ce, Fe, V, and Mo were selected by Dhandapani and Oyama for ozone decomposition on γ-alumina [28]. MnO₂ was the most active metal oxide in the decomposition of ozone followed by Co₃O₄, NiO, Fe₂O₃, and Ag₂O. Conductivity measurement showed that p-type metal oxides such as manganese oxide were more favorable in ozone decomposition comparing to n-type transition metal oxides. Li and Oyama investigated the kinetics and mechanism of catalytic ozone decomposition by in-situ laser Raman spectroscopy, Ab Initio calculations, and isotope exchange techniques [47, 48]. They have proposed that the decomposition of ozone can be described by the following elementary steps:

\[ \text{O}_3 + * \rightarrow \text{O}^* + \text{O}_2 \]  \hspace{1cm} (2.1)
O₃ + O* → O₂* + O₂  \hspace{1cm} (2.2)
O₂* → O₂ + * \hspace{1cm} (2.3)

In which * denotes catalyst active sites which are Mn atoms. Ozone is adsorbed dissociatively on the catalyst surface in step (2.1) to form a surface atomic oxygen (O*) and a gas phase oxygen molecule. The atomic oxygen reacts with the second ozone molecule in step (2.2) to form a surface peroxide (O₂*) and another oxygen molecule. The surface peroxide desorbs in step (2.3) to free the active site. Kinetic studies showed that the surface of the catalysts had a non-uniform nature. The activation energy of step (2.1) was determined to be less than the activation energy of step (2.3), showing that the desorption of ozone is controlling the rate of ozone decomposition.

Oyama and co-workers also worked on the total oxidation of acetone by ozone over silica supported manganese oxides [20, 29, 51]. They focused mainly on the characterization of the catalyst active sites by X-ray absorption spectroscopy, the effect of Mn loading on the catalyst activity, and the kinetics and mechanism of the reaction. They showed that the activation energy of acetone oxidation was reduced by ozone, leading to conversion of acetone to CO₂ below 100 °C [20]. Their results indicated that higher Mn loadings had better catalytic activity. It was mentioned that lower oxidation state of Mn atoms in higher Mn loadings was more favorable in the decomposition of ozone, accelerating the rate of acetone oxidation [29]. They also concluded that the reaction between acetone and ozone was structural sensitive, facilitated on larger manganese oxide clusters. They mentioned that the oxidation reaction needed adjacent Mn atoms to accommodate several atomic oxygen species required for the total oxidation of acetone [29]. Oyama and co-workers also performed fundamental studies on the kinetics and mechanism of acetone oxidation by ozone using in-situ laser Raman spectroscopy [51]. They proposed a
Langmuir-Hinshelwood (L-H) mechanism with the following elementary steps to explain the mechanism of acetone oxidation:

\[
O_3 + * \rightarrow O^* + O_2 \\
O_3 + O^* \rightarrow O_2^* + O_2 \\
O_2^* \rightarrow O_2 + * \\
A + □ ↔ A□ \\
A□ + * \rightarrow A* + □ \\
A* + O* \rightarrow \text{products}
\]

(2.4) to (2.9) refer to the decomposition of ozone on Mn sites while step (2.7) shows quasi equilibrated adsorption of acetone on the support (SiO\(_2\)). Adsorbed acetone molecules migrate from the silica support to Mn active sites, shown by step (2.8). Surface acetone molecules are activated by reacting with atomic oxygen in step (2.9). It was assumed that further steps after step (2.9) are fast and are not important from kinetic point of view. Peroxide species (O\(_2^*\)) were spectators in the mechanism and they were not involved in the oxidation of acetone. Authors derived rate equations for acetone oxidation and ozone decomposition based on the mechanism explained by steps (2.4) to (2.9).

The third research group which has been working on catalytic ozonation of VOCs is led by Einaga and co-workers [21, 22, 52-59]. Einaga and Futamura investigated the performance of alumina-supported metal oxides (Mn, Fe, Co, Ni, Cu, and Ag) for the oxidation of benzene and cyclohexane by ozone at room temperature [21, 52, 53]. Mn showed the highest activity for benzene oxidation at room temperature by converting benzene to CO and CO\(_2\). However, none of the metals showed stable activity in benzene oxidation. MnO\(_x\)/alumina lost its activity within 2
hours, due to the accumulation of reaction intermediates on the surface of the catalyst at room temperature. Weakly bound compounds including formic acid, 2,5-furandione and phenol, and strongly bound species such as formate and carboxylates were found on the catalyst surface as the result of incomplete oxidation of benzene at room temperature [53]. Einaga and Futamura showed that catalyst deactivation at room temperature can be suppressed by introducing water vapor to the gas matrix, inhibiting the accumulation of reaction by-products on the catalyst [60]. Positive effect of water vapor was also observed in the oxidation of formaldehyde by ozone over synthetic MnOₓ catalysts [61]. It was observed that formaldehyde can be completely oxidized to CO₂ under humid condition at room temperature.

Different ion-exchanged zeolites including Mn-Y, Mn-β, Mn-MOR, and Mn-ZSM5 were employed by Einaga and Futamura to oxidize benzene by ozone at 70 °C [54]. Mn-Y had the highest activity among all the catalysts with CO and CO₂ as the main reaction products. No catalyst deactivation was observed at 70 °C, resulting in 80% benzene conversion using Mn-Y catalyst. CO and CO₂ selectivity were 18 and 82%, respectively. The effect of different supports (Al₂O₃, TiO₂, ZrO₂, and SiO₂) in the oxidation of benzene by manganese oxides was studied by Einaga and Ogata [56]. The results show that catalysts with higher surface area and consequently more dispersed Mn atoms result in higher rates of benzene oxidation. SiO₂ with the highest surface area had the highest activity by converting benzene to CO and CO₂ at 100 °C. 60 °C was needed to attain steady state activity of the catalysts. Einaga et al. also studied the effect of Mn loading on the oxidation rate of benzene by ozone [55]. In contrast to the result of Oyama et al. reported for acetone oxidation, Einaga et al. concluded that lower loadings of Mn are more favorable for the oxidation of benzene.
Einaga et al. have reported that manganese oxides impregnated on ultra stable Y (USY) zeolites with high silica to alumina ratio (Mn/USY, SiO$_2$/Al$_2$O$_3$=180) is able to steadily oxidize benzene to CO and CO$_2$ at 22 and 40 °C under humid and dry conditions, respectively [22, 58]. It has been mentioned that high Mn dispersion and hydrophobicity of USY might be responsible for the exceptional activity of Mn/USY. In one of their recent publications, Einaga et al. have studied unsupported perovskite catalysts (LaMnO$_3$ and La$_{0.8}$Sr$_{0.2}$MnO$_3$) in the oxidation of benzene by ozone [59]. They have also investigated the effect of addition of Fe, Co, Ni, and Cu to Mn in order to enhance the activity of unsupported manganese oxide catalysts. The results show that monometallic Mn catalysts have higher activity than the perovskite catalysts. It was indicated that bimetallic catalysts have lower activity than monometallic manganese oxides, even though Mn-Co catalysts improved the selectivity of the reaction to CO$_2$.

There are also works published by other research groups, focusing on model compounds other than acetone, formaldehyde, cyclohexane, and benzene which were discussed above. CO, naphthalene, chlorobenzene, and toluene have also been oxidized by ozone over various catalysts [23-25, 62-71]. The results indicate that catalytic ozonation is an effective method for the oxidation of a wide variety of VOCs. Nevertheless, catalytic oxidation of toluene by ozone has received more attention among the mentioned VOCs. Chao et al. have studied the oxidation of toluene by ozone using zeolite 13X at room temperature [66]. 90% toluene conversion was observed at room temperature but the catalyst did not show stable activity because of the adsorption of CO$_2$, water, and reaction by-products. Zeolites NaX, NaY, and MCM-41 were chosen by Kwong et al. for toluene removal by ozone [67]. The results indicated that MCM-41 had the best performance for the ozonation of toluene at room temperature. No water and carbon dioxide were detected at the reactor outlet. This showed that reaction by-products were
accumulated on the catalyst surface, deactivating the catalyst. Sugasawa and Ogata investigated the performance of transition metals (Fe, Ni, Ag, Co, and Mn) supported on ZSM-5 in catalytic oxidation of toluene at 100 °C. Mn showed superior catalytic activity in the oxidation of benzene. They also mentioned that even though Ag was not as active as Mn, it had higher selectivity toward CO₂ among the transition metals tested [69]. This is in agreement with Einaga and Ogata, reporting higher CO₂ selectivity of Ag supported catalysts in benzene oxidation [57].

Catalytic ozonation of toluene was also studied by Park et al. using Mn supported on TiO₂ ultrafine nano composites [70]. Their results showed that this catalyst was also effective in the oxidation of toluene with the highest conversion of toluene to CO and CO₂ at 100 °C. Li et al. investigated the effect of modification of MCM-41 zeolites by Cu and Co on catalytic oxidation of toluene [25]. Catalysts were prepared by two different methods. In the first method, Cu and Co were loaded on MCM-41 by the conventional impregnation technique. In the second method, Cu and Co were incorporated into the structure of MCM-41 by in-situ synthesis of MCM-41 with the metals. The Co containing catalyst prepared by the in-situ formation of MCM-41 showed higher and more stable activity compared to the other catalysts for low temperature oxidation of toluene. Authors mentioned that high dispersion of Co atoms might be responsible for high activity of the Co incorporated MCM-41 catalyst.

In summary, it can be said that catalytic oxidation of VOCs by ozone is an effective method for low temperature oxidation of air pollutants. CO, benzene, acetone, formaldehyde, naphthalene, chlorobenzene, cyclohexane, and toluene are among the air contaminants removed by ozone. The operating temperature of catalytic ozonation process ranges from 22 to 100 °C, depending on the type of the catalyst, nature of the pollutant, and reaction conditions. Catalysts mostly consist of supported or unsupported transition metal oxides such as Mn, Cu, Co, Ag, and
Ni. This gives an extra advantage to catalytic ozonation processes since it removes the need for the synthesis of catalysts based on noble metals which are more expensive than transition metal oxides. Mn is the most active transition metal oxide for the decomposition of ozone and the oxidation of VOCs. The high activity of Mn is attributed to the electronic configuration of this metal oxide, allowing Mn atoms to exist at different oxidation states during the catalytic cycle of VOCs oxidation. Major reaction by-products of VOCs oxidation are CO and CO$_2$. Catalyst deactivation has been reported in case of aromatic compounds such as benzene and toluene at room temperature. Heating the catalyst bed to 60-70 °C increases the activity of the catalysts and results in steady state oxidation of aromatics. Water vapor has a positive effect on the activity of the catalysts at room temperature, preventing the deactivation of catalysts. The mechanism of the oxidation of VOCs by ozone follows a heterogeneous phenomenon. Ozone is decomposed to surface oxygen atoms which are responsible for the oxidation of adsorbed VOCs. The reaction mechanism can be described by steps (2.4) to (2.9) in which, adsorbed VOC molecules react with adsorbed oxygen atoms to start the oxidation reaction.

There are also technologies developed based on catalytic oxidation by ozone [17]. Toto, Inc. (Japan) has designed units for odor removal in toilets and refrigerators working by ozone and different types of catalysts. Hitachi (Japan) also has developed a catalytic ozonation unit called Dashpure™ for odor removal, disinfection, and oxidation of VOCs. UCAIR™ is the name of the technology developed by Union Carbide for the removal of VOCs, CO, and aerosol microbes from indoor air at ambient pressure and temperature. Nevertheless, there are not many reports on the commercial applications of catalytic ozonation, showing the need for more research on this method.
2.4. Knowledge gaps and objectives

Even though details of the reaction of ozone with VOCs such as acetone and benzene have been investigated widely, there are not many reports on catalytic oxidation of other aromatic compounds such as toluene or xylenes. Toluene has been frequently reported as one of the VOCs in indoor air. Therefore, toluene is considered as the model compound in the present study. Chapter 4 of this thesis elucidates different aspects of the catalytic oxidation of toluene by ozone over manganese oxides. The oxidation behavior of toluene at room and higher temperatures up to 100 °C is presented in this chapter. Characterization of the catalyst active sites, identification of the reaction by-products, and carbon balance of the reactor is discussed in Chapter 4 as well.

As mentioned in section 2.3, nano-particles of manganese oxides are mostly used in the oxidation of VOCs by ozone [26, 29, 55]. Less attention has been paid to the characterization of manganese oxides by novel techniques such as X-ray absorption spectroscopy including X-ray absorption near edge structure (XANES) and especially extended X-ray absorption fine structure (EXAFS). This is mainly due to the complicated unit cell structure of Mn₃O₄ and Mn₂O₃ with two Mn sites and several sets of oxygen atoms, closely located around the Mn sites. There are also controversial reports on the effect of Mn loading and catalyst structure on the activity of supported MnOₓ in the oxidation of VOCs [29, 55]. It is a matter of question whether higher or lower loadings of Mn benefit oxidation of VOCs by ozone. Therefore, Chapter 5 discusses activities of different loading of manganese oxides dispersed on alumina in the oxidation of toluene by ozone. This chapter clarifies the relationship among the structure of manganese oxides, the oxidation states of Mn atoms, and the observed catalytic activities. XANES and EXAFS are used as the main characterization techniques in Chapter 5 to justify the activity of the catalysts.
Investigation of the promotional effect of noble metals on the activity of manganese oxides is another area which has not been explored in the oxidation of VOCs by ozone. There are reports showing that the combination of transition metal oxides and noble metals such as Pt and Pd can enhance the activity of catalysts in the combustion of VOCs by oxygen [72-74]. The majority of researchers have only focused on the ozonation of VOCs over transition metal oxides. Thus, Chapter 6 of this thesis is dedicated to study the effect of addition of Pt and Pd to Mn in order to enhance the activity of supported manganese oxides in the oxidation of toluene by ozone. Atomic interactions between noble metals (Pt and Pd) and manganese oxides are studied using XANES and EXAFS. EXAFS helps to probe the atomic environment around Mn, Pt, and Pd in monometallic and bimetallic catalysts. XANES is also useful to track changes in the oxidation states of Mn, Pt, and Pd in order to explain the promotional effect of noble metals on manganese oxides.

Another issue which is not still completely clear is the mechanism of catalytic ozonation of aromatic molecules and relevant kinetic steps involved in the mechanism. It is believed that the oxidation reaction occurs between adsorbed atomic oxygen and VOC molecules according to Langmuir-Hinshelwood type of mechanism [51]. Even though this mechanism has been applied to explain the kinetics of acetone oxidation by ozone, there has not been any report to explore other possible reaction mechanisms which might better explain the oxidation of aromatic molecules such as toluene. Thus, Chapter 7 investigates the possible elementary steps involved in the catalytic oxidation of toluene by ozone over alumina supported manganese oxides. This is carried out by analysis of the effect of toluene and ozone partial pressures on the initial reaction rates and testing the applicability of the available reaction mechanisms reported in the literature.
to the catalytic ozonation of toluene. Possible alternative elementary steps will also be proposed which can better describe the observed kinetic data of toluene oxidation by ozone.
This chapter presents details of the experiments performed on catalytic ozonation of toluene by manganese oxides. Experimental set-up, analytical methods, and catalyst preparation are introduced first followed by activity tests and catalyst characterization techniques. The characterization methods includes Brunauer–Emmett–Teller (BET), inductively coupled plasma mass spectrometry (ICP-MS), X-ray diffraction (XRD), laser Raman spectroscopy, hydrogen temperature programmed reduction (H$_2$-TPR), X-ray absorption near edge structure (XANES), and extended X-ray absorption fine structure (EXAFS). Deactivated catalysts are studied by Fourier transform infra red (FT-IR) spectroscopy and CHNS analysis.

3.1. Experimental set-up and analysis

The schematic diagram of the experimental set-up is shown in Fig. 3.1. Toluene is supplied from a mixture of toluene in nitrogen cylinder (Praxair) with concentration of 300 ppm of toluene and balance of nitrogen (accuracy of ±2%). Oxygen (Praxair, ultra high purity grade, 99.993%) flows to an ozone generator (AZCO Industries LTD, HTU-500S) to produce ozone. Two mass flow controllers (Brooks, SLA 5850) are used to adjust flow rates of toluene-nitrogen and oxygen streams at desired set points. Accuracy of the mass flow controllers are ±1% of the reading flow rate. Total flow rate entering the reactor is 1000 ml min$^{-1}$.

The ozone and toluene streams are combined to produce toluene and ozone concentration of 90-240 and 1050-3000 ppm, respectively. In a typical experiment, 400 ml min$^{-1}$ of toluene/nitrogen stream is mixed with 600 ml min$^{-1}$ of ozone/oxygen stream to yield 1050 and 120 ppm of ozone and toluene, respectively. This combination results in nitrogen and oxygen
mole fractions of 39.94% and 59.94%, respectively. The reactor operating at ambient pressure is made of a horizontal Pyrex tube (i.d. 1/2 inch). The reactor is installed inside an oven (Binder, FP 115) which accurately controls the reaction temperature between room temperature and 100 °C. Fluctuation in the reaction temperature is less than ± 0.3 °C during the reaction.

Outlet stream of the reactor is passed through an ozone analyzer to measure the concentration of ozone. The ozone analyzer is an API 465M, working based on UV absorption to measure gas phase ozone concentration. The analyzer is set to measure the concentration of ozone from zero to 5000 ppm. The accuracy of the ozone analyzer is ±50 ppm.

Fig. 3.1. Schematic of the experimental set-up.
An on-line gas chromatograph (GC) (Agilent, 7890A) coupled with a mass spectrometer (MS) (Agilent, 5975C) is employed to measure the concentration of toluene, CO, CO₂, and other reaction by-products. The MS is set to work under scan mode. A capillary column (Varian, Select Permanent Gasses/CO₂, CP7429) is utilized to measure toluene, CO, and CO₂ concentrations. This column consists of four sub-columns. The first one is a 5 meter pre-retention gap column with diameter of 0.53 mm connected to two parallel columns. The first column is a CP-PoraBond Q (length: 50 m, dia.: 0.53 mm) for the detection of CO₂ and toluene and the second column is a CP-Molsieve 5A (length: 10 m, dia.: 0.32 mm) for the detection of CO. These two columns reconnect to a 5 meter post-retention gap column with diameter of 0.25 mm. Other oxygenated by-products are identified using an Agilent standard column HP-5MS (length: 30 m, dia.: 0.25 mm). A standard mixture of CO, CO₂, and nitrogen is used to quantify the concentration of CO and CO₂. The concentration of CO and CO₂ in the standard gas mixture is 650 and 630 ppm, respectively. Appendix A presents the GC-MS calibration data of toluene, CO, and CO₂.

3.2. Catalyst preparation

γ-alumina (Alfa Aesar, S_BET=212 m² g⁻¹, pore volume of 0.6 cm³ g⁻¹) is impregnated with aqueous solutions of manganese (II) nitrate tetrahydrate (Sigma-Aldrich, 97%) in order to synthesize alumina supported manganese oxides catalysts (MnOₓ/γ-alumina). The catalyst preparation method is wet incipient impregnation or dry impregnation. The loading of Mn is between 1 and 20%. Mn loading is calculated based on the mass of metallic manganese per mass of the catalysts (support plus manganese oxides). For example, in order to prepare 10 g of MnOₓ/γ-alumina with 10% Mn loading from the nitrate precursor, 42.3 g of manganese (II) nitrate tetrahydrate (Mn(NO₃)₂·4H₂O, MW=251.01) is added to 50 cm³ of deionized water under
vigorous stirring until the precursor is dissolved. This results in a solution with Mn(NO$_3$)$_2$.4H$_2$O concentration of 3.4 mol l$^{-1}$. Then 6 cm$^3$ of the prepared solution is taken by micropipette and added dropwise to γ-alumina which is already dispersed in a beaker. The impregnated support is dried at 100 °C for 10 h and calcined in the furnace under atmospheric air at 500 °C for 4h.

Tetraamineplatinum (II) nitrate salt (Alfa Aesar, 99.99%) and tetraaminepalladium (II) nitrate solution (Alfa Aesar, 99.9%) are used as precursors for the preparation of noble metal containing catalysts. The catalysts are labeled as Pd/γ-alumina and Pt/γ-alumina for monometallic catalysts and Pt-MnO$_x$/γ-alumina and Pd-MnO$_x$/γ-alumina for bimetallic catalysts. Pd/γ-alumina and Pt/γ-alumina are prepared by wet incipient impregnation of γ-alumina while Pt-MnO$_x$/γ-alumina and Pd-MnO$_x$/γ-alumina are synthesized by dry impregnation of the prepared MnO$_x$/γ-alumina catalyst. All the four catalysts are dried for 10 hours at 100 °C and calcined at 550 °C for 4 hours. Loadings of manganese and the noble metals are set at 10 % and 1 %, respectively.

In order to check the effect of particle size on the activity of the catalysts and to inhibit internal mass transfer limitations on the reaction rate, different ranges of particle size are tested for the reaction. The result of toluene conversion versus reaction time is shown in Fig. B.1 of Appendix B. It is observed that the activity of the catalyst, MnO$_x$/γ-alumina (10%), drops when particles larger than 208 μm are used. Thus, all experiments are conducted using catalysts with particle size less than 208 μm. The effect of external mass transfer on the reaction rate is also investigated in order to make sure that the catalytic oxidation of toluene is not controlled by the rate of external mass transfer of toluene from the gas phase. Details of the calculations are presented in Appendix C, showing that the oxidation of toluene proceeds in the absence of external mass transfer limitations.
MnO (Alfa Aesar, 99%, manganosite), MnO$_2$ (Alfa Aesar, 99.9%, pyrolusite), Mn$_2$O$_3$ (Sigma Aldrich, 99%, bixibyte), Mn$_3$O$_4$ (Sigma Aldrich, 97%, hausmannite), and α-PtO$_2$ (Alfa Aesar, 99.95%) are used as reference materials for characterization.

3.3. **Catalyst activity measurements**

Typically 0.2 g of fine powder catalysts is used for activity measurements. Catalysts are sandwiched between two pieces of glass wool and the empty space of the reactor is filled with glass beads in order to help mixing of the reactants before reaching the catalyst bed. The total flow rate of reactant at the inlet of the reactor is 1000 ml min$^{-1}$, corresponding to weight hourly space velocity (WHSV) of 300 l hr$^{-1}$ g$^{-1}$. Experiments are conducted in three phases. The first phase is saturation of catalysts with toluene in the absence of ozone for almost one hour until the outlet concentration of toluene reaches the inlet value. After the saturation of the catalysts, the ozone generator is turned on and the reaction starts at room temperature. Room temperature activity of the catalyst is monitored for 150 minutes. When the catalysts become deactivated at room temperature, reactor temperature is increased to 40, 60, 80, and 100 °C to study the catalyst activity at higher temperatures. At each temperature, the catalyst bed is allowed to stabilize for 80 minutes. Two samples are taken and compared at each temperature step before moving forward to the higher temperatures to test whether the reactor has reached steady state operating conditions. Raw data of a typical repeated experiment is given in Appendix D. The activity of catalysts versus time is obtained in another set of experiments. These experiments also start with the saturation of the catalyst with toluene at room temperature. But catalysts are allowed to work at different temperatures for longer period of time. Catalyst activities are monitored for 6 hrs at temperature levels of 22, 40, 55, 65, 80, and 100 °C. These experiments are suitable to study long term activity of the catalysts and to study the deactivation behavior of the catalysts.
A homogeneous reaction contribution (maximum toluene conversion of 18%) is observed in blank runs at different temperatures inside the empty reactor. Heterogeneous catalytic activities are calculated based on the subtraction of the homogeneous reaction rate from total reaction rate. In order to incorporate ozone concentration in the homogeneous reaction, a first order reaction is assumed for calculation of the homogeneous reaction rate.

3.4. Catalyst characterization

Brunauer–Emmett–Teller (BET) surface area and pore volume of the catalysts is determined by nitrogen adsorption at -196 °C using ASAP 2000 (Micromeritics). Samples are degassed at 200 °C for two hours under vacuum level of 500 μmHg. BET surface area is measured between nitrogen partial pressure ratio (P/P\text{atm}) of 0.05 and 0.3. Total pore volume of the catalysts is determined by measuring the amount of nitrogen adsorbed at P/P\text{atm} = 0.95. Loading of the catalysts is verified by inductively coupled plasma mass spectrometry (ICP-MS) on a PerkinElmer, Nexion 300D instrument. The weight percent of carbon accumulated on the catalysts is measured by CHNS analysis using Vario EL III, Elementar Americans Inc. Uncertainties of BET, ICP-MS, and CHNS analyses are presented in Appendix E.

X-ray diffraction (XRD) is performed on a Brucker diffractometer (D8 Advance) using Cu Kα in the 2θ range of 10-60° with a step size of 0.04°. Raman spectroscopy measurements are carried out on a Renishaw InVia Raman microscope using a 514.5 nm Ar+ laser (Spectra Physics) and a CCD detector. The instrument is operated in the line focus mode with a 10 s detector exposure time, and 32-128 spectra accumulations. Raman spectra are collected in the range of 100 to 1000 cm\(^{-1}\). Hydrogen temperature programmed reduction (H\(_2\)-TPR) is performed with an Autosorb-iQ (Quantachrome Instruments) TPD/TPR instrument equipped with a thermal conductivity detector (TCD). Samples are dried at 100 °C for two hours before the TPR analysis.
Reduction is performed in the range of 50 to 550 °C with the temperature ramp of 10 K/min using a gas mixture of 3.1% hydrogen balanced with nitrogen. The flow rate of the feed gas is 50 ml N min⁻¹.

Fourier transform infra red (FT-IR) spectra are recorded in the range of 400 to 4000 cm⁻¹ using 32 scans on a JASCO FT/IR 4100 spectrometer with DLATGS detector. The resolution of the detector is 4 cm⁻¹. Samples are diluted with potassium bromide (KBr) and pressed into disks before the analysis.

X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) of catalysts are collected at HXMA beamline, Canadian Light Source (CLS) having a storage ring with 200-250 mA current at 2.9 GeV [75]. The experiments are performed in transmission mode for Mn data collection of the catalysts and the reference materials. Fluorescence mode is used for the Pt and Pd catalysts. The samples are diluted with adequate amount of boron nitride (BN), pressed to thin disks and stuck on Kapton tape in order to obtain an approximately one unit edge jump at the absorption edge. Straight ion chamber detectors are used with Si(111) monochromator crystal and Rh mirrors as collimating and focusing mirrors, respectively. The scan step-sizes are 10 eV/step, 0.25 eV/step, and 0.05 Å⁻¹ for the pre-edge, XANES, and EXAFS regions, respectively.

Data processing including background removal, data averaging and normalization, energy calibrations, and determination of absorption energies are performed by Athena [76]. Absorption energies are obtained based on the position of the inflection point in the rising part of the spectra. The inflection point is determined by the energy of the major first peak in the 1st derivative of the normalized absorption spectra. EXAFS fitting is performed by Artemis which is a front-end graphical interface for FEFF [77] and IFEFFIT [78]. EXAFS fitting parameters are
passive electron reduction factor ($S_0^2$), coordination number (CN), change in interatomic distance ($\Delta R$), mean-square displacement of the bond length ($\sigma^2$), and change in energy scale ($\Delta E_o$). Uncertainties in processing EXAFS data in Artemis are estimated by evaluating the root mean square value of the measured signal in the range of 15 to 25 Å assuming that no structure can be measured in this range and any signal is due to shot noise [79]. CNs are fixed based on crystallographic data in EXAFS analysis of the reference materials to obtain $S_0^2$ which can be used as a fixed parameter in EXAFS analysis of the catalysts. This allows obtaining oxygen and manganese CNs at the first and second shells of MnO$_x$ clusters dispersed on γ-alumina. $\Delta R$, $\Delta E_o$, and $\sigma^2$ are allowed to be floating for the reference materials and the catalysts and their values are calculated by the fitting of EXAFS data.
Chapter 4 investigates the performance of manganese oxides dispersed on γ-alumina for the oxidation of toluene by ozone. The catalyst investigated in this chapter is MnOₓ/γ-alumina with manganese loading of 10%. The details of catalyst characterization, activity of the catalyst, identification of reaction by-products and reaction carbon balance are presented and discussed.

4.1. Characterization of MnOₓ/γ-alumina (10%)

4.1.1. BET and ICP-MS

Manganese oxides (MnOₓ) with nominal loading of 10% are impregnated on γ-alumina based on the method explained in section 3.2. The catalyst is called MnOₓ/γ-alumina (10%). Table 4.1 shows the loading of Mn measured by ICP-MS. BET surface area and pore volume of γ-alumina and MnOₓ/γ-alumina (10%) are also presented in Table 4.1. It can be seen that the pore volume and BET surface area of γ-alumina decrease after the impregnation of manganese oxides due to the partial pore plugging by MnOₓ clusters. However, the prepared catalyst still shows high surface area (S_BET=177 m² g⁻¹) and pore volume (0.51 cm³ g⁻¹), indicating that the porous structure of γ-alumina is preserved during the preparation procedure. The measured concentration of manganese by ICP-MS shows that the actual loading of Mn is close to the nominal value which is 10% by weight.
Table 4.1. BET surface area, pore volume, and metal loading of the catalyst

<table>
<thead>
<tr>
<th></th>
<th>γ-alumina</th>
<th>MnO&lt;sub&gt;x&lt;/sub&gt;/γ-alumina (10%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BET surface area (m&lt;sup&gt;2&lt;/sup&gt; g&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>212</td>
<td>177</td>
</tr>
<tr>
<td>Pore volume (cm&lt;sup&gt;3&lt;/sup&gt; g&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>0.60</td>
<td>0.51</td>
</tr>
<tr>
<td>Mn Loading wt%</td>
<td>-</td>
<td>9.5</td>
</tr>
</tbody>
</table>

4.1.2. X-ray diffraction (XRD)

Fig. 4.1 shows the powder diffraction files (PDF) of manganese oxides (MnO, MnO<sub>2</sub>, Mn<sub>2</sub>O<sub>3</sub>, and Mn<sub>3</sub>O<sub>4</sub>) [80] while Fig. 4.2 indicates the XRD spectra of the reference materials. The most intense peaks of MnO (manganosite) are located at 2θ of 34.9, 40.4, and 58.8 degrees as shown in Fig. 4.1 (A). PDF of Pyrolusite MnO<sub>2</sub> is shown in Fig. 4.1 (B) with the most intense peaks at 28.7, 37.3, 42.8, and 56.7 degrees. Mn<sub>2</sub>O<sub>3</sub> (bixbyite) has the major diffraction peaks at 23.1, 32.9, 38.2, 49.3, and 55.2 degrees as it can be seen in Fig 4.1 (C). PDF of Mn<sub>3</sub>O<sub>4</sub> (hausmannite) is shown in Fig. 4.1 (D) with the major peaks at 18.0, 28.9, 32.3, 36.1, and 59.9 degrees. One can identify the phases of the reference materials, shown in Fig. 4.2, by matching the diffraction pattern of the reference materials to the powder diffraction files presented in Fig. 4.1. Based on peak-matching, it can be concluded that MnO, MnO<sub>2</sub>, Mn<sub>2</sub>O<sub>3</sub>, and Mn<sub>3</sub>O<sub>4</sub> reference materials are manganosite, pyrolusite, bixbyite, and hausmannite phases of manganese oxides.
Fig. 4.1. Powder diffraction file (PDF) of reference materials, (A) MnO: manganosite, (B) MnO$_2$: pyrolusite, (C) Mn$_2$O$_3$: bixbyite, and (D) Mn$_3$O$_4$: hausmannite.
The XRD of MnOₓ/γ-alumina (10%) is shown in Fig. 4.3, indicating the formation of MnO₂ and Mn₂O₃ phases on γ-alumina. MnO₂ (pyrolusite) is identified with the most intense peaks at 28.7, 37.3, 42.8, and 56.7 degrees. In addition, Mn₂O₃ (bixbyite) has distinct peaks at 23.1, 32.9, 38.2, 49.3, and 55.2 degrees. Therefore, it can be concluded that MnO₂ (pyrolusite) and Mn₂O₃ (bixbyite) are formed on γ-alumina with Mn loading of 10%. Average particle sizes of MnO₂ and Mn₂O₃ on γ-alumina can be estimated by applying Scherrer’s equation [81]:

\[
t = \frac{K\lambda}{\beta \cos \theta}
\]  

(4.1)
In which, \( t \) is the thickness of the particle in nm, \( \lambda \) is the X-ray wavelength (0.154 nm), \( \beta \) is the peak width at half maximum in radians, \( K \) is a constant (0.9), and \( \theta \) is the angle between the beam and the normal on the reflecting lattice plane in radians. For particle size analysis, the most intense XRD peaks of MnO\(_2\) (28.7 degree) and Mn\(_2\)O\(_3\) (32.9 degree) are selected. It is estimated by Eq. 4.1 that MnO\(_2\) and Mn\(_2\)O\(_3\) have average metal cluster size of 17 and 23 nm on \( \gamma \)-alumina, respectively.

![XRD of the MnO\(_x\)/\( \gamma \)-alumina (10%).](image)

Table 4.2 contains the unit cell dimensions of MnO\(_2\) (pyrolusite) and Mn\(_2\)O\(_3\) (bixbyite) crystals along with the calculated dispersion of MnO\(_2\) and Mn\(_2\)O\(_3\) based on the particle sizes obtained from XRD [82]. It is determined that there are 8 and 20 atoms of Mn on the external
surface of the unit cell structure of MnO$_2$ and Mn$_2$O$_3$, respectively. This results in surface Mn atom density of $1.80 \times 10^{19}$ and $7.53 \times 10^{18}$ atom m$^{-2}$ for MnO$_2$ and Mn$_2$O$_3$, respectively. Therefore, a 17 nm cubic cluster of MnO$_2$ will have approximately 31130 Mn surface atoms. Similarly, a 23 nm cubic particle of Mn$_2$O$_3$ roughly contains 23910 surface Mn atoms. If one considers the density of MnO$_2$ and Mn$_2$O$_3$, reported in Table 4.2, and the number of Mn surface atoms in MnO$_2$ and Mn$_2$O$_3$ clusters, dispersion of MnO$_2$ and Mn$_2$O$_3$ can be estimated to be 6.1 and 3.8%, respectively.

Table 4.2. Mn dispersion in MnO$_x$/$\gamma$-alumina (10%)

<table>
<thead>
<tr>
<th></th>
<th>Lattice constants (nm)</th>
<th>Density (cm$^3$ g$^{-1}$)</th>
<th>Dispersion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MnO$_2$ (pyrolusite)</td>
<td>a=b=0.4396</td>
<td>5.03</td>
<td>6.1</td>
</tr>
<tr>
<td></td>
<td>c= 0.2871</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn$_2$O$_3$ (bixibyte)</td>
<td>a=b=c= 0.9408</td>
<td>4.50</td>
<td>3.8</td>
</tr>
</tbody>
</table>

4.1.3. Laser Raman spectroscopy

Raman spectra of the reference materials (MnO, MnO$_2$, Mn$_2$O$_3$, and Mn$_3$O$_4$), $\gamma$-alumina, and MnO$_x$/$\gamma$-alumina (10%) are depicted in Fig. 4.4. Mn$_3$O$_4$ is the most Raman sensitive phase of manganese oxides with a strong peak at 658 cm$^{-1}$. There are also two other peaks at 318 and 372 cm$^{-1}$ in the spectra of Mn$_3$O$_4$. These peaks have also been reported by Buciuman and coworkers as fingerprints of Mn$_3$O$_4$ [83]. MnO is the second Raman active phase of manganese. Raman spectra of MnO show two peaks at 552 and 652 cm$^{-1}$ which is in agreement with MnO Raman spectra reported by Kapteijn et al. [84]. MnO$_2$ and Mn$_2$O$_3$ are less Raman active oxides of manganese with small signal to noise ratios. MnO$_2$ generates two peaks at 531 and 662 cm$^{-1}$. Buciaman et al. have observed three peaks at 510, 580, and 640 cm$^{-1}$ for MnO$_2$ [83] whereas,
Kapteijn et al. did not identify any peaks in Raman spectra of this oxide [84]. In another work, Ferrandon et al. have obtained a signal at 532 cm\(^{-1}\) [85] which is the same as MnO\(_2\) peak at 531 cm\(^{-1}\) reported in our work. Mn\(_2\)O\(_3\) has four peaks at 198, 311, 644, and 699 cm\(^{-1}\), respectively. Peak locations of Mn\(_2\)O\(_3\) are in agreement with the work of Kapteijn et al. except the one at 198 cm\(^{-1}\) which cannot be observed in their results [84]. The reason that they have not detected this peak might be due to the fact that they collected the spectra from 200 to 1200 cm\(^{-1}\), leading to the absence of the peak at 198 cm\(^{-1}\).

Fig. 4.4. Raman spectra of the samples, (1) MnO, (2) MnO\(_2\), (3) Mn\(_2\)O\(_3\), (4) Mn\(_3\)O\(_4\), (5) \(\gamma\)-alumina, and (6) MnO\(_x\)/\(\gamma\)-alumina (10%).
As emphasized by Buciuman et al., there is a high potential for the formation of Mn$_3$O$_4$ during data collection of Raman spectra due to the local heating of samples [83]. Therefore, peaks at 311 and 644 cm$^{-1}$ might be assigned to in-situ formation of Mn$_3$O$_4$ during the analysis of Mn$_2$O$_3$ sample. The same point also applies to MnO$_2$, suggesting that the peak at 662 cm$^{-1}$ should not be mistaken as characteristic of MnO$_2$. Thus we consider the peak at 531 cm$^{-1}$ for the identification of MnO$_2$ and two peaks at 198 and 699 cm$^{-1}$ as characteristic of Mn$_2$O$_3$.

Raman spectra of γ-alumina and MnO$_x$/γ-alumina (10%) are also shown in Fig. 4.4. γ-alumina does not show any Raman signal, showing that γ-alumina is a Raman inactive material. On the other hand, there are two distinct peaks of Mn$_2$O$_3$ at 198 and 696 cm$^{-1}$ on the spectra of MnO$_x$/γ-alumina (10%), implying the presence of Mn$_2$O$_3$ in this catalyst. No MnO$_2$ peak is detected in MnO$_x$/γ-alumina (10%) which might be expected to be detected at around 531 cm$^{-1}$. It is noteworthy to mention that the peak at 640 cm$^{-1}$ is due to the in-situ formation of Mn$_3$O$_4$ and it is not considered as an indication of Mn$_3$O$_4$ on MnO$_x$/γ-alumina (10%). The reason for the absence of MnO$_2$ Raman signal in the spectra of MnO$_x$/γ-alumina (10%) is likely due to the low fraction of MnO$_2$ in comparison to the fraction of Mn$_2$O$_3$ on the surface of the catalyst.

4.1.4. X-ray absorption near edge structure (XANES)

Fig. 4.5 contains the normalized absorption XANES spectra of Mn K-edge for manganese oxide reference materials and MnO$_x$/γ-alumina (10%). XANES spectra of manganese oxides contain a weak pre-edge feature due to the quadruple (dipole-forbidden) transition of electrons from 1s to 3d orbital. The main absorption happens due to the dipole-allowed transition of electrons from 1s to 4p orbital of Mn giving the rising part of the spectra. It can be seen that the spectra of MnO$_x$/γ-alumina (10%) is located between that of Mn$_2$O$_3$ and MnO$_2$, suggesting that
manganese has an average oxidation state between +4 and +3. Linear combination fitting (LCF) of MnOₓ/γ-alumina (10%) with MnO₂ and Mn₂O₃ reference materials shows that 91.0% of Mn is in +3 and 9.0% of Mn is in +4 oxidation states.

Fig. 4.5. Mn K-edge XANES spectra, (1) MnO, (2) Mn₃O₄, (3) Mn₂O₃, (4) MnOₓ/γ-alumina (10%), and (5) MnO₂.

It can be concluded from the result of XRD, laser Raman spectroscopy, and XANES that MnO₂ and Mn₂O₃ are the two phases of manganese oxides which are generated by using the nitrate precursor of Mn for the catalyst preparation. XANES analysis reveals that Mn₂O₃ is the dominant phase of manganese oxides on MnOₓ/γ-alumina (10%) with weight fraction of 91.0%.
4.2. Activity of MnOₓ/γ-alumina (10%)

4.2.1. Oxidation of toluene by MnOₓ/γ-alumina (10%)

Fig. 4.6 and 4.7 show the activity of MnOₓ/γ-alumina (10%) in terms of toluene and ozone conversions, respectively. Concentration of ozone and toluene are 1050 and 120 ppm, respectively. The conversions are obtained at 22, 40, 55, 65, 80, and 100 °C for 6 hours at each temperature level using 0.2 g of MnOₓ/γ-alumina (10%). The flow rate of toluene/nitrogen and ozone/oxygen streams are 400 and 600 ml min⁻¹, respectively. It can be observed in Fig. 4.6 that the catalyst shows high initial activity at all temperatures. But the activity of catalysts decreases at temperatures below 65 °C, implying the deactivation of the catalyst.

Fig. 4.6. Toluene conversion versus time over MnOₓ/γ-alumina (10%), (1) 22 °C, (2) 40 °C, (3) 55 °C, (4) 65 °C, (5) 80 °C, and (6) 100 °C.
MnOx/\(\gamma\)-alumina (10\%) shows stable activity in the oxidation of toluene at 65, 80, and 100\(^\circ\)C after almost 150 minutes reaction. Maximum toluene conversion is around 95\% which is obtained at 80\(^\circ\)C. A decrease in toluene conversion can be seen by increasing the temperature from 80 to 100\(^\circ\)C.

Ozone conversions at different temperatures are shown in Fig. 4.7, in which the conversions increase by increasing the temperature. No steady state ozone conversion is obtained below 65\(^\circ\)C. Ozone conversion reaches 100\% at 80\(^\circ\)C and remains constant by increasing the temperature from 80 to 100\(^\circ\)C. In fact, the 80\(^\circ\)C is the temperature, at which ozone is completely consumed under the given reaction condition. Therefore, the increase of temperature from 80 to 100\(^\circ\)C does not improve the ozone conversion. This can be considered as the reason
for the small decrease in toluene conversion above 80 °C, since all ozone is already decomposed at 80 °C. The same behavior has been reported in catalytic ozonation of acetone by MnO$_x$/γ-alumina and MnO$_x$/SiO$_2$, in which a fluctuating plateau occurs above 80 °C due to complete conversion of ozone, keeping acetone conversion more or less constant [20].

Carbon content of the deactivated catalyst, MnO$_x$/γ-alumina (10%), is determined by CHNS analysis. Data is obtained after the catalyst becomes deactivated at room temperature after 150 minutes. Bare support, γ-alumina, is also tested without exposure to the reaction medium to determine a baseline for the initial carbon content of the support. The fresh support contains negligible carbon (0.20%), whereas the deactivated catalyst showed 10.25% carbon content after the deactivation. Thus, it can be inferred that the accumulation of carbon containing species on the catalyst blocks active sites and reduces the catalyst activity at room temperature. Einaga and coworkers studied the deactivated alumina supported manganese oxides in the oxidation of benzene by ozone [53, 56]. They reported that two types of by-products were formed on the surface of the catalysts. The first type is weakly bonded molecules such as carboxylic acids (formic acid, acetic acid, and oxalic acid) and the second type is strongly attached species such as formate and carboxilates. Thus, it can be assumed that oxygenated by-products formed as the result of incomplete oxidation of toluene are responsible for the deactivation of the catalysts at room temperature. By increase of the reaction temperature, oxidation of these by-products is promoted and the catalyst regains its activity at temperatures higher than 65 °C.

In order to clarify the contribution of γ-alumina in the activity of MnO$_x$/γ-alumina (10%), 0.2 g of bare γ-alumina (not impregnated by manganese oxides) was tested for the oxidation of toluene by ozone. It was observed that γ-alumina resulted in constant conversions of 20 and 10% for toluene and ozone in the range of 20-100 °C, respectively. By comparing the low activity of
\(\gamma\)-alumina (not more than 20\% in terms of toluene conversion) with the activity of manganese oxides impregnated \(\gamma\)-alumina (Fig. 4.6 and 4.7), one can conclude that the active phase of the catalyst is manganese oxides. As it was revealed in section 4.1, Mn\(_2\)O\(_3\) is the dominant phase of manganese oxides in MnO\(_x\)/\(\gamma\)-alumina (10\%) which can be considered as the active component of the catalyst in this work.

The oxidation mechanism of VOCs by ozone is considered to be based on the decomposition of ozone to atomic oxygen and peroxide species on manganese oxide clusters which can be expressed by the following steps [20]:

\[
\begin{align*}
O_3 + * & \rightarrow O_2 + O^* \quad (4.2) \\
O_3 + O^* & \rightarrow O_2 + O_2^* \quad (4.3) \\
O_2^* & \rightarrow O_2 + * \quad (4.4)
\end{align*}
\]

In equations 1 to 3, * corresponds to Mn sites. It is believed that the generated atomic oxygen is responsible for the oxidation of VOCs [20, 56].

In another study, the conversion of toluene is monitored over 0.2 g of MnO\(_x\)/\(\gamma\)-alumina (10\%) in the absence of ozone. 120 ppm of toluene in nitrogen and oxygen (toluene/nitrogen flow rate of 400 ml min\(^{-1}\) and oxygen flow rate of 600 ml min\(^{-1}\)) is passed over the catalyst bed while the ozone generator is kept off. The reaction performance of toluene conversion versus temperature is shown in Fig. 4.8 (a). No toluene conversion is observed in the temperature range of 22-100 °C, indicating the crucial role of ozone in low temperature oxidation of toluene over the catalyst. In order to explore the applicability of this method to actual air composition, the flow rates of toluene/nitrogen and ozone/oxygen streams are changed to 800 and 200 ml/min, respectively. These flow rates set the concentration of N\(_2\) and O\(_2\) at 80 and 20\%, respectively. This arrangement results in toluene concentration of 240 ppm. Accordingly, the power of ozone
generator is set to produce 2500 ppm ozone at the inlet of the reactor in order to have enough ozone to react with toluene. Activity of the catalyst with actual air composition (toluene/nitrogen = 800 ml min$^{-1}$ and ozone/oxygen = 200 ml min$^{-1}$) is shown in Fig. 4.8 (b). Fig. 4.8 (c) shows the activity of the catalyst with 120 ppm toluene and 1050 ppm ozone obtained by mixing 400 ml min$^{-1}$ of toluene/nitrogen mixture with 600 ml min$^{-1}$ of ozone/oxygen mixture for comparison. It has to be noted that 0.2 g of MnO$_x$/γ-alumina is used to obtain all activities in Fig. 4.8.

![Graph showing toluene conversion versus temperature](image)

**Fig. 4.8.** Toluene conversion versus temperature, (a) in the absence of ozone, N$_2$ mole fraction ≈ 40%, O$_2$ mole fraction ≈ 60% (b) actual air composition with N$_2$ mole fraction ≈ 80% and O$_2$ mole fraction = 20%, and (c) N$_2$ mole fraction ≈ 40%, O$_2$ mole fraction ≈ 60%.

The same trend of activity is observed when toluene was oxidized in actual air composition. However, slightly higher activities are obtained in actual air composition due to higher ratio of ozone to toluene. This shows that the oxidation of toluene is not very sensitive to
the composition of air and it is mostly related to the concentration of ozone which is assumed to be the main oxidant in this system.

4.2.2. Reaction by-products

CO and CO\(_2\), (CO\(_x\)), are the main products of the catalytic ozonation of toluene over 0.2 g of MnO\(_x\)/γ-alumina (10%). Fig. 4.9 presents the concentration of CO and CO\(_2\) measured in the temperature range of 22-100 °C. It can be seen that CO and CO\(_2\) concentrations are very low at room temperature due to the low conversion of toluene and the accumulation of carbon species on the catalysts. As the reaction temperature increases, the yield of the reaction to CO\(_x\) increases and reaches a maximum (≈ 90%) at 80 °C.

The carbon balance of the reactor is shown in Fig. 4.10. The carbon balance is around 90% at temperatures that the catalyst, MnO\(_x\)/γ-alumina (10%), works steadily (above 65 °C) and around 80% at room temperature. The difference in the carbon balance is attributed to the production of by-products other than CO and CO\(_2\) and the accumulation of carbon species on the catalysts below 65 °C.
Fig. 4.9. Concentration of CO and CO₂ at different temperatures.

Fig. 4.10. Carbon balance of the reactor.
Fig. 4.11 shows a total ion chromatograph (TIC) of reaction products of catalytic ozonation of toluene on MnO$_x$/γ-alumina (10%) at 80 °C, collected by the mass spectrometer. Oxalic acid, acetic acid, benzene, and maleic anhydride are detected as the minor reaction by-products. The difference in the carbon balance (10% carbon loss during the steady state operation of the catalyst) is due to the formation of these by-products produced as the result of incomplete oxidation of toluene. Monneyron et al. have studied a combined adsorption-ozonation process for the removal of toluene by zeolites [65]. Using GC-MS, they showed that CO$_2$, water, formic acid, oxalic acid, acetic acid, and acetaldehyde are the main reaction by-products of the ozonation of adsorbed toluene on Fau Y zeolite. Although they have not used any metal oxides impregnated on Fau Y and their process is based on saturation of zeolites followed by the ozonation of the adsorption bed, oxalic acid and acetic acid are two common by-products found in the present work and the work by Monneyron et al.

If one assumes that only atomic oxygen is involved in the oxidation of toluene and the only reaction byproduct is CO$_2$, the following equation can describe the maximum ozone to toluene ratio required for complete oxidation of toluene:

$$\text{C}_7\text{H}_8 + 18\text{O}_3 \rightarrow 18\text{O}_2 + 7\text{CO}_2 + 4\text{H}_2\text{O}$$  \hspace{1cm} (4.5)

In this work ozone to toluene ratio is 8.75 which is almost half of the theoretical reaction stoichiometry. In fact, ozone is the limiting reactant in our work and the limitation of oxygen species can explain the formation of byproducts indicated in Fig. 4.11. Einaga and Futamura hypothesized that not only ozone but also oxygen can participate in the oxidation of benzene. They proposed that oxygen is involved in some form of auto-oxidation reactions to oxidize radical reaction intermediates to CO and CO$_2$ [56]. This hypothesis along with the production of CO besides CO$_2$ can explain how ozone to toluene ratio of 8.75, which seems to be far from the
reaction stoichiometry, leads to 90% carbon balance. Increase of ozone concentration to 2200 (ozone to toluene ratio of 18.3) ppm led to near 100% carbon balance at 100 °C with CO and CO₂ concentrations of 205 and 633 ppm, respectively. This shows that when enough oxygen atoms are provided, CO and CO₂ are the only byproducts of reaction. However, even when ozone to toluene ratio is close to the theoretical stoichiometric value, there is still some CO formation indicating incomplete oxidation of toluene.

Fig. 4.11. Total ion chromatograph (TIC) of reaction byproducts at 80 °C obtained on MnOₓ/γ-alumina (Nit), (1) N₂-O₂, (2) oxalic acid, (3) acetic acid, (4) benzene, (5) toluene, and (6) maleic anhydride.
4.3. Summary

Manganese oxides were impregnated on γ-alumina ($S_{BET} = 212 \text{ m}^2 \text{ g}^{-1}$ and $V_{pore} = 0.60 \text{ cm}^3 \text{ g}^{-1}$) from the nitrate precursor of Mn which is manganese (II) nitrate tetrahydrate. The loading of Mn was 10% by weight. The catalyst was called MnO$_x$/γ-alumina (10%). The final catalyst was characterized by X-ray diffraction (XRD), laser Raman spectroscopy, and X-ray near edge structure (XANES). The results of XRD analysis showed that MnO$_2$ (pyrolusite) and Mn$_2$O$_3$ (bixbyite) formed on γ-alumina. Dispersion of Mn was calculated to be 6.1 and 3.8% for MnO$_2$ and Mn$_2$O$_3$, respectively. Laser Raman spectroscopy also confirmed the presence of Mn$_2$O$_3$ particles in the catalyst, even though no MnO$_2$ was detected by Raman spectroscopy. XANES analysis showed that the near edge spectra of the catalyst was located between that of MnO$_2$ and Mn$_2$O$_3$ reference materials, indicating that two phases of manganese oxides can be found in the catalyst. Linear combination fitting of the catalyst with MnO$_2$ and Mn$_2$O$_3$ reference materials revealed that 9% of manganese oxides are in the form of MnO$_2$ and 91% are in the form of Mn$_2$O$_3$.

Toluene oxidation was carried out in the temperature range of 22-100 °C using 0.2 g of the catalyst. Results show that the catalyst lost its activity below 65 °C due to the accumulation of carbon containing species on the catalyst surface. Almost complete conversion of toluene was achieved at 80 °C. Experiments with the support alone (γ-alumina) revealed that the most active part of the catalyst was manganese oxides. Considering the characterization results, Mn$_2$O$_3$ can be suggested to be the catalyst active site in MnO$_x$/γ-alumina (10%). CO and CO$_2$ were determined to be the major reaction products, leading to 90% carbon balance at 80 °C. Minor reaction by-products were identified by the mass spectrometer. Oxalic acid, acetic acid, benzene, and maleic anhydride were detected in addition to CO and CO$_2$, implying incomplete oxidation.
of toluene. The mechanism of the reaction was assumed to be based on the decomposition of ozone on manganese oxides to peroxide and atomic oxygen. The produced surface oxygen atoms are highly reactive capable of oxidizing toluene molecule. It was observed that at higher ozone to toluene ratios, carbon balance approaches 100% with CO and CO₂ as the only reaction products.
CHAPTER 5
EFFECT OF CATALYST LOADING ON ACTIVITY OF ALUMINA SUPPORTED MANGANESE OXIDES

As it was observed in chapter 4, MnO₂ and Mn₂O₃ are the oxide phases of manganese found in MnOₓ/γ-alumina (10%). In order to investigate the relationship between the phases of manganese oxides and the activity of alumina supported manganese oxides, four catalysts with different Mn loadings (1, 5, 10, and 20%) are studied in this chapter. The catalysts activities are presented in the temperature range of 22-80 °C for the oxidation of toluene. Catalyst active sites are characterized by X-ray diffraction (XRD), X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS). There are a few studies on the atomic structure of manganese oxides performed by EXAFS [29, 58, 86-88]. The EXAFS study of MnO₂ is rather straightforward since it has only one Mn site. But difficulty arises in the EXAFS analysis of Mn₂O₃ and Mn₃O₄ due to the existence of two Mn sites and several sets of oxygen atoms located closely at the first shell around the Mn sites. These oxygen atoms are usually 0.1 to 0.3 Å apart, making it difficult for them to be distinguished by the sole EXAFS fitting procedure. Therefore, more emphasis is put on the EXAFS study of the reference materials (MnO₂ and Mn₂O₃) and the catalysts in this chapter. EXAFS of Mn₂O₃ is presented in more detail in order to build rigorous theoretical models to perform EXAFS analysis of this oxide. A brief introduction on X-ray absorption spectroscopy is also given at the beginning of Chapter 5 to provide readers with the basics of XANES and EXAFS. The goal of chapter 5 is to relate the observed catalyst activities to the electronic and structural properties of the catalysts with different Mn loadings.
5.1. X-ray absorption spectroscopy (XAS)

The atoms of elements can be excited by the absorption of the energy of X-ray photons. If the energy of the emitting X-ray is less than the binding energy of electrons, the electrons are promoted to unoccupied higher energy level orbitals, creating core holes in their original orbitals, from which they have been removed. The transition of electrons from lower energy orbitals to higher energy orbitals follows the dipole selection rule, by which only certain transitions are permitted between orbitals [89]. The excited electrons are called photoelectrons. The core hole will be filled with electrons in higher energy levels in a femto-second ($10^{-15}$ s) [89]. The transition of electrons from higher to lower energy levels causes release of Auger electrons, fluorescence radiation, and photons. The absorption of X-ray by a material can be expressed by Beer-Lambert’s law [89]:

$$I_t = I_o e^{-\mu x}$$  \hspace{1cm} (5.1)

$I$ is the intensity of the X-ray and subscripts $t$ and $o$ are used for transmitted and incident X-ray, respectively. $\mu$ is the absorption coefficient and $x$ is the thickness of the matter. The amount of the X-ray energy absorbed by a specific element depends on the density of the sample, the atomic mass and the atomic number of the atoms within the material [90]. As an example, the normalized absorption spectrum of Mn $K$-edge of Mn$_2$O$_3$ is shown in Fig. 5.1, in which the normalized absorption is plotted as a function of the energy of X-ray. The spectrum consists of two main parts which are called X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS). The XANES part starts a few hundred electronvolts below the excitation energy of the electrons in Mn 1S orbital. By increasing the energy of the X-ray, electrons are excited from the 1S orbital to 4P orbitals, making the rising part (white line) of
the spectrum. XANES data can be used to determine the formal oxidation state of an atom. It may also provide information about the geometry of atoms around the absorbing atom.

By further increase of the X-ray energy, the photoelectrons are removed from the atom and sent to the continuum. Photoelectrons propagate as three-dimension waves among atoms and are scattered from neighboring atoms back to the absorbing atom. Therefore, interactions occur among the outgoing photoelectron waves and the backscattered waves at the absorbing atom, affecting the absorption of X-ray. If the two waves (emitted and backscattered) are in phase, a maximum can be seen in the absorption spectrum. On the other hand, if the waves are out of phase, a minimum occurs in the absorption spectrum. The oscillatory part of the spectrum above the edge is called extended X-ray absorption fine structure (EXAFS). Usually there are

Fig. 5.1. X-ray absorption spectrum of Mn K-edge of Mn$_2$O$_3$. 
several atoms at different distances around the absorbing atom which can back scatter the photoelectrons. Each set of these atoms at equal distances from the absorbing atom is called a shell. The journey of the photoelectrons from the absorbing atom to the neighboring atoms and back to the absorbing atom is identified as a path. A single scattering path contains only one neighboring atom while a multiple scattering path may consist of several neighboring atoms at different shells. This makes the EXAFS an atomic scale probe to investigate the atomic environment of an atom. The atomic range investigated by the EXAFS is about 6 Å [89]. The EXAFS oscillations are defined from the experimental data by [90]:

\[
\chi(E) = \frac{\mu(E) - \mu_0(E)}{\Delta\mu_0(E)} \tag{5.2}
\]

\(\mu(E)\) is the absorption coefficient at X-ray energy of E and \(\mu_0(E)\) is the smoothed background in the EXAFS region. \(\Delta\mu_0(E)\) is the absorption jump at the absorption edge. Photo-electrons’ wave number \(k\) is usually used instead of the X-ray energy \(E\) to define the EXAFS function. The \(k\) dependent EXAFS function is shown by \(\chi(k)\) in which, \(k\) is related to \(E\) by [90]:

\[
k = \sqrt{\frac{2m(E - E_o)}{\hbar^2}} \tag{5.3}
\]

\(m\) is the mass of the electron and \(\hbar\) is the Plank’s constant. \(E_o\) is the energy of absorption in electronvolts (eV) and \(k\) is in Å⁻¹.

The theory of the EXAFS phenomenon has been well developed now and it is possible to extract information such as coordination numbers, inter-atomic bond lengths and bond disorders from the EXAFS experimental data [91-93]. The theoretical EXAFS equation for the \(i^{th}\) path can be written as [89]:

\[
\chi_i(k) = \frac{(CN_i^2)F_{\text{eff}}(k)}{kR_i^2} e^{-2\sigma_i^2k^2} e^{-2R_i/\lambda(k)} \sin[2kR_i + \phi_i(k)] \tag{5.4}
\]
k and $R_i$ are given by:

$$k = \sqrt{\frac{2m(E - E_o + \Delta E_o)}{\hbar^2}}, \quad R_i = R_{Reff, i} + \Delta R_i$$

(5.5)

The total EXAFS oscillation can be obtained by considering the contributions of all paths as a sum over all $\chi_i(k)$:

$$\chi(k) = \sum_i \chi_i(k)$$

(5.6)

$\lambda(k)$, $\varphi_i(k)$ and $F_{ef, i}$ in Eq. (5.4) are the photoelectrons mean free path, phase shift, and the effective amplitude of the backscattered photoelectron, respectively. The phase shift occurs since the photoelectron encounters the atomic potentials of the absorbing and the neighboring atoms as they travel in a path. These three parameters ($\lambda(k)$, $\varphi_i(k)$ and $F_{ef, i}$) are calculated from theoretical quantum mechanics. FEFF is a computer code which is usually used to estimate these parameters [77]. (CN, $S_o^2$) is multiplied to $F_{ef, i}$ to correct the EXAFS signal amplitude. CN is the coordination number for single scattering paths or the number of similar paths in multiple scattering paths. $S_o^2$ is the amplitude reduction factor which is related to the relaxation of electrons after the core hole is generated. Typical values of $S_o^2$ are between 0.7 and 1.0. $\sigma^2$ is the mean square displacement in the inter-atomic bond distance, representing the thermal and structural disorders in a crystal structure. $R_{Reff, i}$ is the inter-atomic distance obtained from the crystallography data and $\Delta R_i$ is the change in $R_{Reff, i}$ which is less than 0.1 Å [89]. $\Delta E_o$ is the difference in the energy scale between the theory and data.

Since the EXAFS signal is made of the interaction of sine waves, Eq. (5.6), the Fourier transform is used to extract the frequency of each wave at distance $R$ (Å) from the absorbing atom [89]. In fact, by taking the Fourier transform, one can move from energy space (k) to R-
space in order to find the amplitude and phase of each shell. The amplitude and the phase are related to the coordination number of atoms at each shell, the type of neighboring atoms, and the distance of atoms from the absorbing atom [89].

In order to model the experimental EXAFS signal by Eq. (5.6), one can evaluate $\Delta E_0$, $\Delta R$, $\sigma^2$, and $(CN, S^o_0)$ for each path by fitting the experimental EXAFS spectrum to Eq. (5.6). This is done by fitting the real and imaginary parts of the EXAFS Fourier transformed signal in available computer programs such as Artemis [77, 94]. The fitting consists of non-linear minimization of the sum of squared errors between data and the theory in R space [79]:

$$\chi^2 = \frac{N_{idp}}{N_{pts} \varepsilon^2 R^2} \sum_{i=1}^{N_{pts}} \left\{ \left| \text{Re}(\tilde{\chi}_{data}(R_i) - \tilde{\chi}_{theory}(R_i)) \right|^2 + \left| \text{Im}(\tilde{\chi}_{data}(R_i) - \tilde{\chi}_{theory}(R_i)) \right|^2 \right\}$$  \hspace{1cm} (5.7)

$\chi^2$ is the objective function which should not be mistaken as the EXAFS signal expressed by Eq. (5.6). Re and Im denote the real and imaginary parts of the Fourier transformed EXAFS signal. The tilde above $\chi$ shows the Fourier transformation of the EXAFS. $N_{pts}$ and $N_{idp}$ are the number of fitting parameters and the number of independent points, respectively. $N_{idp}$ is determined by:

$$N_{idp} = \frac{2 \Delta k \Delta R}{\pi} + 1$$  \hspace{1cm} (5.8)

$\Delta k$ is the range used to take the Fourier transform and $\Delta R$ is the range of the fitting in R space. $\varepsilon$ is the measurement uncertainty. It can be estimated by calculating the mean square of $\tilde{\chi}(R)$ in the range of 15-25 Å, assuming that there is no contribution from atomic shells in this range.

The goodness of EXAFS fits is usually expressed by R factor ($R_f$) which is the absolute misfit between the experimental data and the theory. $R_f$ is given by [79]:

$$R_f = \sqrt{\frac{\sum_{i=1}^{N_{pts}} \left[ \left| \text{Re}(\tilde{\chi}_{data}(R_i) - \tilde{\chi}_{theory}(R_i)) \right|^2 + \left| \text{Im}(\tilde{\chi}_{data}(R_i) - \tilde{\chi}_{theory}(R_i)) \right|^2 \right]}{\left[ \left| \text{Re}(\tilde{\chi}_{data}(R_i)) \right|^2 + \left| \text{Im}(\tilde{\chi}_{data}(R_i)) \right|^2 \right]}}$$  \hspace{1cm} (5.9)
Smaller values of $R_f$ show that fits are better performed. In order to compare the fitting of different models for the same data set, reduced $\chi^2$ is used. It is shown by $\chi^2_\nu$ and is given by:

$$\chi^2_\nu = \frac{\chi^2}{\nu}$$

(5.10)

$v$ is the degree of freedom of the data which is defined by the difference between $N_{pts}$ and $N_{idp}$. $\chi^2$ is given by Eq. (5.7). A fit can be said to be improved if $\chi^2_\nu$ is reduced by a factor of $1 + 2\sqrt{2/v}$ [79]. The precision of EXAFS in the determination of bond distances is about 0.02 Å while the resolution ($\Delta R$) can be estimated by [93]:

$$\Delta R = \frac{\pi}{2\Delta k}$$

(5.11)

Most often, the resolution of EXAFS fittings is between 0.1 to 0.2 Å.

### 5.2. EXAFS study of MnO$_2$ and Mn$_2$O$_3$ reference materials

As it can be seen in Eq. (5.4), the coordination numbers (CN) and the amplitude reduction factor ($S_o^2$) are 100% correlated. Therefore, it is not possible to determine the coordination number of oxygen and manganese shells in the catalysts without having a prior knowledge of $S_o^2$. $S_o^2$ is usually obtained from the EXAFS spectra of reference materials by fixing the coordination numbers from crystallography data. The $S_o^2$ is assumed to be the same in the data collection period of catalysts and reference materials. Therefore, $S_o^2$ obtained from the reference materials can be used as a fixed parameter for the EXAFS analysis of the catalysts to find the coordination numbers. Thus, EXAFS fittings of the reference materials (MnO$_2$ and Mn$_2$O$_3$) are explained in this section, resulting in $S_o^2$ which can be used in section 5.3 to investigate the changes in coordination numbers as the result of the change in Mn loading.
5.2.1. MnO$_2$

Unit cell dimensions of MnO$_2$ are presented in Table 4.2. The space group of MnO$_2$ is P42/m n m. A spatial structure of MnO$_2$ can be observed in Fig. 5.2 generated by PowderCell 2.4. MnO$_2$ has originally two oxygen atoms at 1.877 Å and four oxygen atoms at 1.891 Å [29]. But due to their close atomic positions, 6 oxygen atoms at an average distance of 1.887 Å are considered for MnO$_2$ EXAFS analysis. Results of EXAFS study of MnO$_2$ are presented in Fig. 5.3 and Table 5.1. The fitting is performed using k-weight of 1 in k and R ranges of 3-13 Å$^{-1}$ and 1-4 Å, respectively. Sine windows are used for k and R windows with $\Delta k$ and $\Delta R$ of 1 Å$^{-1}$ and 0 Å, respectively.
Three paths including the first oxygen shell and the first and second Mn shells are included in the model to simulate the EXAFS signal of MnO$_2$. The fitting results in an amplitude reduction factor of $0.68 \pm 0.06$. It has to be noted that coordination numbers (CN) are fixed from the crystallography information of MnO$_2$.

<table>
<thead>
<tr>
<th>Path</th>
<th>CN (fixed)</th>
<th>$\sigma^2 \times 10^{-3}$ Å$^2$</th>
<th>R (Å)</th>
<th>$E_o$ (eV)</th>
<th>$R_f$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>6</td>
<td>$1.8 \pm 1.0$</td>
<td>$1.872 \pm 0.007$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MnO$_2$</td>
<td>Mn-Mn</td>
<td>2</td>
<td>$2.871 \pm 0.013$</td>
<td>$0.15 \pm 1.0$</td>
<td>1.8</td>
</tr>
<tr>
<td></td>
<td>Mn-Mn</td>
<td>8</td>
<td>$3.424 \pm 0.010$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5.2. Unit cell structure of MnO$_2$. Numbers are in Å.
Fig. 5.3. Magnitude (a) and real part (b) of the EXAFS spectra of MnO₂ reference material. Solid lines represent the data and the symbols are the fittings.

Fig. 5.3 shows the magnitude and real part of MnO₂ EXAFS and the fitting results. The first peak in Fig. 5.3(a) is due to the first shell oxygen atoms. The second peak is due to the first Mn shell while the third peak can be related to farther atomic shells. It can be seen that the fitting curves follow closely the data, resulting in absolute percentage misfit ($R_f$) of 1.8%.

5.2.2. Mn₂O₃

The unit cell dimensions of Mn₂O₃ are available in Table 4.2 with the space group of Ia3. Mn₂O₃ has a more complicated unit cell structure than MnO₂. There are two Mn sites in Mn₂O₃ with occupancy of 25 and 75%. The crystal structure information of Mn₂O₃ up to ca. 3.5 Å is
given in Table 5.2. Mn site 1 (Mn1) with 25% occupancy, has 6 oxygen atoms at 1.993 Å while Mn site 2 (Mn2) has three sets of oxygen atoms at different bond lengths. There are 2 short oxygen bonds at 1.899 Å and two long oxygen bonds at 2.248 Å in Mn site 2. In addition to these 4 oxygen atoms, there are two oxygen atoms at 1.985 Å in Mn site 2 which are extremely close to the oxygen bond length in Mn site 1 at 1.993 Å. The spatial schematic of the two Mn sites can be seen in Fig. 5.4 produced by PowderCell 2.4.

Table 5.2. Crystal structure of Mn₂O₃

<table>
<thead>
<tr>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>6</td>
<td>1.993</td>
<td>Mn-O</td>
<td>2</td>
<td>1.899</td>
<td>Mn-O</td>
<td>1.5</td>
<td>1.899</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>3.104</td>
<td>Mn-O</td>
<td>2</td>
<td>1.985</td>
<td>Mn-O</td>
<td>3</td>
<td>1.987</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>3.565</td>
<td>Mn-O</td>
<td>2</td>
<td>2.248</td>
<td>Mn-O</td>
<td>1.5</td>
<td>2.248</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.115</td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.112</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.575</td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.572</td>
</tr>
</tbody>
</table>

*Mn₂O₃*

<table>
<thead>
<tr>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
<th>Bond</th>
<th>CN</th>
<th>R_{eff} (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>6</td>
<td>1.993</td>
<td>Mn-O</td>
<td>2</td>
<td>1.899</td>
<td>Mn-O</td>
<td>1.5</td>
<td>1.899</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>3.104</td>
<td>Mn-O</td>
<td>2</td>
<td>1.985</td>
<td>Mn-O</td>
<td>3</td>
<td>1.987</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>3.565</td>
<td>Mn-O</td>
<td>2</td>
<td>2.248</td>
<td>Mn-O</td>
<td>1.5</td>
<td>2.248</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.115</td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.112</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.575</td>
<td>Mn-Mn</td>
<td>6</td>
<td>3.572</td>
</tr>
</tbody>
</table>

*a* Averaged based on Mn site occupancies.

*b* Obtained from crystallography data.
The presence of two Mn sites in Mn$_2$O$_3$ makes the EXAFS analysis complicated due to the close atomic coordinates of oxygen in the two sites. There are two methods which can be used to model the EXAFS signal of Mn$_2$O$_3$. The first method simplifies the structure of Mn$_2$O$_3$ by averaging the two Mn sites based on their site occupancies. This integrates the Mn sites into one single site which can be used for EXAFS analysis. The second method is Bond Valence Model which is based on Pauling’s 2nd rule of electrostatic valence [95]. The Bond Valence Model relates the oxidation state of Mn atoms to the CN of oxygen atoms, adding constrains on the EXAFS fitting procedure. This allows modeling the structure of Mn$_2$O$_3$ in a more general way with no simplifying assumption. The remaining of this section represents and compares the results of EXAFS fitting of Mn$_2$O$_3$ reference material by the two mentioned methods in order to
model complicated crystals such as Mn$_2$O$_3$. EXAFS fitting are carried out with $k$-weight of 1 for the Mn$_2$O$_3$ reference material. Sine window function with $\Delta k$ and $\Delta R$ of 1 Å$^{-1}$ and 0 Å is used, respectively. The Fourier transforms are taken between 3-13 Å$^{-1}$. EXAFS Fittings are performed in R space from 1 to 3.7 Å.

**Averaging method:**

The simplified structure of Mn$_2$O$_3$ by averaging Mn site 1 (Mn1) and Mn site 2 (Mn2) is shown in the right column of Table 5.2. The averaging method results virtually in 1.5, 3, and 1.5 oxygen atoms at ca. 1.90, 1.99, and 2.25 Å, respectively. First and second shells of Mn atoms are located at ca. 3.11 and 3.57 Å, respectively. Table 5.3 contains the result of EXAFS fitting of Mn$_2$O$_3$ based on the averaging method and Fig. 5.5 shows the magnitude and the real part of the Fourier transform of the data and the fitting.

Table 5.3. EXAFS fitting result of Mn$_2$O$_3$ based on the averaging method

<table>
<thead>
<tr>
<th>Path</th>
<th>CN</th>
<th>$\sigma^2$ ($\times 10^{-3}$ Å$^2$)</th>
<th>R (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>1.5</td>
<td>6.8 ± 2.1</td>
<td>1.883 ± 0.012</td>
</tr>
<tr>
<td>Mn-O</td>
<td>3</td>
<td>6.8 ± 2.1</td>
<td>1.969 ± 0.012</td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.5</td>
<td>6.8 ± 2.1</td>
<td>2.233 ± 0.012</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>7.7 ± 1.5</td>
<td>3.113 ± 0.011</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>18.9 ± 4.9</td>
<td>3.617 ± 0.036</td>
</tr>
</tbody>
</table>
Referring to Table 5.3, it can be seen that the averaging method results in an amplitude reduction factor \( S_o^2 \) of 0.83 ± 0.12 and locates the first shell oxygen atoms at ca. 1.88, 1.97, and 2.33 Å with CN of 1.5, 3, and 1.5, respectively. Two Mn atoms are also resolved at the second and third shells at ca. 3.11 and 3.62 Å, respectively. There are almost 17 independent points in the fitting range and 8 fitting parameters in the averaging method, resulting in 9 degree of freedom.

**Bond Valence Model:**

In order to construct a more general model and also to check the validity of the averaging method, the Bond Valence Model is used to model the EXAFS of Mn\(_2\)O\(_3\) [95]. According to this
model which is based on Pauling’s 2\(^{nd}\) rule of electrostatic valence, formal oxidation state of the cation \(i\) is equal to the sum of valence of the bond between the cation \(i\) and surrounding anions \(j\):

\[
V_i = \sum_j s_{ij}
\]  

(5.12)

in which the bond valence \((s_{ij})\) is defined by:

\[
s_{ij} = \exp\left(\frac{R'_{ij} - R_{ij}}{b}\right)
\]  

(5.13)

\(R'_{ij}\) and \(b\) are empirical parameters and \(R_{ij}\) is the inter-atomic distance between the cation \(i\) and the anion \(j\) which can be coupled to EXAFS models. It has been shown that \(b\) has a universal value of 0.37 [95]. If there is only one type of anions around the cation, \(R_{ij}\) can be extracted from Eq. (5.13) and expressed as:

\[
R_{ij} = R'_{ij} + 0.37\ln\left(\frac{CN_{ij}}{V_i}\right)
\]  

(5.14)

Where \(CN_{ij}\) is the coordination number of the anion \(j\) around the cation \(i\) at distance \(R_{ij}\). Eq. (5.14) correlates the inter-atomic bond length to the coordination number of anions and oxidation state of the cation. This allows using a prior knowledge of the cation oxidation state, available from XANES analysis, in the EXAFS fitting which results in the reduction of the number of fitting parameters. Even though it is possible to directly use Eq. (5.14) in the EXAFS analysis to remove \(\Delta R\) or \(CN\) from the fitting parameters, it is recommended to use Eq. (5.12) as a restraint to model EXFAS signals [95]. The reason relies on the inaccuracy of Eq. (5.12) and (5.13), leading to unreliable fitting parameters in EXAFS, when their exact definitions are used as constraints. Therefore, it is suggested to use the Bond Valence Model as a restraint in the EXAFS fitting of \(\text{Mn}_2\text{O}_3\). This adds a penalty term on the EXAFS minimization objective function to keep the oxidation state of Mn cations close to their formal valence which is +3.
Considering the value of $R'_ij$ (1.760) for Mn$^{3+}$ in Mn$_2$O$_3$ [96], application of Eq. (5.12) and (5.13) results in the following equations for the two Mn sites:

\[
V_{\text{Mn,sitel}} = \text{CN}_O \exp\left(\frac{1.76 - R_{O,site1}}{0.37}\right)
\]  
(5.15)

\[
V_{\text{Mn,site2}} = \left(\frac{\text{CN}_O}{3}\right) \exp\left(\frac{1.76 - R_{O,site2}}{0.37}\right) + \left(\frac{\text{CN}_O}{3}\right) \exp\left(\frac{1.76 - R_{O,site1}}{0.37}\right) + \left(\frac{\text{CN}_O}{3}\right) \exp\left(\frac{1.76 - R_{O2,site2}}{0.37}\right)
\]  
(5.16)

Where $\text{CN}_O$ represents the total number of oxygen atoms at the first shell of Mn$_2$O$_3$ which can be set at 6. $R_{O,site}$ is used to represent the inter-atomic distance of the oxygen atoms at 1.993 and 1.985 Å in the first and second Mn sites, respectively. The distances of the other two oxygen atoms at 1.899 and 2.248 Å in Mn site 2 (Mn2) are denoted by $R_{O1,site2}$ and $R_{O2,site2}$, respectively. Eq. (5.15) and (5.16) are defined as restraints in Artemis with restricted values between 0 and 3 using a moderate penalty coefficient. It has to be noted that no simplification is performed in EXAFS fitting of Mn$_2$O$_3$ by the Bond Valence Model, allowing incorporation of both Mn sites in the model. The amplitude reduction factor is multiplied by Mn site occupancies in order to consider paths in each Mn sites.

Results of the fitting of Mn$_2$O$_3$ using the Bond Valence Model are shown in Table 5.4 and Fig. 5.6. The Bond Valence Model gives an amplitude reduction factor of 0.88 ± 0.16 with roughly 17 independent points, 10 fitting variables and 7 degree of freedom. There are 6 oxygen atoms at ca. 1.99 Å from Mn atom in site one whereas three sets of two oxygen atoms are located at ca. 1.87, 1.98, and 2.25 Å around the second Mn site. The first Mn shell is found around the first and second Mn sites at ca. 3.11 and 3.12 Å, respectively with CN of 6. The second Mn shell also has 6 Mn atoms which are located at ca. 3.61 and 3.62 Å away from Mn site 1 and site 2,
respectively. CNs reported in Table 5.4 are fixed parameters obtained from crystallography data of Mn$_2$O$_3$.

Table 5.4. EXAFS fitting result of Mn$_2$O$_3$ based on the Bond Valence Model

<table>
<thead>
<tr>
<th>Path</th>
<th>CN</th>
<th>$\sigma^2$ ($\times 10^{-3}$ Å$^2$)</th>
<th>R (Å)</th>
<th>Path</th>
<th>CN</th>
<th>$\sigma^2$ ($\times 10^{-3}$ Å$^2$)</th>
<th>R (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>6</td>
<td>6.5 ± 2.6</td>
<td>1.986 ± 0.031</td>
<td>Mn-O</td>
<td>2</td>
<td>6.5 ± 2.6</td>
<td>1.870 ± 0.064</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>8.3 ± 1.9</td>
<td>3.110 ± 0.012</td>
<td>Mn-O</td>
<td>2</td>
<td>6.5 ± 2.6</td>
<td>1.977 ± 0.031</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>21.0 ± 6.1</td>
<td>3.606 ± 0.041</td>
<td>Mn-O</td>
<td>2</td>
<td>6.5 ± 2.6</td>
<td>2.251 ± 0.035</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>6</td>
<td>8.3 ± 1.9</td>
<td>3.118 ± 0.012</td>
<td>Mn-Mn</td>
<td>6</td>
<td>21.0 ± 6.1</td>
<td>3.616 ± 0.041</td>
</tr>
</tbody>
</table>

By comparing the results of the two methods in Table 5.3 and 5.4, it can be seen that the obtained parameters are very close to each other. For example, $\sigma^2$ of the first shell of oxygen atoms obtained by the averaging method is $6.8 \times 10^{-3}$ Å$^2$ while the one from the Bond Valence Model is $6.5 \times 10^{-3}$ Å$^2$. The same similarities can also be seen in $\sigma^2$ of the Mn shells, the amplitude reduction factor (0.83 versus 0.88) and inter-atomic distances. The $R_f$ (percentage of absolute misfit between the data and theory) of the averaging method and the Bond Valence Model are 1.9 and 1.7%, respectively. The reduced chi-square values, 3416 for the averaging method and 4573 for the Bond Valence Model, are also close to each other, showing that the two methods have almost the same statistical significance. This shows that both models can be used for EXAFS analysis of Mn$_2$O$_3$ containing catalysts. However, the Bond Valence Model can be
considered as a general model for the EXAFS fitting of complicated crystal structures such as Mn$_2$O$_3$ since it does not impose any simplification on the unit cell structure. Nevertheless, the Bond Valence Model proves that the averaging method results in feasible and valid solutions and can be used to model the EXAFS signal of oxides of manganese such as Mn$_2$O$_3$ and Mn$_3$O$_4$. An average value of 0.85 is assumed for the amplitude reduction factor of Mn$_2$O$_3$ from the result of this section. The averaging method will be used in order to study the EXAFS of alumina supported manganese oxides due to less complexity of this method.

![EXAFS spectra](image)

**Fig. 5.6.** Magnitude (a) and real part (b) of the EXAFS spectra of Mn$_2$O$_3$ reference material based on the Bond Valence Model. Solid lines represent the data and the symbols are the fittings.

It has to mentioned that the results of EXAFS studies reported in Table 5.3 and 5.4, are comparable with the EXAFS study of Mn$_2$O$_3$ (bixbyite) performed by Longo et al. [86].
Basically, the authors have modeled the theoretical EXAFS signal of Mn$_2$O$_3$ by considering 6 oxygen atoms at ca. 1.88 Å, 6 manganese atoms at ca. 3.12 Å and 6 manganese atoms at ca. 3.60 Å. They have fixed the oxygen and manganese coordination numbers along with a predetermined value of 0.74 for $S_0^2$. Their model has resulted in $\sigma^2$ of $7 \times 10^{-3}$ Å$^2$ for the first oxygen shell which is very close to $\sigma^2$ reported in this work for Mn$_2$O$_3$ reference material, i.e. $6.8 \times 10^{-3}$ Å$^2$ and $6.5 \times 10^{-3}$ Å$^2$ for the averaging method and the Bond Valence Model, respectively.

5.3. *Characterization of MnO$_x$/γ-alumina (1, 5, 10 and 20%)*

Four loadings of Mn (1, 5, 10, and 20%) are used to prepare MnO$_x$/γ-alumina with different manganese loadings. Catalysts are characterized with X-ray diffraction (XRD) and X-ray absorption spectroscopy (XAS). More emphasis is put on XAS including XANES and EXAFS in order to find the relationship among the electronic and structural properties of manganese oxides and Mn loading.

5.3.1. *X-ray diffraction (XRD)*

Fig. 5.7 shows XRD of the catalysts and the reference materials including Mn$_2$O$_3$ and MnO$_2$. Peaks corresponding to the reference materials are marked with two dashed lines in order to track the peak locations on XRD of the catalysts. The catalyst with Mn loading of 1% does not show any diffraction peaks due to high dispersion of manganese oxides on alumina. But MnO$_x$/γ-alumina (5%) shows peaks, matching the spectra of Mn$_2$O$_3$. More strong peaks of Mn$_2$O$_3$ can be seen in XRD of MnO$_x$/γ-alumina (10%) along with weak peaks of MnO$_2$ showing that MnO$_2$ is coexisting with Mn$_2$O$_3$ in this catalyst. With the increase of Mn loading to 20%, more intense peaks of MnO$_2$ are observed while Mn$_2$O$_3$ peaks become smaller in XRD spectra of MnO$_x$/γ-
alumina (20%). This shows that by increasing Mn loading, different phases of manganese oxides are formed. At lower loading up to 10%, the dominant phase is Mn$_2$O$_3$ while at higher loadings, both MnO$_2$ and Mn$_2$O$_3$ phases coexist.

![XRD spectra](image)

Fig. 5.7. XRD of reference materials and catalysts, (a) Mn$_2$O$_3$, (b) MnO$_2$, (C), MnO$_x$/γ-alumina (1%), (d) MnO$_x$/γ-alumina (5%), (e) MnO$_x$/γ-alumina (10%), and (f) MnO$_x$/γ-alumina (20%).

5.3.2. X-ray absorption near structure (XANES)

Mn K-edge XANES spectra of the reference materials (MnO$_2$ and Mn$_2$O$_3$) and the catalysts are shown in Fig. 5.8. It can be seen that the rising part of the spectra of the catalysts with 1 and 5% Mn loading are located at the left side of Mn$_2$O$_3$. XANES Spectrum of MnO$_x$/γ-alumina (10%) is closely located at the right side of Mn$_2$O$_3$, suggesting that this catalyst is
mainly composed of Mn₂O₃. The XANES spectra of the catalyst with 20% are well located between that of Mn₂O₃ and MnO₂, indicating that MnOₓ/γ-alumina (20%) has considerable amount of MnO₂ in addition to Mn₂O₃.

![XANES spectra](image)

Fig. 5.8. Mn K-edge XANES spectra of the samples, (a) MnOₓ/γ-alumina (1%), (b) MnOₓ/γ-alumina (5%), (c) MnOₓ/γ-alumina (10%), (d) MnOₓ/γ-alumina (20%), (e) MnO₂, and (f) Mn₂O₃.

Mn K-edge absorption energy (Eₒ) of the reference materials are 6548.1 and 6552.7 eV for Mn₂O₃ and MnO₂, respectively. Absorption energies of the catalysts are presented in Table 5.5. MnOₓ/γ-alumina (1 and 5%) catalysts have absorption energies less than that of Mn₂O₃ most probably due to the presence of highly dispersed manganese atoms and incomplete growth of Mn₂O₃ unit cells. On the other hand, catalysts with Mn loadings of 10 and 20% have absorption
energies higher than that of Mn$_2$O$_3$. The absorption energy of MnO$_x$/γ-alumina (10%) is 0.3 eV higher than $E_o$ of Mn$_2$O$_3$ while the sample with 20% Mn has an absorption energy at 6552.7 eV which is the same as MnO$_2$ absorption energy.

Table 5.5. Mn K-edge absorption energies of the catalysts

<table>
<thead>
<tr>
<th></th>
<th>MnO$_x$/γ-alumina (1%)</th>
<th>MnO$_x$/γ-alumina (5%)</th>
<th>MnO$_x$/γ-alumina (10%)</th>
<th>MnO$_x$/γ-alumina (20%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn K-edge $E_o$ (eV)</td>
<td>6547.4</td>
<td>6547.8</td>
<td>6548.4</td>
<td>6552.7</td>
</tr>
</tbody>
</table>

Linear combination fitting of the catalysts (10 and 20% Mn loading) with the reference materials (MnO$_2$ and Mn$_2$O$_3$) shows that the weight percent of MnO$_2$ increases from 9.0 to 55.3% by increasing Mn loading from 10 to 20%. Meanwhile, the fraction of Mn$_2$O$_3$ drops from 91.0 to 44.7% from MnO$_x$/γ-alumina (10%) to MnO$_x$/γ-alumina (20%). This is in agreement with the XRD results, showing that more MnO$_2$ is formed in MnO$_x$/γ-alumina by increasing the loading from 10 to 20%.

5.3.3. Extended X-ray absorption fine structure (EXAFS)

Fig. 5.9 shows Mn K-edge EXAFS oscillation of the samples. The EXAFS spectra of the catalysts with 5 and 10% loading matches the spectra of Mn$_2$O$_3$ reference material. On the other hand, EXAFS oscillation of 1% Mn loaded catalyst is different from the spectra of MnO$_2$ and Mn$_2$O$_3$ due to the high dispersion of Mn. EXAFS of the catalyst with 20% Mn is similar to the spectra of MnO$_2$ since the major fraction of manganese oxides in this catalyst is composed of MnO$_2$. 
Fig. 5.9. Mn K-edge EXAFS spectra of the samples, (a) MnO$_x$/$\gamma$-alumina (1%), (b) MnO$_x$/$\gamma$-alumina (5%), (c), MnO$_x$/$\gamma$-alumina (10%), (d) MnO$_x$/$\gamma$-alumina (20%), (e) MnO$_2$, and (f) Mn$_2$O$_3$.

Using the model explained in section 5.2, $S^2_o$ of MnO$_2$ and Mn$_2$O$_3$ were estimated to be 0.68 and 0.85, respectively. The results of the EXAFS fitting of the catalysts are summarized in Table 5.6 by the averaging method. Fig. 5.10 and 5.11 depict the magnitude and the real part of the Fourier transform (FT) of Mn K-edge spectra of the catalysts, respectively. Fig. 5.10 and 5.11 also contain the magnitude and the real part of the FT of MnO$_2$ and Mn$_2$O$_3$ reference materials in order to provide a guideline for comparison between the catalysts and the reference materials.
Table 5.6. EXAFS fitting results of the catalysts by the averaging method

<table>
<thead>
<tr>
<th>Path</th>
<th>CN</th>
<th>$\sigma^2$ ($\times 10^{-3}$ Å$^2$)</th>
<th>R (Å)</th>
<th>$E_o$ (eV)</th>
<th>$R_t$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>1.7 ± 0.5</td>
<td>1.886 ± 0.017</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.2 ± 0.2</td>
<td>7.0 ± 1.6</td>
<td>1.985 ± 0.017</td>
<td>0.2 ± 1.6</td>
<td>0.9</td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.4 ± 0.3</td>
<td>2.236 ± 0.017</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>1.0 ± 0.5</td>
<td>17.4 ± 9.3</td>
<td>3.105 ± 0.026</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.7 ± 0.6</td>
<td>1.904 ± 0.018</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-O</td>
<td>2.1 ± 0.5</td>
<td>5.4 ± 1.8</td>
<td>1.990 ± 0.018</td>
<td>4.5 ± 1.1</td>
<td>2.4</td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.4 ± 0.1</td>
<td>2.254 ± 0.018</td>
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<tr>
<td>Mn-Mn</td>
<td>4.2 ± 1.2</td>
<td>10.5 ± 3.0</td>
<td>3.117 ± 0.014</td>
<td></td>
<td></td>
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<tr>
<td>Mn-O</td>
<td>1.7 ± 0.4</td>
<td>1.894 ± 0.015</td>
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</tr>
<tr>
<td>Mn-O</td>
<td>2.5 ± 0.5</td>
<td>4.6 ± 1.6</td>
<td>1.985 ± 0.015</td>
<td>5.0 ± 0.9</td>
<td>1.6</td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.4 ± 0.1</td>
<td>2.249 ± 0.015</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>5.9 ± 1.1</td>
<td>9.6 ± 1.9</td>
<td>3.127 ± 0.011</td>
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<tr>
<td>Mn-O</td>
<td>3.5 ± 0.5</td>
<td>1.884 ± 0.009</td>
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<tr>
<td>Mn-O</td>
<td>1.1 ± 0.1</td>
<td>3.3 ± 1.3</td>
<td>1.970 ± 0.009</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-O</td>
<td>0.9 ± 0.1</td>
<td>2.234 ± 0.009</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>1.4 ± 0.2</td>
<td>2.882 ± 0.012</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>2.1 ± 0.1</td>
<td>4.7 ± 1.2</td>
<td>3.115 ± 0.012</td>
<td></td>
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</tr>
<tr>
<td>Mn-Mn</td>
<td>4.0 ± 0.3</td>
<td>3.435 ± 0.012</td>
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</tr>
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</table>
Fig. 5.10. Magnitude of the Fourier transform of Mn K-edge data (solid lines) and fitting (symbols), (a) MnOₓ/γ-alumina (1%), (b) MnOₓ/γ-alumina (5%), (c) MnOₓ/γ-alumina (10%), (d) MnOₓ/γ-alumina (20%), (e) MnO₂, and (f) Mn₂O₃.
Fig. 5.11. Real part of the Fourier transform of Mn $K$-edge data (solid lines) and fitting (symbols), (a) MnO$_x$/γ-alumina (1%), (b) MnO$_x$/γ-alumina (5%), (c), MnO$_x$/γ-alumina (10%), (d) MnO$_x$/γ-alumina (20%), (e) MnO$_2$, and (f) Mn$_2$O$_3$.

Starting with MnO$_x$/γ-alumina (1%) in Table 5.6, it can be seen that CN of oxygen and manganese atoms at the first and second shells are smaller than the proposed model for Mn$_2$O$_3$ with one single site in Table 5.2. Low oxygen and manganese CNs can be observed by considering the magnitude and the real part of the FT of MnO$_x$/γ-alumina (1%) shown in Fig. 5.10 and Fig. 5.11, respectively. The FT magnitude and the real part of the first shell oxygen atoms are smaller than that of the catalysts with higher loadings. Mn also shows a very broad weak peak at the second shell of this catalyst, confirming the low CN of manganese.
By increasing the loading to 5%, oxygen CN at ca. 1.99 Å and manganese CN at ca. 3.11 Å increase to 2.1 and 4.2, respectively. Oxygen CNs are basically close to the values considered for \( \text{Mn}_2\text{O}_3 \) in Table 5.2 (1.5 at 1.899 Å, 3 at 1.987 Å and 1.5 at 2.248 Å) while Mn CN is smaller than 6 which is the expected CN of Mn at the second shell of \( \text{Mn}_2\text{O}_3 \) crystal.

Increasing loading to 10% does not significantly change oxygen CNs while it results in Mn CN of 5.9 which is very close to Mn CN in the second shell of \( \text{Mn}_2\text{O}_3 \). Fig. 5.10 (b) and 5.10 (c) also qualitatively show that the intensity of oxygen at the first shell remains unchanged by increasing the loading from 5% to 10%. However, Mn signal intensity increases at the second shell, implying higher CN of Mn in \( \text{MnO}_x/\gamma\text{-alumina (10%)} \) than \( \text{MnO}_x/\gamma\text{-alumina (5%)} \). It should be noted that the EXAFS of \( \text{MnO}_x/\gamma\text{-alumina (10%)} \) was modeled based on the structure of \( \text{Mn}_2\text{O}_3 \) and no contribution from \( \text{MnO}_2 \) is considered in the fitting of this catalyst. This is mainly because of the fact that only 9.0% of \( \text{MnO}_x \) is in the form of \( \text{MnO}_2 \) in \( \text{MnO}_x/\gamma\text{-alumina (10%)} \). This results in the EXAFS oscillation of this catalyst to be similar to that of \( \text{Mn}_2\text{O}_3 \) shown in Fig. 5.9, allowing fitting EXAFS of \( \text{MnO}_x/\gamma\text{-alumina (10%)} \) by \( \text{Mn}_2\text{O}_3 \) structure.

In order to model the EXAFS spectra of the catalyst with 20% Mn loading which mostly contains \( \text{MnO}_2 \) (55.3%), two Mn backscattering atoms from \( \text{MnO}_2 \) at ca. 2.87 and 3.42 Å are added to the model in order to accommodate \( \text{MnO}_2 \) contributions. Comparing the common paths (Mn-O at ca. 1.90, 1.99, and 2.25 Å and Mn-Mn at ca. 3.11 Å) in \( \text{MnO}_x/\gamma\text{-alumina (10%)} \) and \( \text{MnO}_x/\gamma\text{-alumina (20%)} \), some changes can be seen in coordination numbers due to the coexistence of \( \text{MnO}_2 \) and \( \text{Mn}_2\text{O}_3 \). By increase of Mn loading from 10 to 20%, Oxygen CN at ca. 1.90 Å increases from 1.7 to 3.5. On the other hand, oxygen CNs of \( \text{MnO}_x/\gamma\text{-alumina (20%)} \) at ca. 1.99 and 2.25 Å are 1.1 and 0.9, respectively which are smaller than their corresponding values in \( \text{MnO}_x/\gamma\text{-alumina (10%)} \), i.e. 2.5 and 1.4 at ca. 1.99 and 2.25 Å, respectively. CN of Mn
at ca. 3.11 decreases from 5.9 in MnO_x/γ-alumina (10%) to 2.1 in MnO_x/γ-alumina (20%). In addition to these changes, two new manganese atoms were resolved at ca. 2.88 Å and 3.44 Å with CN of 1.4 and 4.0 which are the coordinates of manganese atoms in the structure of MnO_2.

If one makes an average over oxygen and manganese CNs based on the weight percent of MnO_2 (55.3%) and Mn_2O_3 (44.7%) in MnO_x/γ-alumina (20%) and uses CNs available in Table 5.1 and Table 5.2 for MnO_2 and Mn_2O_3 (with one single Mn site), a simplified model can be obtained for the catalysts with 20% Mn loading which is shown in Table 5.7. Based on this approach, one would expect 4.0, 1.4, and 0.7 oxygen atoms at ca. 1.90, 1.99, and 2.25 Å, respectively. In addition, this method results in 1.1, 2.7 and 4.4 of manganese atoms at ca. 2.87, 3.11, and 3.42 Å, respectively. It can be seen that the result of EXAFS fitting of MnO_x/γ-alumina (20%) reported in Table 5.6, follows the trend of CN changes shown in Table 5.7, implying that EXAFS fitting results are fairly reasonable.

Table 5.7. Averaged coordination numbers of MnO_x/γ-alumina (20%)

<table>
<thead>
<tr>
<th>Path</th>
<th>R_{eff} (Å)</th>
<th>Coordination number (CN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn-O</td>
<td>1.892</td>
<td>4.0</td>
</tr>
<tr>
<td>Mn-O</td>
<td>1.987</td>
<td>1.4</td>
</tr>
<tr>
<td>Mn-O</td>
<td>2.248</td>
<td>0.7</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>2.871</td>
<td>1.1</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>3.112</td>
<td>2.7</td>
</tr>
<tr>
<td>Mn-Mn</td>
<td>3.424</td>
<td>4.4</td>
</tr>
</tbody>
</table>
R factors (R_t) reported in Table 5.6 represent the absolute percentage misfit between the data and theory. All fittings show low R_t values around 2% which shows that the models have been fitted closely to the EXAFS data.

5.4. Relationship between the structure and the activity of the catalysts

Four loadings of Mn (1, 5, 10, and 20%) are used for the oxidation of toluene by ozone at room temperature and higher temperatures up to 80 °C. 0.2 g of the catalysts is used to obtain the activities. Concentrations of toluene and ozone are 120 and 1050 ppm, respectively. The flow rates of toluene/nitrogen and ozone/oxygen streams are 400 and 600 ml min⁻¹, respectively. Toluene and ozone conversion at room temperature are shown in Fig. 5.12 and Fig. 5.13, respectively. It can be seen that the activity of the catalysts decreases with the increases of Mn loading. MnOₓ/γ-alumina (1%) shows the highest activity followed by the catalysts with 5, 10 and 20% loadings. COₓ (CO and CO₂) are the major by-product of the reaction. All catalysts lose their activity during the reaction course in terms of toluene and ozone conversions in accordance with the activity of MnOₓ/γ-alumina (10%) reported in Chapter 4. CHNS analysis is performed on the catalysts after running the reaction for 150 minutes at room temperature. The results, shown in Table 5.8, indicate that all the four catalysts gain considerable amount of carbon which can be considered as the reason for the catalyst deactivation. Table 5.8 also shows that the most active catalyst, MnOₓ/γ-alumina (1%), gains higher carbon content than the other catalysts. The amount of accumulated carbon decreases with the increase of Mn loading in all catalysts. This trend can be explained with respect to the fact that the nature of the ongoing reaction at room temperature is mostly partial oxidation of toluene to carbonaceous compounds. Therefore, the catalysts with lower loading and higher activity produce more accumulated carbon species at room temperature.
Fig. 5.12. Toluene conversion at room temperature, (a) MnO$_x$/$\gamma$-alumina (1%), (b) MnO$_x$/$\gamma$-alumina (5%), (c) MnO$_x$/$\gamma$-alumina (10%), and (d) MnO$_x$/$\gamma$-alumina (20%).

Fig. 5.13. Ozone conversion at room temperature, (a) MnO$_x$/$\gamma$-alumina (1%), (b) MnO$_x$/$\gamma$-alumina (5%), (c) MnO$_x$/$\gamma$-alumina (10%), and (d) MnO$_x$/$\gamma$-alumina (20%).
Table 5.8. Accumulated carbon content on the catalysts after 150 minutes reaction at room temperature

<table>
<thead>
<tr>
<th></th>
<th>MnO₅/γ-alumina (1%)</th>
<th>MnO₅/γ-alumina (5%)</th>
<th>MnO₅/γ-alumina (10%)</th>
<th>MnO₅/γ-alumina (20%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon content (wt%)</td>
<td>11.16</td>
<td>11.01</td>
<td>10.25</td>
<td>9.00</td>
</tr>
</tbody>
</table>

Fourier transform infrared (FT-IR) is used to study the surface of the deactivated catalysts. FT-IR spectra of fresh γ-alumina and the catalysts after 150 minutes of the reaction at room temperature are shown in Fig. 5.14.

![FT-IR spectra](image)

Fig. 5.14. FT-IR spectra of the deactivated catalysts, (a) fresh γ-alumina, (b) MnO₅/γ-alumina (1%), (c) MnO₅/γ-alumina (5%), (d) MnO₅/γ-alumina (10%), and (e) MnO₅/γ-alumina (20%).
Spectra of fresh $\gamma$-alumina not used in the reactor, Fig. 5.14 (a), show peaks at 1622, 2350 cm$^{-1}$ and a broad signal between 2500 and 3800 cm$^{-1}$ which are due to adsorbed water, CO$_2$, and hydroxyl groups on alumina, respectively. All the four catalysts show new bands at 1410 and 1718 cm$^{-1}$ which are indicative of C-O-H bending of alcohols and carbonyl group (C=O) stretch of carboxylic acids, respectively [97]. In addition, a broad peak in the range 2500 to 3800 cm$^{-1}$ is observed on all catalysts which can be due to the overlapping of OH stretching of carboxylic acids and alcohols with that of adsorbed water [6]. The intensity of these new peaks (1410, 1718, and 2500-3800 cm$^{-1}$) becomes smaller with the increase of Mn loading, implying that the amount of deposited alcohols and carboxylic acids has a reverse relationship with the manganese loading. This is also qualitatively in agreement with the result of CHNS analysis suggesting that the order of carbon content of the used catalysts has reverse relationship with the Mn loading. Acetic acid and oxalic acid, benzene, and maleic anhydride were identified as the minor by-products of the reaction in Chapter 3 [98]. The presence of acetic acid and oxalic acid also supports the identification of carboxylic acids on the catalysts at room temperature due to the incomplete oxidation of toluene.

Results (toluene and ozone conversion versus temperature) of high temperature activity of the catalysts up to 80 °C are shown in Fig. 5.15 and Fig. 5.16. Activity orders are similar to the room temperature activities, indicating that lower Mn loadings result in higher conversion of toluene and ozone. It can be seen that increasing the temperature from room temperature (22 °C) enhances toluene and ozone conversions up to 80 °C. All the four catalysts show the same activity at 80 °C, at which maximum conversion of toluene occurs. Increasing the temperature from 80 to 100 °C, data not shown, decreases toluene conversion to some extent while the conversion of ozone remains constant at 100%.
Fig. 5.15. Toluene conversion versus temperature, (a) MnO$_x$/γ-alumina (1%), (b) MnO$_x$/γ-alumina (5%), (c) MnO$_x$/γ-alumina (10%), and (d) MnO$_x$/γ-alumina (20%).

Fig. 5.16. Ozone conversion versus temperature, (a) MnO$_x$/γ-alumina (1%), (b) MnO$_x$/γ-alumina (5%), (c) MnO$_x$/γ-alumina (10%), and (d) MnO$_x$/γ-alumina (20%).
The overall observation of the activities at room and higher temperatures suggests that catalysts with lower Mn loadings are more active for the oxidation of toluene by ozone. Einaga et al. have studied the effect of Mn loading in the oxidation of benzene by ozone at room temperature over MnOx/γ-alumina [55]. They reported that catalysts with lower loadings (1 to 7.5%) have slightly higher activities and isolated Mn atoms in lower loadings and aggregated Mn3O4 clusters in higher loadings (10 to 15%) show almost similar activities. On the other hand, Reed et al. have showed that higher loadings of Mn on silica result in higher oxidation rate of acetone by ozone [29]. They proposed that lower oxidation states of Mn and more adjacent Mn sites in catalysts with higher Mn loadings are two factors, controlling the rate of acetone oxidation by ozone. They have also mentioned that the rate controlling step of acetone ozonation is the oxidation of the catalyst by ozone and not the oxidation of acetone by the catalyst.

As it was mentioned in Chapter 2, the mechanism of oxidation of VOCs by ozone is based on ozone decomposition expressed by Eq. (5.17) to Eq. (5.19), generating atomic oxygen to react with adsorbed VOCs:

\[ O_3 + Mn \rightarrow O_2 + O-Mn \]  
\[ O_3 + O-Mn \rightarrow O_2 + O_2-Mn \]  
\[ O_2-Mn \rightarrow O_2 + Mn \]  

It is believed that higher dispersion of Mn atoms in catalysts with lower loading levels decreases the oxidation state of Mn. As the oxidation state of Mn decreases, the ability of Mn atoms in transferring electrons to ozone in Eq. (5.17) \( O_3 + Mn^{n+} \rightarrow O_2 + O^{2-} + Mn^{(n+2)+} \) increases, leading to faster decomposition rate of ozone to atomic oxygen and consequently higher rate of toluene oxidation. In fact, activity orders of the catalysts can be related to the
reverse of Mn *K-edge* absorption energy orders reported in Table 5.5. Einaga and Ogata also found that higher dispersion of Mn is more effective in benzene ozonation at room temperature [56] but it seems that changes in Mn oxidation state as the result of variation in Mn dispersion is controlling the rate of aromatics oxidation such as benzene and toluene by ozone.

5.5. Summary

The effect of manganese loading on the activity of MnO$_x$/γ-alumina was studied in this chapter. Four loadings of Mn (1, 5, 10, and 20%) were used to obtain the activity of the catalysts in the oxidation of toluene in the range of 22-100 °C. Characterization of the catalysts was performed by XRD, XANES, and EXAFS analysis in order to elucidate the relationship between the structure of manganese oxides and Mn loading. XRD analysis showed that Mn$_2$O$_3$ was the dominant phase at Mn loading up to 5%, whereas a mixture of MnO$_2$ and Mn$_2$O$_3$ was observed at higher catalyst loadings. The fraction of MnO$_2$ and the absorption energy of the catalysts increased with the increase of Mn loading. The catalyst with 10% Mn loading had 91.0% of Mn$_2$O$_3$ and 9.0% of MnO$_2$. On the other hand, the catalyst with 20% Mn loading contained 55.7% of MnO$_2$ and 44.7% of Mn$_2$O$_3$. The formal oxidation state of Mn in the catalysts with 1 and 5% Mn loading was less than that of Mn$_2$O$_3$ reference material due to the incomplete growth of Mn$_2$O$_3$ clusters. The oxidation state of MnO$_x$/γ-alumina (10%) was slightly higher than Mn$_2$O$_3$, showing the presence of MnO$_2$ in addition to Mn$_2$O$_3$ nano-particles. The catalyst with 20% loading of manganese had the same absorption energy as that of MnO$_2$, suggesting that the major component of this catalyst is MnO$_2$. EXAFS fitting was in agreement with the result of XRD and XANES, showing the growth of Mn$_2$O$_3$ particles by increasing the loading from 1 to 10%. EXAFS signals and coordination numbers were changed by further increase of the loading from 10 to 20% due to the formation of more MnO$_2$ phase.
Activity tests of the catalysts indicated that lower Mn loadings were more favorable in the oxidation of toluene up to 80°C, at which the catalysts showed the same activity under the reaction conditions. All the catalysts became deactivated at room temperature due to the accumulation of alcohols and carboxylic acids on their surface indicated by CHNS and FT-IR analyses. It was suggested that activity orders of the catalysts could be related to the oxidation state of Mn on the catalyst. Catalysts with lower Mn loadings had higher activity in transferring electrons to ozone to initiate the ozone decomposition reaction due to their lower oxidation state. This increased the decomposition rate of ozone to atomic oxygen which consequently enhanced the oxidation rate of toluene.
CHAPTER 6
PROMOTIONAL EFFECT OF NOBLE METALS ON ACTIVITY OF MANGANESE
OXIDES

It was discussed in Chapters 4 and 5 that manganese oxides are effective catalysts for low
temperature oxidation of toluene. Toluene can be completely oxidized at around 80 °C in the
presence of ozone and MnOₓ dispersed on γ-alumina. One possible approach to increase the
catalyst activity is to add promoters to manganese oxides. There are reports in the literature,
showing that combination of transition metal oxides with noble metals can lead to a synergy in
the activity of catalysts in the combustion of VOCs by oxygen. Andreeva et al. have shown that
addition of gold improves the catalytic activity of vanadium oxides in the oxidation of benzene
[99]. The activity of vanadium oxides was also promoted by palladium for benzene combustion
by Garcia et al. [73]. Galvan and co-workers added palladium to alumina-supported manganese
oxides to reduce the oxidation temperature of formaldehyde-methanol mixture [100-102].
Incorporation of small amounts of palladium improved the activity of manganese oxides, causing
the oxidation of formaldehyde and methanol to occur at room temperature. Similar synergy was
also observed between manganese and platinum in catalytic oxidation of toluene, n-hexane, and
ethanol [72, 74, 103].

Even though the combination of transition metal oxides with noble metals has been
studied in the oxidation of VOCs by oxygen, this issue has not been explored in catalytic
oxidation of VOCs by ozone. The majority of researchers have focused on the ozonation of
VOCs over transition metal oxides. Chapter 6 explores the effect of addition of Pt and Pd to Mn
in order to enhance the activity of alumina-supported manganese oxides in catalytic ozonation of
toluene. This chapter also aims to elaborate atomic interactions between the noble metals and
manganese oxides in order to relate the structure of the monometallic and bimetallic catalysts to the observed catalytic activities.

6.1. Characterization of monometallic and bimetallic catalysts

Platinum and palladium are the two noble metals studied in this chapter. The monometallic catalysts are called Pt/γ-alumina, Pd/γ-alumina, and MnOx/γ-alumina for the catalysts containing Pt, Pd, and Mn, respectively. The bimetallic catalysts are prepared by the impregnation of the prepared MnOx/γ-alumina with Pt and Pd. They are called Pt-MnOx/γ-alumina and Pd-MnOx/γ-alumina for Pt-Mn and Pd-Mn bimetallic catalysts, respectively. Loadings of Pt, Pd, and Mn are 1%, 1%, and 10%, respectively. Catalysts are characterized by X-ray diffraction (XRD), hydrogen temperature programmed reduction (H2-TPR), and X-ray absorption spectroscopy (XAS) including XANES and EXAFS. XAS will be mainly used to clarify the atomic interactions between the noble metals and manganese oxides.

6.1.1. XRD and H2-TPR

Fig. 6.1 presents the XRD spectra of different catalysts studied in this chapter. Fig. 6.1 (a) shows the diffractogram of MnO2 (pyrolusite) and Mn2O3 (bixbyite) on MnOx/γ-alumina as it was presented in Chapters 4 and 5. The addition of Pt and Pd to MnOx/γ-alumina (Fig. 6.1 (b) and 6.1 (c)) does not change the XRD spectra of MnOx/γ-alumina except for the appearance of a platinum peak at 2θ value of 39.8 degree in Fig. 6.1 (b). The same peak is also present in Fig 6.1 (d) which is the XRD of Pt/γ-alumina catalyst with the same loading. This suggests that Pt is likely present in metallic state in Pt-MnOx/γ-alumina and Pt/γ-alumina catalysts. Fig. 6.1 (c) and (e) show no distinct peaks for Pd or PdO, suggesting high dispersion of palladium species on Pd-MnOx/γ-alumina and Pd/γ-alumina.
Fig. 6.1. XRD spectra of the catalysts, (a) MnO\textsubscript{x}/\gamma-alumina, (b) Pt-MnO\textsubscript{x}/\gamma-alumina, (c) Pd-MnO\textsubscript{x}/\gamma-alumina, (d) Pt/\gamma-alumina, and (e) Pd/\gamma-alumina, *: \gamma-alumina, -: Pt, ○: MnO\textsubscript{2}, ●: Mn\textsubscript{2}O\textsubscript{3}.

H\textsubscript{2}-TPR of the catalysts is shown in Fig. 6.2. TPR Profile of Pt/\gamma-alumina, shown in Fig. 6.2 (a), does not show any distinct reduction peak which is likely due to the presence of metallic platinum on the catalyst surface. Fig. 6.2 (b) shows a small broad peak for the alumina-supported palladium catalyst at 121 °C. This implies that palladium oxide species are present on \gamma-alumina, resulting in a broad and weak reduction peak at 121 °C. TPR pattern of MnO\textsubscript{x}/\gamma-alumina catalyst is indicated in Fig. 6.2 (c) with two main peaks at 394 and 508 °C along with a smaller one at
423 °C. The first peak at 394 °C is attributed to the reduction of MnO₂ and Mn₂O₃ to Mn₃O₄ through the sequence of MnO₂ → Mn₂O₃ → Mn₃O₄ [102, 104]. The second and third peaks at 394 and 508 °C are related to the reduction of small and large particles of Mn₃O₄ to MnO, respectively. [102]. TPR of the bimetallic catalysts are depicted in Fig. 6.2 (d) and (e). Addition of platinum and palladium to MnOₓ/γ-alumina greatly reduces the reduction temperature of MnOₓ and causes the total reduction of MnOₓ (MnO₂ and Mn₂O₃ to MnO) to take place at 134 °C in one step. This phenomenon can be explained by the spillover effect of hydrogen from noble metals to manganese oxides, facilitating the reduction reaction at low temperatures [104, 105].

![Graph showing TPR of catalysts](image_url)

Fig. 6.2. H₂-TPR of the catalysts, (a) Pt/γ-alumina, (b) Pd/γ-alumina (4 times zoomed), (c) MnOₓ/γ-alumina, (d) Pt-MnOₓ/γ-alumina, and (e) Pd-MnOₓ/γ-alumina.
6.1.2. XANES and EXAFS analysis

Results of X-ray absorption spectroscopy (XAS) are explained in order to develop a better understanding about atomic interactions between Mn and Pd or Pd in the bimetallic catalysts. XANES analysis is presented quantitatively while EXAFS analysis is demonstrated qualitatively as the magnitude of Fourier transforms.

Pd $K$-edge normalized XANES spectra are shown in Fig. 6.3, in which an absorption edge occurs due to 1s to 5p dipole-allowed excitation. Pd/$\gamma$-alumina and Pd-MnO$_x$/\(\gamma\)-alumina catalysts have the same edge shape with the absorption edge of 24354.0 eV, which is 4.0 eV higher than the edge energy of Pd foil at 24350.0 eV. This value is the same as the absorption energy reported for PdO [106], suggesting that Pd exists in the form of PdO. Fourier transformed EXAFS spectra of Pd/$\gamma$-alumina and Pd-MnO$_x$/\(\gamma\)-alumina are included in Fig. 6.4. The Fourier transform is taken in the range of 2-14 Å$^{-1}$. The first peak at ca. 1.55 Å corresponds to Pd-O backscattering in Pd/$\gamma$-alumina and Pd-MnO$_x$/\(\gamma\)-alumina [106, 107]. No Pd-Pd peak was observed for either of the catalysts. This is due to the high dispersion of Pd on these catalysts, implying that Pd is most likely in amorphous form. The Pd-O peak in Pd/$\gamma$-alumina and Pd-MnO$_x$/\(\gamma\)-alumina is the same in terms of shape and intensity, showing that the oxidation state and the first shell nearest neighbor coordination of Pd do not change in the presence of manganese oxides.
Fig. 6.3. Pd K-edge XANES spectra, (a) Pd/γ-alumina, (b) Pd-MnOₓ/γ-alumina, and (c) Pd foil.

Fig. 6.4. Magnitude of the Fourier transform of Pd K-edge, (a) Pd/γ-alumina, (b) Pd-MnOₓ/γ-alumina, and (c) Pd foil.
Fig. 6.5 shows XANES spectra of Pt _L_{III}-edge_ of the reference materials (Pt foil and α-PtO₂) and Pt containing catalysts. Pt foil does not show an intense white line due to filled d orbital of Pt (5d⁹ 6s¹). On the other hand, α-PtO₂ with vacancies in the d orbital (5d⁶ 6s⁰) has a sharp white line due to 2p_{3/2} to 5d_{3/2} and 5d_{5/2} excitations. It can be seen that the white line intensity of Pt/γ-alumina is higher than that of Pt foil, indicating the presence of platinum oxide species on alumina. The white line intensity of Pt _L_{III}-edge_ is even higher in the presence of manganese oxide, indicating the presence of more Pt oxide species in Pt-MnOₓ/γ-alumina than Pt/γ-alumina. Pt _L_{III}-edge_ absorption energy is 11564.5 and 11565.1 eV for Pt/γ-alumina and Pt-MnOₓ/γ-alumina, respectively. These absorption energies are 0.5 and 1.1 eV higher than the absorption energy of Pt foil at 11564.0 eV.

---

Fig. 6.5. Pt _L_{III}-edge_ XANES spectra, (a) Pt foil, (b) Pt/γ-alumina, (c) Pt-MnOₓ/γ-alumina, and (d) α-PtO₂.
Using linear combination fitting with Pt foil and α-PtO₂ reference materials, it can be found that 84.4% of Pt exists in metallic form and 15.6% of Pt is in +4 oxidation state on Pt/γ-alumina. Addition of Pt to manganese causes the fraction of metallic Pt to drop to 78.7% while the fraction of Pt⁴⁺ increases to 21.3%. This shows that the presence of manganese oxides increases the portion of Pt oxide species when Pt is added to MnOₓ/γ-alumina instead of γ-alumina. Fig. 6.6 shows the magnitude of Fourier transforms of Pt LIII-edge EXAFS of Pt containing catalysts and the Pt reference materials. The Fourier transform is taken in the range of 2-14 Å⁻¹. Basically, the results of XANES study are in agreement with the EXAFS of Pt. It can be seen that the addition of platinum to MnOₓ/γ-alumina reduces the intensity of Pt-Pt peak at ca. 2.70 Å, which mostly corresponds to lower Pt-Pt coordination number (CN). But Pt-O signal at ca. 1.62 Å increases, implying higher Pt-O CN in Pt-MnOₓ/γ-alumina. Decrease in Pt CN and increase in the first shell oxygen CN after the addition of Pt to MnOₓ/γ-alumina indicate higher dispersion of Pt in the presence of Mn, leading to the attachment of some portion of Pt to manganese oxide clusters. The percent of the attached platinum to MnOₓ is estimated to be at least 5.7% which is the difference between the percent of platinum oxides in Pt/γ-alumina (15.6%) and Pt-MnOₓ/γ-alumina (21.3%) quantified by XANES.
It is clear in Fig. 6.5 and 6.6 that platinum oxide species are present on alumina while they are not detectable by H2-TPR and XRD techniques. This indicates that in addition to metallic platinum particles on alumina, platinum oxide species are most likely present as small particles with size of less than 5 nm since there is no platinum oxide peaks in XRD of Pt/γ-alumina and Pt-MnOx/γ-alumina. It also suggests that these small platinum species have a strong interaction with alumina, resulting in no H2 reduction peak for Pt/γ-alumina in Fig. 6.2. This structural model for Pt on alumina is also in agreement with the work of Pan and Cowley and Schneider et al., in which they have observed the presence of separate metallic Pt and Pt oxide particles on alumina [108, 109].

Fig. 6.7 shows the near edge spectra of MnOx/γ-alumina, Pt-MnOx/γ-alumina, Pd-MnOx/γ-alumina, and Mn2O3 reference material. Even though the addition of noble metals
causes a small increase in the white line area of MnO₅/γ-alumina, no changes are observed in the absorption energy of MnO₅/γ-alumina after the addition of noble metals. Basically, MnO₅/γ-alumina, Pt-MnO₅/γ-alumina, and Pd-MnO₅/γ-alumina have the same absorption energies at 6548.4 eV which is 0.3 eV higher than that of Mn₂O₃ at 6548.1 eV.

Fig. 6.7. Mn K-edge XANES spectra, (a) Mn₂O₃, (b) MnO₅/γ-alumina, (c) Pd-MnO₅/γ-alumina, and (d) Pt-MnO₅/γ-alumina.

Fig 6.8 shows the magnitude of Fourier transforms of Mn K-edge. The Fourier transform is taken in the range of 2-14 Å⁻¹. EXAFS peak locations of MnO₅/γ-alumina (Fig. 6.8 (b)), Pd-MnO₅/γ-alumina (Fig. 6.8 (c)), and Pt-MnO₅/γ-alumina (Fig. 6.8 (d)) are similar to that of Mn₂O₃ (Fig. 6.8 (a)) since 91.0% of manganese oxides have Mn₂O₃ structure determined in Chapter 4. It can be seen that signal intensities of Mn-O peak at ca. 1.47 Å and Mn-Mn peak at ca. 2.73 Å are similar in MnO₅/γ-alumina and Pd-MnO₅/γ-alumina, whereas an increase in Mn-Mn signal is
observed in Pt-MnO\textsubscript{x}/γ-alumina by comparing Fig. 6.8 (b) and Fig. 6.8 (d). This shows that Mn coordination number (CN) slightly increases as the result of atomic interaction between Pt and manganese oxides clusters on alumina. On the other hand, Mn-Mn signal at 2.73 Å remains unchanged in the presence of Pd, showing that Mn and Pd do not have atomic interaction.

![Fourier Transform of Mn K-edge](image)

**Fig. 6.8.** Magnitude of the Fourier transform of Mn *K-edge*, (a) Mn\textsubscript{2}O\textsubscript{3}, (b) MnO\textsubscript{x}/γ-alumina, (c) Pd-MnO\textsubscript{x}/γ-alumina, and (d) Pt-MnO\textsubscript{x}/γ-alumina.

Table 6.1 presents a summary of the absorption energies determined in this work by XANES analysis while Table 6.2 gives X-ray photoelectron spectroscopy (XPS) binding energies of similar bimetallic systems reported in the literature in order to compare the results obtained by XANES in this work (Table 6.1) with the other reports (Table 6.2). It can be seen in Table 6.2 that binding energies of Mn 2p\textsubscript{1/2} and Mn 2p\textsubscript{3/2} are more or less constant in the
presence or absence of noble metals. Pd is essentially in oxide form regardless of the fact that it is alone or it coexists with manganese on the support. Pt is slightly oxidized in Pt/γ-alumina and its degree of oxidation increases when manganese is present in the system. Therefore, by comparing the absorption energies of the monometallic and bimetallic catalysts given in Tables 6.1 and 6.2, one can conclude that electronic interactions found by XANES is consistent with those obtained by an alternative approach such as XPS.
Table 6.1. Summary of Mn, Pd and Pt absorption energies determined by XANES

<table>
<thead>
<tr>
<th>Manganese samples</th>
<th>Mn K-edge</th>
<th>Palladium samples</th>
<th>Pd K-edge</th>
<th>Platinum samples</th>
<th>Pt L_{III}-edge</th>
</tr>
</thead>
<tbody>
<tr>
<td>E_0 (eV)</td>
<td>E_0 (eV)</td>
<td>E_0 (eV)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn</td>
<td>6539.0</td>
<td>Pd</td>
<td>24350.0</td>
<td>Pt</td>
<td>11564.0</td>
</tr>
<tr>
<td>Mn_2O_3</td>
<td>6548.1</td>
<td>PdO [106]</td>
<td>24354.0</td>
<td>PtO_2</td>
<td>11566.7</td>
</tr>
<tr>
<td>MnO_2</td>
<td>6552.7</td>
<td>Pd/γ-alumina</td>
<td>24354.0</td>
<td>Pt/γ-alumina</td>
<td>11564.5</td>
</tr>
<tr>
<td>MnO_x/γ-alumina</td>
<td>6548.4</td>
<td>Pd- MnO_x/γ-alumina</td>
<td>24354.0</td>
<td>Pt- MnO_x/γ-alumina</td>
<td>11565.1</td>
</tr>
<tr>
<td>Pd- MnO_x/γ-alumina</td>
<td>6548.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt- MnO_x/γ-alumina</td>
<td>6548.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Catalytic system</td>
<td>Mn 2p&lt;sub&gt;1/2&lt;/sub&gt; (L&lt;sub&gt;III&lt;/sub&gt;)</td>
<td>Mn 2p&lt;sub&gt;3/2&lt;/sub&gt; (L&lt;sub&gt;III&lt;/sub&gt;)</td>
<td>Pt 4d&lt;sub&gt;5/2&lt;/sub&gt; (N&lt;sub&gt;5&lt;/sub&gt;)</td>
<td>Pd 3d&lt;sub&gt;5/2&lt;/sub&gt; (M&lt;sub&gt;5&lt;/sub&gt;)</td>
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<td></td>
<td>BE (eV)</td>
<td>BE (eV)</td>
<td>BE (eV)</td>
<td>BE (eV)</td>
<td></td>
</tr>
<tr>
<td>[104] Mn/alumina</td>
<td>642.8</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt/alumina</td>
<td>-</td>
<td>316.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt-Mn/alumina</td>
<td>642.7</td>
<td>317.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>[74] Mn/alumina</td>
<td>654.0</td>
<td>642.4</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt/alumina</td>
<td>-</td>
<td>-</td>
<td>315.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt-Mn/alumina</td>
<td>654.3</td>
<td>642.5</td>
<td>317.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[101, 102, 110] Mn/alumina</td>
<td>641.7</td>
<td></td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pd/alumina</td>
<td>-</td>
<td></td>
<td>336.6, 336.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pd-Mn/alumina</td>
<td>641.7 to 641.9</td>
<td></td>
<td>336.5 to 336.8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
It is also important to explain the reason why palladium tends to be in oxide form while the majority of platinum is in metallic state on $\gamma$-alumina. This relates to the decomposition temperature of palladium and platinum oxides which are as follows: $\text{PtO (325 }^\circ\text{C)}$, $\text{PtO}_2 (450 \degree\text{C})$, $\text{PdO (750 }^\circ\text{C)}$, and $\text{PdO}_2 (200 \degree\text{C})$ [111]. The calcinations temperature of the noble metal containing catalysts is 550 °C. Therefore, Pd exists as PdO while Pt is mostly in metallic state even though a small portion of Pt is in oxide form as it was revealed by XAS.

6.2. Activity of monometallic and bimetallic catalysts

Catalytic activities of monometallic and bimetallic catalysts at room temperature are shown in Fig. 6.9 and 6.10 in terms of toluene and ozone conversions, respectively. It can be seen in Fig. 6.9 that the activity of the catalysts for the oxidation of toluene by ozone is in the following order: $\text{Pt-MnO}_x/\gamma$-alumina (Fig. 6.9 (b)) $>$ $\text{Pd-MnO}_x/\gamma$-alumina (Fig. 6.9 (c)) $\approx$ $\text{MnO}_x/\gamma$-alumina (Fig. 6.9 (a)) $>$ $\text{Pd}/\gamma$-alumina (Fig. 6.9 (e)) $>$ $\text{Pt}/\gamma$-alumina (Fig. 6.9 (d)). All catalysts lose their activity at room temperature due to the accumulation of carbonaceous species. The same trend of activity order and deactivation behavior is also seen for ozone conversion in Fig. 6.10.
Fig. 6.9. Toluene conversion at room temperature, (a) MnOₓ/γ-alumina, (b) Pt-MnOₓ/γ-alumina, (c) Pd- MnOₓ/γ-alumina, (d) Pt/γ-alumina, and (e) Pd/γ-alumina. Error bars are standard deviations.
Fig. 6.10. Ozone conversion at room temperature, (a) MnO\textsubscript{x}/\gamma-alumina, (b) Pt-MnO\textsubscript{x}/\gamma-alumina, (c) Pd- MnO\textsubscript{x}/\gamma-alumina, (d) Pt/\gamma-alumina, (e) Pd/\gamma-alumina. Error bars are standard deviations.

The activity of the catalysts at higher temperatures of 40, 60, 70, 80, and 100 °C is presented in Fig. 6.11 and 6.12. Increase of the reaction temperature improves toluene conversion and prevents the deactivation of the catalysts. Pt-MnO\textsubscript{x}/\gamma-alumina (Fig. 6.11 (b)) has the highest toluene conversion while Pd-MnO\textsubscript{x}/\gamma-alumina (Fig. 6.11 (c)) and MnO\textsubscript{x}/\gamma-alumina (Fig. 6.11 (a)) have almost the same activity. Pd-MnO\textsubscript{x}/\gamma-alumina and MnO\textsubscript{x}/\gamma-alumina catalysts result in maximum conversion of toluene at 80 °C, after which their activities drop slightly. But complete conversion of toluene is possible at 70 °C with Pt-MnO\textsubscript{x}/\gamma-alumina. This temperature is 10 °C lower than the reaction temperature of Pd-MnO\textsubscript{x}/\gamma-alumina and MnO\textsubscript{x}/\gamma-alumina for the total oxidation of toluene. Pd/\gamma-alumina (Fig. 6.11 (e)) shows higher activity than Pt/\gamma-alumina (Fig. 6.11 (d)) but manganese containing catalysts are more effective than Pd/\gamma-alumina and Pt/\gamma-alumina.
Fig. 6.11. Toluene conversion above room temperature, (a) MnO\textsubscript{x}/\gamma-alumina, (b) Pt-MnO\textsubscript{x}/\gamma-alumina, (c) Pd-MnO\textsubscript{x}/\gamma-alumina, (d) Pt/\gamma-alumina, and (e) Pd/\gamma-alumina. Error bars are standard deviations.

Ozone conversion of the catalysts is shown in Fig. 6.12, in which the activity order of the catalysts is the same as that of toluene conversion. Ozone conversion increases in manganese containing catalysts with the increase of temperature and reaches almost 100% at 80 °C. The plateau in toluene conversion above 80 °C for Pt-MnO\textsubscript{x}/\gamma-alumina, Pd-MnO\textsubscript{x}/\gamma-alumina, and MnO\textsubscript{x}/\gamma-alumina is related to the complete consumption of ozone at 80 °C. Increasing the reaction temperature from 80 °C to 100 °C does not increase the toluene conversion since all ozone is already consumed at 80 °C.
Fig. 6.12. Ozone conversion above room temperature, (a) MnO$_x$/γ-alumina, (b) Pt-MnO$_x$/γ-alumina, (c) Pd-MnO$_x$/γ-alumina, (d) Pt/γ-alumina, and (e) Pd/γ-alumina. Error bars are standard deviations.

From the results discussed on Fig. 6.9 to Fig. 6.12, it can be seen that the addition of platinum to manganese enhances the activity of MnO$_x$/γ-alumina, while Pd does not have any effect on the performance of MnO$_x$/γ-alumina. This result can be explained by XANES and EXAFS studies of Pt-MnO$_x$/γ-alumina and Pd-MnO$_x$/γ-alumina catalysts. Basically, the fraction of Pt oxide species and Pt atomic environment change when Pt coexists with Mn while such changes are not found in the case of Pd. This means that Pt can interact with manganese oxides to improve Mn activity while Pd does not have this capability. Most probably Pd is mainly attached to alumina surface in Pd/γ-alumina and Pd-MnO$_x$/γ-alumina catalysts, resulting in high dispersion of Pd on alumina without atomic interaction with manganese oxide clusters. On the other hand, as found by XANES, the fraction of oxidized Pt in Pt-MnO$_x$/γ-alumina is higher than
that in Pt/γ-alumina. The EXAFS study of Pt also supports the results found from XANES, indicating the decrease in platinum coordination number and the increase in oxygen CN when Pt is added to MnOₓ/γ-alumina.

There are reports on the effect of noble metals on transition metal oxides for catalytic combustion of VOCs in the literature [73, 74, 99, 103]. It can be inferred that noble metals can interact with transition metal oxides via their lattice oxygen, changing electronic environment around transition metals. It is also believed that Pt is most likely attached to manganese oxide clusters via MnOₓ lattice oxygen, altering Pt and Mn coordination numbers.

Higher activity of Pt-MnOₓ/γ-alumina than MnOₓ/γ-alumina is attributed to the possibility that the addition of Pt changes electronic properties of Mn, enhancing the activity of manganese oxides by formation of Mn-O-Pt bonds. Basically, attachment of Pt to MnOₓ results in more electron occupancy of Mn 3d orbital which is considered to be more favorable in the decomposition of ozone and the oxidation of VOCs [29]. In fact, as it was shown in chapter 5, lower oxidation states of Mn can facilitate the transfer of electrons to ozone. Therefore, one can relate higher activity of Pt-MnOₓ/γ-alumina to higher Mn 3d orbital electron occupancy, accelerating the rate of ozone decomposition and consequently the oxidation rate of toluene.

In order to examine the effect of manganese 3d valence electron occupancy on catalytic ozonation of toluene, Pt-MnOₓ/γ-alumina and Pd-MnOₓ/γ-alumina catalysts are reduced with hydrogen. The catalysts are reduced at 250 °C for two hours with pure H₂ at flow rate of 100 ml/min. Fig. F.1 of Appendix F shows XRD of the catalysts after the reduction. Reduced Pt-MnOₓ/γ-alumina and Pd-MnOₓ/γ-alumina show only MnO peaks (manganosite), corresponding to the reduction of MnO₂ and Mn₂O₃ to MnO. This is in agreement with the TPR profile of Pt-MnOₓ/γ-alumina and Pd-MnOₓ/γ-alumina shown in Fig.6.2, in which manganese oxides (MnO₂
and Mn₂O₃) are reduced to MnO in one step at 134 °C. In fact, the reduction of the bimetallic catalysts with pure hydrogen at 250 °C ensures that the only remaining phase of Mn is MnO after the reduction. Activities of the reduced and unreduced catalysts (Pd-MnOₓ/γ-alumina, Pt-MnOₓ/γ-alumina) are shown in Fig. F.2 and Fig. F.3 of Appendix F in terms of toluene and ozone conversions, respectively. It can be seen that the activity of Pd-MnOₓ/γ-alumina increases after the reduction with hydrogen and reaches to a similar level as the unreduced Pt-MnOₓ/γ-alumina in terms of ozone and toluene conversions. Activity of the reduced Pt-MnOₓ/γ-alumina overlaps with the activity of the unreduced Pt-MnOₓ/γ-alumina at room temperature and higher temperatures. Therefore, it can be concluded that lower oxidation state of manganese oxides are more favorable for catalytic ozonation of toluene in agreement with the conclusion made in Chapter 5. Thus, the effect of platinum addition to MnOₓ/γ-alumina can be explained by increasing electron occupancy of 3d orbital of Mn since the reduced Pd-MnOₓ/γ-alumina showed the same activity as the unreduced Pt-MnOₓ/γ-alumina catalyst.

6.3. Summary

This chapter studied the promotional effect of two noble metals (Pt and Pd) on the activity of MnOₓ/γ-alumina in the oxidation of toluene. Catalysts were characterized by XRD, H₂-TPR, XANES, and EXAFS. Catalysts activities were obtained in the range of 22-100 °C. It was observed that the addition of Pd did not improve the activity of alumina-supported manganese oxides catalysts, whereas the addition of Pt led to 10 °C reduction in the required temperature for the complete oxidation of toluene. XANES and EXAFS studies showed no atomic interaction between Pd and Mn while XANES and EXAFS of Pt changed in the presence of Mn. It was concluded that the interaction between Pt and Mn occurs via surface oxygen of manganese oxides, leading to the formation of Mn-O-Pt bonds. The higher activity of Pt-
MnO$_x$/γ-alumina can be related to the higher electron occupancy of 3d orbital in Mn, accelerating the decomposition rate of ozone and also the rate of toluene oxidation. Activity tests of the reduced catalysts confirmed that lower oxidation states of Mn enhances the activity of the catalysts. This indicated that higher 3d orbital electron occupancy of Mn in Pt-MnO$_x$/γ-alumina can be the reason for better activity of Pt promoted manganese oxides than the other catalysts.
As it has been mentioned in Chapter 2, one of the unclear issues about the catalytic oxidation of VOCs by ozone is the kinetics and mechanism of the reaction. It is believed that the reaction is a surface phenomenon, following a Langmuir-Hinshelwood mechanism between adsorbed VOC molecules and atomic oxygen. Ozone is decomposed to atomic oxygen and peroxide species on manganese oxides. VOC molecules are assumed to be adsorbed on the catalyst and react with atomic oxygen to start the oxidation reaction. This mechanism has been originally proposed by Oyama and co-workers in order to study the kinetics and mechanism of the oxidation of acetone by ozone over MnOₓ/SiO₂ [51]. They have used in-situ laser Raman spectroscopy and transient kinetics techniques to determine the reactive surface species and the reaction mechanism. The elementary steps of this mechanism have been presented through Eq. (2.4) to (2.9) in Chapter 2. To the best of the author’s knowledge, this is the only systematic work done on the kinetics and mechanism of catalytic ozonation of VOCs. Basically, most of the researchers have simply used Oyama’s mechanism in order to explain the mechanism of ozonation of several VOCs such as toluene, benzene, and cyclohexane [21, 25, 53, 58, 69]. This has been done without any attempt to test the validity of this mechanism on VOCs other than acetone. In fact, the generality of the mechanism proposed by Oyama and co-workers has not been studied yet. Therefore, there is need to further study the kinetics of the oxidation of VOCs by ozone in order to explore possible elementary steps involved in the reaction mechanism.

This chapter investigates the kinetics of the catalytic oxidation of toluene by ozone over MnOₓ/γ-alumina (10%). This is achieved by obtaining initial reaction rates at constant ozone or
toluene partial pressures. The obtained reaction rates are first fitted to a power law model in order to obtain the apparent activation energy of the reactions and the reaction orders with respect to ozone and toluene. The initial rates are also fitted to a rate equation derived based on the mechanism proposed by Oyama and co-workers in order to check the applicability of this mechanism to catalytic oxidation of toluene. A more detailed mechanism based on the evidence reported on the oxidation of VOCs (especially toluene) is presented in this chapter to show an alternative reaction mechanism for the catalytic ozonation of toluene.

7.1. Data collection and analysis

Oxidation of toluene is carried out in the same experimental set-up explained in Chapter 3. Kinetics data is obtained at four different temperature levels (70, 80, 90, and 100 °C) since catalyst deactivation occurs below 70 °C. Experiments are conducted with either constant toluene partial pressure (P_{tol.}) of $240 \times 10^{-6}$ atm while ozone partial pressure (P_{ozo.}) is changed between 1050 to $3000 \times 10^{-6}$ atm or constant ozone partial pressure of $1050 \times 10^{-6}$ atm with toluene partial pressure of 90 to $180 \times 10^{-6}$ atm. The space time of the reactor is between 0-1212 kg_{cat.}mol_{tol.}^{-1} s. MnO_{x/\gamma}-alumina (10%) is diluted with silica carbide, inert in the reaction, to prevent gas phase channeling and temperature gradients along the catalyst bed.

Plots of toluene conversion versus space time are used in order to obtain the initial reaction rates of toluene oxidation ($r_{tol.}$) based on the following equation:

$$r_{tol.} = \frac{dX_{tol.}}{d\left(\frac{W_{cat.}}{F_{tol.,in}}\right)}$$

(7.1)

in which, $X_{tol.}$, $W_{cat.}$, and $F_{tol.,in}$ represent toluene conversion, catalyst weight, and inlet molar flow rate of toluene, respectively. Two numerical methods are used to estimate the initial
reaction rates based on Eq. (7.1). The first one is fitting the conversion data by smoothing splines in MATLAB R2006a curve fitting toolbox and obtaining the initial reaction rates by differentiating the splines at zero space time. The second method is based on the fitting of the data with Eq. (7.2) and calculating the initial rates by product of A and B [112]:

$$X_{\text{tol.}} = A \left(1 - \exp \left( - \frac{B}{X_{\text{tol.}, \text{in}}} \right) \right)$$  \hspace{1cm} (7.2)

A and B are the fitting parameters evaluated by fitting Eq. (7.2) to the conversion versus space time data. The initial rates calculated from the two methods, smoothing splines and Eq. (7.2), are averaged and used for kinetics analysis.

Kinetics models are fitted to the experimental data by non-linear least squares technique in MATLAB R2006a using lsqlin function to minimize the sum of squared differences between experimental and calculated reaction rates:

$$\min \sum_i \sum_j \left( r_{\text{exp}, ij} - r_{\text{cal}, ij} \right)^2$$  \hspace{1cm} (7.3)

$r_{\text{exp}}$ and $r_{\text{cal}}$ denote experimental and calculated initial rates of toluene oxidation, respectively. Index i refers to different temperature levels (70, 80, 90, and 100 °C) and index j refers to $j^{th}$ reaction rate at temperature level i, corresponding to specific toluene and ozone partial pressures. In fact, model parameters including rate constants, toluene adsorption entropy, toluene adsorption enthalpy, and activation energies are simultaneously estimated by Eq. (7.3) which allows non-isothermal fitting of the data. Toluene rate constant and toluene adsorption pressure equilibrium constant are defined by Eq. (7.4) and (7.5), respectively:

$$k = k_o \exp \left( \frac{-E}{RT} \right)$$  \hspace{1cm} (7.4)
\[
K_{\text{tol.}} = \frac{K}{P^o} = \exp\left(\frac{\Delta S^o_{\text{tol.}} - \Delta H^o_{\text{tol.}}}{R \cdot T}\right)
\]  

(7.5)

k, \(k_0\), and K are toluene reaction rate constant, pre-exponential factor, and dimensionless adsorption equilibrium constant respectively. \(K_{\text{tol.}}\) is the toluene adsorption pressure equilibrium constant. \(\Delta S^o_{\text{tol.}}\) and \(\Delta H^o_{\text{tol.}}\) are standard entropy and enthalpy of toluene adsorption. R is the universal gas constant and \(P^o\) is the pressure of standard state for gases which is 1 atm. Values of \(k_0\) and E are constrained to positive values while \(\Delta S^o_{\text{tol.}}\) and \(\Delta H^o_{\text{tol.}}\) are allowed to float in negative and positive ranges to minimize the least square objective function expressed by Eq. (7.3).

**Goodness of fits** is evaluated based on the coefficient of determination (\(R^2\)) [113]:

\[
R^2 = 1 - \frac{\sum_i \sum_j (r_{\text{exp,ij}} - r_{\text{cal,ij}})^2}{\sum_i \sum_j (r_{\text{exp,ij}} - r_{\text{mean}})^2}
\]  

(7.6)

\(r_{\text{mean}}\) is the average reaction rate over all data points. \(i\) and \(j\) are the same as Eq. (7.3). Kinetic models are compared by Fisher’s F-test in order to compare different fits [114]:

\[
F_c = \frac{\sum_i \sum_j \left(\frac{r_{\text{cal,ij}}}{P}\right)^2}{\sum_i \sum_j \left(\frac{r_{\text{exp,ij}} - r_{\text{cal,ij}}}{DF}\right)^2}
\]  

(7.7)

Definitions of \(i\) and \(j\) indices are the same as Eq. (7.3). DF is the degree of freedom, defined as the difference between the number of experimental data points and the number of fitting parameters (\(P\)). The best model among the rival models can be distinguished by using Eq. (7.7), suggesting that models with higher \(F_c\) values have higher regression significant.
7.2. Effect of toluene and ozone partial pressures on toluene oxidation

The effect of toluene and ozone partial pressures on toluene conversion at different reactor space times and temperatures is shown in Fig. 7.1 and 7.2. Fig. 7.1 indicates changes in toluene conversion at constant ozone partial pressure when toluene partial pressure is increased. Four toluene partial pressures are considered at 90, 120, 180, and $240 \times 10^{-6}$ atm while the partial pressure of ozone is kept at $1050 \times 10^{-6}$ atm. Toluene conversions ($X_{\text{tol.}}$) versus space time ($W_{\text{cat.}}/F_{\text{tol.}}$) at each toluene partial pressure are shown in Fig. 7.1 (A), (B), (C), and (D). Conversions are obtained at four temperature levels of 70, 80, 90, and 100 °C. Space time of the reactor ranges from 0 to 1212 kg$_{\text{cat.}}$ mol$_{\text{tol.}}^{-1}$ s. Results show that at a constant reactor space time, toluene conversion decreases by the increase of toluene partial pressure while toluene conversions increase by the increasing the temperature.
Fig. 7.1. Toluene conversion versus space time at constant ozone partial pressure ($P_{ozo}$) of $1.05 \times 10^{-6}$ atm and variable toluene partial pressures ($P_{tol}$) at different temperatures; (A) $P_{tol}=9 \times 10^{-6}$ atm, (B) $P_{tol}=12 \times 10^{-6}$ atm, (C) $P_{tol}=18 \times 10^{-6}$ atm, and (D) $P_{tol}=24 \times 10^{-6}$ atm; (1) 70 °C, (2) 80 °C, (3) 90 °C, and (4) 100 °C. Dotted lines show the trend of data.

By visual inspection of Fig. 7.1 (A) to Fig. 7.1 (D) at different $P_{tol}$, it can be seen that the slope of toluene conversion versus space time decreases at zero space time by increasing toluene partial pressure. This implies that the initial rate of toluene oxidation decreases by the increase of toluene concentration. Fig. 7.2 shows the effect of $P_{ozo}$ on toluene conversion when $P_{tol}$ is kept constant at $24 \times 10^{-6}$ atm.
Fig. 7.2. Toluene conversion versus space time at constant toluene partial pressure ($P_{\text{tol.}}$) of $240 \times 10^{-6}$ atm and variable ozone partial pressures ($P_{\text{ozo.}}$) at different temperatures; (A) $P_{\text{ozo.}}$=1050 $\times 10^{-6}$ atm, (B) $P_{\text{ozo.}}$=2000 $\times 10^{-6}$ atm, and (C) $P_{\text{ozo.}}$=3000 $\times 10^{-6}$ atm; (1) 70 °C, (2) 80 °C, (3) 90 °C, and (4) 100 °C. Dotted lines show the trend of data.

Toluene conversions were obtained at three ozone partial pressures of 1050, 2000, and $3000 \times 10^{-6}$ atm, shown in Fig. 7.2 (A), (B), and (C), respectively. In contrast to the result of fixed $P_{\text{tol.}}$, toluene conversions increase by increasing $P_{\text{ozo.}}$ at a fixed reactor space time. It can also be observed that the slope of $X_{\text{tol.}}$ versus $\frac{W_{\text{cat}}}{F_{\text{tol.}}}$ at zero space time and consequently the rate of toluene oxidation increases by the increase of ozone partial pressure.

7.3. Kinetics modeling

Initial reaction rates can be estimated from Fig. 7.1 and Fig. 7.2 in order to fit different kinetics models to the experimental data. Power law model is the first model used in section
7.3.1 to determine the apparent activation energy of the reaction and toluene and ozone reaction orders. The second reaction rate equation is based on the mechanism proposed by Oyama and co-workers for ozonation of acetone [51]. This model is based on a Langmuir-Hinshelwood (LH1) mechanism, in which adsorbed toluene molecules are activated by surface atomic oxygen. The result of the fitting of LH1 is presented in section 7.3.2, in which the goal is to examine the applicability of Oyama’s mechanism to VOCs other than acetone. Eventually, a new mechanism is proposed in section 7.3.3 based on recently published papers on the oxidation of VOCs by atomic oxygen. This mechanism is also developed based on a Langmuir-Hinshelwood mechanism which will be referred to as LH2. LH2 mechanism helps to introduce an alternative mechanism scheme to LH1 which might better explain the catalytic oxidation of toluene by ozone.

7.3.1. Power law model

The power law model is expressed as:

\[ r_{\text{tol}} = k P_{\text{tol}}^n P_{\text{ozo}}^m \]  

(7.8)

The result of fitting Eq. (7.8) to the initial reaction rates is included in Table 7.1 and Fig. 7.3. The reaction orders with respect to toluene and ozone are -1 and 2, respectively. Fig. 7.3 (A) indicates that \( P_{\text{tol}} \) has a negative effect on the initial reaction rate, causing the reaction rate to decrease by increasing toluene partial pressure from 90 to \( 240 \times 10^{-6} \) atm. This results in a reaction order of -1 in Eq. (7.8) for toluene. On the other hand, Fig. 7.3 (B) shows that the initial rate of toluene oxidation increases by increasing ozone partial pressure from 1050 to \( 3000 \times 10^{-6} \) atm which leads to an ozone reaction order of 2. The apparent activation energy (E) of the
reaction is determined to be 31.0 kJ mol\(^{-1}\) which is close to the value reported by Naydenov and Mehandjiev for the oxidation of benzene over MnO\(_2\) by ozone, i.e. 30 kJ mol\(^{-1}\) [41].

Table 7.1. Fitting results of the power law model \(^{a}\)

\[
\begin{align*}
\n r_{\text{tol}} &= k P_{\text{tol}}^n P_{\text{ozo}}^m, \quad k = k_0 \exp\left(-\frac{E}{RT}\right) \\
 n &\quad m \quad k_0 \text{ (mol\_tol kg\_cat s\_1 atm\_1 \times 10^{-4})} \quad E \text{ (kJ mol\_1)} \quad R^2 \quad F_c \\
 -1.0 \pm 0.2 &\quad 2.0 \pm 0.1 &\quad 1.1 \pm 0.8 &\quad 31.0 \pm 0.3 &\quad 0.996 &\quad 1705
\end{align*}
\]

\(^{a}\) Uncertainties are based on 95\% confidence intervals.
Fig. 7.3. Initial reaction rates and the power law model; (A) Constant ozone partial pressure ($P_{\text{ozone}}$) at $1050 \times 10^{-6}$ atm; (B) Constant toluene partial pressure ($P_{\text{tol.}}$) at $240 \times 10^{-6}$ atm; (1) 70 °C, (2) 80 °C, (3) 90 °C, (4) 100 °C. Solid lines show the predicted initial reaction rates by the power law model.

The apparent activation energy of the reaction (31.0 kJ mol$^{-1}$) is lower than the activation energy of catalytic total oxidation of toluene by oxygen which has been reported to be between 75 and 106 kJ mol$^{-1}$ [115]. The decrease of the activation energy can be considered to be the main reason for the effectiveness of the oxidation of VOCs by ozone which makes it possible to oxidize VOCs below 100 °C. The activation energy of the reaction (31.0 kJ mol$^{-1}$) is also lower than the activation energy of the homogeneous reaction between toluene and ozone (55.6 kJ mol$^{-1}$) reported for partial oxidation of toluene by ozone [116]. This indicates that the oxidation of toluene by ozone over MnO$_x$/γ-alumina is a heterogeneous reaction catalyzed by manganese oxides.
7.3.2. **Langmuir-Hinshelwood mechanism (LH1) based on toluene activation by atomic oxygen**

As mentioned before, this mechanism (LH1) is originally suggested by Oyama and co-workers for catalytic oxidation of acetone by ozone [51]. The elementary steps of LH1 mechanism are shown in Table 7.2. The mechanism consists of two catalytic cycles. The first cycle is ozone decomposition which is shown by steps (1) to (3) in Table 7.2. These three steps are considered for the decomposition of ozone to atomic oxygen and peroxide species which eventually decompose ozone to molecular oxygen. The second cycle is composed of quasi equilibrated adsorption of toluene by step (4) and reaction with atomic oxygen in step (5) to activate the adsorbed toluene molecule and to start the oxidation reaction. It is assumed that further steps after step (5) are fast and are not significant from kinetics point of view. The two cycles are coupled to each other by consuming atomic oxygen. If n oxygen atoms are consumed in step (5) in Table 7.2, the rate of toluene oxidation will be [51):

\[
\begin{align*}
    r_{\text{tol.}} &= K_{\text{tol.}} \frac{k'_1 P_{\text{tol.}} P_{\text{o2.}}}{(1 + k'_2 P_{\text{o2.}} + K_{\text{tol.}} P_{\text{tol.}})^{n+1}} \\
    & \quad \text{where } K_{\text{tol.}} \text{ is expressed by Eq. (7.5). } k'_1 \text{ and } k'_2 \text{ are defined as follows:}
    \end{align*}
\]

\[
\begin{align*}
    k'_1 &= \frac{k_1}{k_3} = k'_{1o} \exp \left( \frac{-E'_1}{RT} \right) \\
    k'_2 &= \frac{k_2}{k_3} = k'_{2o} \exp \left( \frac{-E'_2}{RT} \right)
\end{align*}
\]
Table 7.2. Elementary steps of LH1 mechanism

<table>
<thead>
<tr>
<th>Elementary steps based on [51]</th>
<th>Reaction rate/Equilibrium constant</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) ( \text{O}_3 + \ast \rightarrow \text{O}_2 + \text{O*} )</td>
<td>( k_1 )</td>
</tr>
<tr>
<td>(2) ( \text{O}_3 + \text{O*} \rightarrow \text{O}_2 + \text{O}_2^* )</td>
<td>( k_2 )</td>
</tr>
<tr>
<td>(3) ( \text{O}_2^* \rightarrow \text{O}_2 + \ast )</td>
<td>( k_3 )</td>
</tr>
<tr>
<td>(4) ( \text{C}_6\text{H}_5\text{-CH}_3^* \leftrightarrow \text{C}_6\text{H}_5\text{-CH}_3^* )</td>
<td>( K_{\text{tol.}} )</td>
</tr>
<tr>
<td>(5) ( \text{C}_6\text{H}_5\text{-CH}_3^* + n\text{O*} \rightarrow \text{products} )</td>
<td>( k_5 )</td>
</tr>
</tbody>
</table>

The first term in the denominator of Eq. (7.9) corresponds to the fraction of vacant catalyst active sites (\( \theta^* \)) which are proposed to be Mn atoms. The second term in the denominator (\( k'_1 P_\text{o2} \)) is related to the fraction of active sites covered by peroxide species (\( \theta_{\text{o2}} \)) and the last term (\( K_{\text{tol.}} P_{\text{tol.}} \)) is due to the fraction of active sites covered by toluene molecules (\( \theta_{\text{tol.}} \)). The concentration of surface atomic species is assumed to be negligible due to the high reactivity and short lifetime of these species. Oyama and co-workers have shown that peroxide species are spectators in the catalytic oxidation of acetone by ozone and atomic oxygen is responsible for the oxidation reaction [51]. However, they have considered the peroxide species in the calculation of the fraction of vacant active sites.

Results of the fitting of Eq. (7.9) to the initial reaction rates are shown in Table 7.3. It is found that 2 oxygen atoms are necessary to start the oxidation reaction by step (5) in Table 7.2. The first pre-exponential factor (\( \kappa'_{1o} \)) is much larger than the second pre-exponential factor (\( \kappa'_{2o} \)). The high value of \( \kappa'_{1o} \) shows that step (5) is probably a fast step with high reaction rate constant since \( \kappa'_1 \) is defined as the product of \( k_5 \) and \( k'_2 \). The value of \( E'_2 \) (89.3 kJ mol\(^{-1}\)) is
positive which shows that the activation energy of the desorption of oxygen molecule in step (3) is larger than the activation energy of ozone adsorption in step (1).

Table 7.3. Fitting result of LH1 mechanism based on Eq. (7.9)

<table>
<thead>
<tr>
<th>n</th>
<th>( k'_{1o} )</th>
<th>( E'_{1} )</th>
<th>( k'_{2o} )</th>
<th>( E'_{2} )</th>
<th>( \Delta S_{\text{tol}}^{o} )</th>
<th>( \Delta H_{\text{tol}}^{o} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>(mol_{\text{tol.}} \cdot kg_{\text{cat.}}^{-1} \cdot s^{-1} \cdot atm^{-2})</td>
<td>(kJ \cdot mol^{-1})</td>
<td>(atm^{-1})</td>
<td>(kJ \cdot mol^{-1})</td>
<td>(J \cdot mol^{-1} \cdot K^{-1})</td>
<td>(kJ \cdot mol^{-1})</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>( 3 \times 10^{10} )</td>
<td>41</td>
<td>0.2</td>
<td>89</td>
<td>100</td>
<td>8</td>
</tr>
</tbody>
</table>

By calculating \( K_{\text{tol.}} \) using Eq. (7.5) and the values reported in Table 7.3, one can conclude that \( K_{\text{tol.}} P_{\text{tol.}} \gg k'_{2o} P_{\text{oon.}} \). This shows that fraction of active sites covered by peroxide species are negligible in comparison with the ones occupied by toluene. This is in agreement with the findings of Oyama and co-workers, showing that peroxide species are spectators in the reaction and should not appear in the rate equation. Therefore, Eq. (7.9) can be reduced to a simpler rate equation by excluding the fraction of peroxide species:

\[
r_{\text{tol.}} = K_{\text{tol.}} \frac{k'_{1o} P_{\text{tol.}} P_{\text{oon.}}^{n}}{(1 + K_{\text{tol.}} P_{\text{tol.}})^{n+1}} \quad (7.12)
\]

The results of fitting Eq. (7.12) to the initial reaction rates are shown in Fig. 7.4 and Table 7.4. The same values are estimated for \( n, k'_{1o}, E'_{1}, \Delta S_{\text{tol.}}, \) and \( \Delta H_{\text{tol.}} \), showing that the surface peroxide species are most likely not involved in the reaction mechanism.
Fig. 7.4. Initial reaction rates and LH1 model; (A) Constant ozone partial pressure ($P_{\text{ozo.}}$) at $1050 \times 10^{-6}$ atm; (B) Constant toluene partial pressure ($P_{\text{tol.}}$) at $240 \times 10^{-6}$ atm; (1) 70 °C, (2) 80 °C, (3) 90 °C, (4) 100 °C. Solid lines show the predicted initial reaction rates by the power law model.
Table 7.4. Fitting result of LH1 mechanism based on Eq. (7.12) \(^a\)

<table>
<thead>
<tr>
<th>n</th>
<th>(k'<em>n \times 10^{10}) (mol(</em>{\text{tol.}}) kg(_{\text{cat.}})(^{-1}) s(^{-1}) atm(^{-2}))</th>
<th>(E'_i) (kJ mol(^{-1}))</th>
<th>(\Delta S^o_{\text{tol.}}) (J mol(^{-1}) K(^{-1}))</th>
<th>(\Delta H^o_{\text{tol.}}) (kJ mol(^{-1}))</th>
<th>(R^2)</th>
<th>(F_c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0 ± 0.1</td>
<td>3 ± 1.4</td>
<td>41.2 ± 19.4</td>
<td>100.1 ± 46.3</td>
<td>7.9 ± 16.2</td>
<td>0.997</td>
<td>1402</td>
</tr>
</tbody>
</table>

\(^a\)Uncertainties are based on 95% confidence intervals.

The value of standard entropy \((\Delta S^0_{\text{tol.}})\) and enthalpy \((\Delta H^0_{\text{tol.}})\) of toluene adsorption are positive in Table 7.4. Considering the confidence intervals reported for \(\Delta S_{\text{tol.}}\), it can be seen that the fitting of LH1 mechanism to experimental data results in a positive value for entropy of toluene adsorption. It has been mentioned by Froment and Vannice that adsorption entropy should satisfy the following criterion in catalytic reactions [114, 117]:

\[
0 < -\frac{\Delta S^o_{\text{ads}}}{S^o_g}
\]  

\((7.13)\)

\(\Delta S^o_{\text{ads}}\) is the standard entropy of adsorption and \(S^o_g\) is the standard entropy in the gas phase with the value of 320.7 J mol\(^{-1}\) K\(^{-1}\) [115]. It can be observed that the positive value of toluene entropy of adsorption does not satisfy Eq. (7.13), indicating that LH1 mechanism most probably cannot adequately explain the mechanism of catalytic oxidation of toluene by ozone.

Fisher’s \(F_c\) values can be used to compare the power law model and LH1 mechanisms from statistical point of view. The \(F_c\) corresponding to the power law model (1705) is larger than that of the LH1 mechanism (1402) since the number of fitting parameters is less in the power law model. However, both models seem to perfectly follow the trend of the experimental data.
7.3.3. Langmuir-Hinshelwood mechanism (LH2) based on toluene activation by hydrogen abstraction

Even though the LH1 mechanism, presented in Table 7.2, agrees well with the experimental data, it results in a positive value for the entropy of toluene adsorption which is not acceptable for catalytic reactions following Langmuir-Hinshelwood mechanisms. In fact, LH1 mechanism has been originally proposed for acetone oxidation and it might not be necessarily suitable for the oxidation of other VOCs such as toluene. Thus, it was attempted to modify the LH1 mechanism based on evidences reported on the mechanism of catalytic oxidation of hydrocarbons and especially toluene by atomic oxygen.

It has been proposed that toluene interacts dissociatively with the surface of transition metal oxides via abstraction of one H atom from the methyl group producing one adsorbed proton and adsorbed radical benzyl species [118-120]. This step is considered to be the first step in the oxidation of toluene. Iglesia and co-workers also proved with isotopic data and density functional theory (DFT) calculations that C-H bond cleavage is the kinetically relevant step in the oxidation of alkanes (C\textsubscript{n}H\textsubscript{2n+2}) which occurs on catalyst active site pairs [121, 122]. The research group of Marin has also studied the reaction network of catalytic toluene oxidation by temporal analysis of products (TAP) and isotopic techniques [123]. They have observed that hydrogen abstraction from the methyl and phenyl groups occurs before the oxidation of carbon atoms, releasing H\textsubscript{2}O before the detection of CO\textsubscript{2}. The application of C-H bond cleavage model for catalytic oxidation of hydrocarbons by ozone has also been used by Li and Oyama for catalytic ozonation of ethanol, even though the authors did not propose a rate equation based on this mechanism [18].

Therefore, a second Langmuir-Hinshelwood mechanism (LH2) is proposed for the oxidation of toluene by ozone based on the initiation of the reaction by hydrogen abstraction.
from the methyl group [51, 122, 123]. The scheme of the mechanism is shown in Table 7.5. No by-products other than CO, CO₂ and water vapor are considered in the scheme.

Table 7.5. Scheme of the Langmuir-Hinshelwood mechanism (LH2) based on C-H activation

<table>
<thead>
<tr>
<th>Step</th>
<th>Reaction rate/Equilibrium constant</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1)</td>
<td>O₃ + * → O₂ + O*</td>
</tr>
<tr>
<td>(2)</td>
<td>O₃ + O*→ O₂ + O₂*</td>
</tr>
<tr>
<td>(3)</td>
<td>O₂* → O₂ + *</td>
</tr>
<tr>
<td>(4)</td>
<td>C₆H₅-CH₃ + * ⇌ C₆H₅-CH₃*</td>
</tr>
<tr>
<td>(5)</td>
<td>C₆H₅-CH₃* + * → C₆H₅-CH₂* + H*</td>
</tr>
<tr>
<td>(6)</td>
<td>C₆H₅-CH₂* + 7* → C₆ –C* + 7H*</td>
</tr>
<tr>
<td>(7)</td>
<td>H* + O* ⇌ OH* + *</td>
</tr>
<tr>
<td>(8)</td>
<td>2OH* ⇌ H₂O* + O*</td>
</tr>
<tr>
<td>(9)</td>
<td>H₂O* ⇌ H₂O + *</td>
</tr>
<tr>
<td>(10)</td>
<td>C₆ –C* +O* → C₆* + CO*</td>
</tr>
<tr>
<td>(11)</td>
<td>C₆* + 6O* → 6CO*</td>
</tr>
<tr>
<td>(12)</td>
<td>CO* + O* → CO₂* + *</td>
</tr>
<tr>
<td>(13)</td>
<td>CO* ⇌ CO +*</td>
</tr>
<tr>
<td>(14)</td>
<td>CO₂ ⇌ CO₂ + *</td>
</tr>
</tbody>
</table>

The first three steps in Table 7.5 are considered for the decomposition of ozone. The fourth step is quasi equilibrated adsorption of toluene on the first catalyst site. The fifth step is release of one proton on the second catalyst site from the adsorbed toluene molecule to produce C₆H₅-CH₂* and H⁺. C₆H₅-CH₂* undergoes successive H abstractions to produce more H⁺ and a
hydrogen free carbon structure (C$_6$-C$^*$) shown in step 6. The protons react with adsorbed oxygen atoms to produce OH$^*$ at the seventh step. The produced OH$^*$ species combine and produce water through steps 8 and 9 in Table 7.5. The next step would be the reaction of the carbon atom from the methyl group in step 10 to produce an adsorbed CO molecule, leaving the C$_6$ ring (C$_6^*$). The ring opening occurs in step (11) of Table 7.5, in which adsorbed CO molecules are formed. The adsorbed CO molecules can be oxidized to CO$_2$ in step (12) or desorb via step (13). The generated CO$_2$ molecules can leave the surface in step (14).

It has to be noted that step (6) and (11) are not elementary steps. Step (6) represents successive abstraction of hydrogen from C$_6$H$_5$-CH$_2^*$ while step (11) indicates successive oxidation of carbon atoms in the C$_6$ ring by atomic oxygen. Fig. 7.5 shows a simple illustration of the mechanism shown in Table 7.5. It indicates two catalytic cycles, consisting of ozone decomposition and the oxidation of toluene. Toluene is assumed to be adsorbed by the abstraction of one hydrogen atoms on two adjacent sites. The adsorbed toluene reacts with atomic oxygen to produce oxidation by-products.
Assuming that CO, CO₂, and H₂O are the only by-products of the reaction and all O* atoms react with toluene, the steady state mass balance for O* gives:

\[ r_1 = r_7 - r_8 + r_{10} + 6r_{11} + r_{12} \]  \hspace{1cm} (7.14)

In which, \( r_1 \) is the rate of generation of surface atomic oxygen (O*) by step (1) in Table 7.5. \( r_7, r_{10}, r_{11}, \) and \( r_{12} \) are the rate of consumption of ozone by step (7), (10), (11), and (12), respectively. \( r_8 \) is the rate of recombination of OH* in step (8). It should be noted that \( r_{11} \) is the reaction rate between one carbon atom from the C₆ ring (C₆*) with one adsorbed oxygen atom (O*) which can be expressed by:

\[ r_{11} = k_{11} \theta_O \cdot \theta_{C_n}, \, n = 1, ..., 6 \]  \hspace{1cm} (7.15)

Therefore, \( r_{11} \) is multiplied by 6 in Eq. (7.14) assuming that reaction rates of carbon atoms from the C₆* ring with O* are equal. Steady state mass balance for C₆H₅-CH₂* and H* results in:

\[ r_6 = r_5 \]  \hspace{1cm} (7.16)
\[ r_7 = r_5 + 7r_6 = 8r_5 \quad (7.17) \]

\( r_6 \) is the rate of successive abstraction of one hydrogen atom from \( \text{C}_6 \text{H}_5\text{-CH}_2^+ \):

\[ r_6 = k_6 \theta_{c,H_n^+}^n, n = 1, \ldots, 7 \quad (7.18) \]

\( \text{C}_6 \text{H}_5\text{-CH}_2^+ \) is shown by \( \text{C}_7 \text{H}_n^+ \) in Eq. (7.18) for simplicity. Steady state mass balance for \( \text{OH}^+ \) gives:

\[ r_7 = 2r_8 \quad (7.19) \]

Therefore \( r_8 \) becomes:

\[ r_8 = \frac{r_7}{2} = 4r_5 \quad (7.20) \]

Steady state mass balance for \( \text{C}_6\text{-C}^+, \text{C}_6^+ \) and \( \text{CO}^+ \) leads to Eq. (7.21), (7.22), and (7.23), respectively:

\[ r_{10} = r_6 = r_5 \quad (7.21) \]

\[ r_{11} = r_{10} = r_6 = r_5 \quad (7.22) \]

\[ r_{12} + r_{13} = r_{10} + 6r_{11} = 7r_5 \quad (7.23) \]

Selectivity of \( \text{C}_6\text{H}_5\text{-CH}_2^+ \) to \( \text{CO} \) can be defined as:

\[ S_{\text{CO}} = \frac{r_{13}}{7r_5} \quad (7.24) \]

Inserting (7.24) into (7.23):

\[ r_{12} = 7(1 - S_{\text{CO}})r_5 \quad (7.25) \]

By substituting Eq. (7.17), (7.20), (7.21), (7.22), and (7.25) into Eq. (7.14), the rate of ozone decomposition by step (1) in Table 7.5 will be:

\[ r_1 = [11 + 7(1 - S_{\text{CO}})]r_5 = v_{\text{ozone}}r_5 \quad (7.26) \]
\(v_{\text{ozo}}\) is ozone theoretical stoichiometric coefficient which is expressed by:

\[v_{\text{ozo}} = 11 + 7(1 - S_{\text{CO}})\]  
(7.27)

\(r_1\) and \(r_5\) are defined by Eq. (7.28) and (7.29), respectively:

\[r_1 = k_1 P_{\text{ozo}} \theta^*\]  
(7.28)

\[r_5 = k_5 \theta_{\text{C}_6\text{H}_5-\text{CH}_3^*} \theta^*\]  
(7.29)

\(\theta_{\text{C}_6\text{H}_5-\text{CH}_3^*}\) can be determined from step (4) in Table 7.5 by using pressure equilibrium constant of toluene:

\[\theta_{\text{C}_6\text{H}_5-\text{CH}_3^*} = K_{\text{tol}} P_{\text{tol}} \theta^*\]  
(7.30)

Combining Eq. (7.28), (7.29), and (7.30) with Eq. (7.26), we have:

\[r_1 = k_1 P_{\text{ozo}} \theta^* = v_{\text{ozo}} r_5 = v_{\text{ozo}} k_5 K_{\text{tol}} P_{\text{tol}} \theta^{*2}\]  
(7.31)

Eq. (7.31) can be used to determine \(\theta^*\):

\[\theta^* = \frac{k_1 P_{\text{ozo}}}{v_{\text{ozo}} k_5 K_{\text{tol}} P_{\text{tol}}}\]  
(7.32)

Substitution of Eq. (7.32) into Eq. (7.31), the rate of toluene oxidation becomes:

\[r_{\text{tol}} = r_5 = \frac{r_1}{v_{\text{ozo}}} = \frac{k_1^2 P_{\text{ozo}}^2}{v_{\text{ozo}}^2 k_5 K_{\text{tol}} P_{\text{tol}}} = k_{\text{eff}}^{-1} P_{\text{tol}}^{-1} P_{\text{ozo}}^2\]  
(7.33)

\[k_{\text{eff}} = \frac{k_1^2}{v_{\text{ozo}}^2 k_5 K_{\text{tol}}}\]  
(7.34)

Eq. (7.33) is the rate equation of the LH2 mechanism showing that order of the reaction is -1 and 2 with respect to toluene and ozone, respectively. The rate equation of the LH2 model is the same as the rate expression obtained by the power law model via Eq. (7.8). Both models result in the same reaction orders for toluene and ozone. Therefore, one can estimate \(k_{\text{eff}}\) from the value
of pre-exponential factor ($k_o$) and the activation energy ($E$) reported in Table 7.1 for the power law model. Eq. (7.33) also shows that the rate of toluene oxidation is proportional to the square of $k_1$, indicating that higher rates of ozone decomposition accelerates the rate of toluene oxidation in agreement with the findings of Chapters 5 and 6. It can be seen that LH2 mechanism can be considered as the second Langmuir-Hinshelwood mechanism, proposing the elementary steps involved in oxidation of toluene. However, LH2 mechanism has been suggested based on what has been reported in the literature for similar reactions and more experimental and theoretical details are needed to prove the validity of the LH2 mechanism.

7.4. Summary

Chapter 7 presents a study on the kinetics and mechanism of catalytic oxidation of toluene by ozone. Kinetics data were analyzed based on the initial reaction rates. It was revealed that increasing the toluene partial pressure reduced the reaction rate while higher ozone partial pressures increased the rate of toluene oxidation. The apparent activation energy of the reaction was determined to be $31 \text{ kJ mol}^{-1}$, explaining the low temperature activity of the catalyst in the presence of ozone. The power law reaction orders were -1 and 2 for toluene and ozone, respectively.

A Langmuir-Hinshelwood mechanism (LH1) was used to explain the elementary steps involved in the oxidation of toluene by ozone. This mechanism was originally proposed for the catalytic oxidation of acetone by ozone over silica supported manganese oxides. Results showed that two surface oxygen atoms might be involved to start the oxidation of toluene. It was also observed that the concentration of surface peroxide species was low and these species could be considered as spectators in the reaction as it was suggested by Oyama and co-workers [51].
However, LH1 model resulted in a positive value for the standard entropy of toluene adsorption which is not commonly observed in catalytic reactions.

Therefore, a second Langmuir-Hinshelwood mechanism (LH2) was proposed to introduce an alternative approach to the reaction mechanism. The LH2 model explained the oxidation mechanism in more detail which was developed by considering the C-H bond cleavage from the methyl group. It was assumed that the adsorbed toluene molecule was dehydrogenated first to produce water vapor. The hydrogen free carbon structure was further oxidized to produce CO and CO$_2$. The LH2 model resulted in a simple power law model with the reaction orders of -1 and 2 for toluene and ozone, respectively. In fact, the LH2 mechanism is the same as the power law model with the same reaction orders.

The judgment about rejecting one of these models (LH1 or LH2) is possible by evaluating the reaction rates at toluene partial pressures close to zero while ozone partial pressure is kept constant. Basically, the LH1 mechanism predicts the rate of toluene oxidation to approach zero when the partial pressure of toluene decreases to zero. On the other hand, the LH2 (the power law model) predicts large reaction rates when toluene partial pressure is moving toward zero. However, the discrimination between the two models has not been carried out in this research work due to analytical limitations. Nevertheless, the goal of Chapter 7 was to introduce an alternative mechanism scheme which could also be considered to explain the mechanism of toluene oxidation. It is clear that further evidence is needed to fully understand the mechanism of this reaction which can be the subject of a future work.
8.1. Summary of the thesis

Catalytic ozonation of toluene by ozone over alumina supported manganese oxides was investigated in this thesis. The motivation behind this research was studying the effectiveness of catalysts in low temperature oxidation of indoor VOCs, contributing to sick building syndrome. It was concluded in Chapter 2 that sorption filtration was the most efficient method for indoor air purification. However, deactivation was reported in sorption filtration, making the regeneration of adsorbents necessary. Other methods such as photo-catalytic oxidation (PCO) and non-thermal plasma (NTP) needed more study in order to overcome the deficiencies related to these methods. Oxidation could be considered as an alternative method for the removal of VOCs from air. However, homogenous oxidation of VOCs was considered to be highly energy consuming, occurring at elevated temperatures. Catalysis might be applied to oxidation reactions in order to lower the reaction temperature. More energy saving in oxidation processes could also be achieved by replacing oxygen with ozone and using appropriate catalysts. A wide range of molecules such as ethanol, acetone, benzene, cyclohexane, formaldehyde, and toluene was oxidized by this method. Results showed that it was possible to convert these pollutants to CO, CO$_2$, and water vapor in the temperature range of 22-100 °C, depending on the reaction conditions. In addition to the energy saving, catalytic ozonation of VOCs had the advantage of using transition metals which were cheaper than noble metals used in the catalytic oxidation of VOCs by oxygen. Manganese oxides were reported to be the most active transition metal oxides in the decomposition of ozone and the oxidation of VOCs. The high activity of manganese oxides was related to the capability of these oxides to have different oxidation states during
catalytic cycle of VOCs oxidation. Therefore, the current research work was dedicated to study different aspects of catalytic oxidation of VOCs by ozone. Toluene was selected as the model compound since it was reported to be one of the major indoor air VOCs. In addition, there were a few reports on the catalytic oxidation of toluene by ozone published in the literature, making toluene to be a suitable model compound. The catalyst of interest was alumina supported manganese oxides due to the high activity of these oxides in ozonation reactions.

Oxidation of toluene was performed in a continuous fixed bed reactor. The pressure of the reactor was atmospheric and the temperature range was between 22-100 °C. Typical concentrations of ozone and toluene were 1050 and 120 ppm, respectively. The method of catalyst preparation was dry impregnation. The detail of the experimental set-up, analysis, and the catalyst preparation was given in Chapter 3. Catalyst characterization and activity tests were presented in Chapter 4. Characterization techniques such as XRD and laser Raman spectroscopy showed that MnO$_2$ and Mn$_2$O$_3$ were formed on the surface of the catalyst with 10% Mn loading. The weight percent of MnO$_2$ and Mn$_2$O$_3$ were determined by XANES analysis to be 9.0 and 91.0%, respectively. Activity tests indicated that the catalyst deactivated at room temperature. It was observed that 65 °C was the minimum temperature, at which the catalyst worked without deactivation. The main cause of the catalyst deactivation was the accumulation of carbonaceous species on the surface of the catalyst due to the incomplete oxidation of toluene. FT-IR analysis identified alcohols and carboxylic acids as the deposited species on the surface of the catalyst. Near 95% toluene conversion was achieved at 80 °C with CO and CO$_2$ as the major reaction products, resulting in 90% carbon balance. The minor by-products of the reaction were determined to be oxalic acid, acetic acid, benzene, and maleic anhydride.
Chapter 5 studied the effect of Mn loading (1, 5, 10, and 20%) on the oxidation of toluene by ozone. X-ray absorption spectroscopy including XANES and EXAFS was used in order to relate the electronic properties and the structure of the catalysts to Mn loading. Results showed that the catalysts with Mn loading less than 10% mostly contained Mn$_2$O$_3$ while a mixture of MnO$_2$ and Mn$_2$O$_3$ was observed at higher Mn loading. It was also found that the formal oxidation state of Mn increased with the increase of catalyst loading. EXAFS studies were first carried out on the reference materials (MnO$_2$ and Mn$_2$O$_3$) in order to estimate the amplitude reduction factors ($S_o^2$). EXAFS of Mn$_2$O$_3$ was studied in more detail due to the complicated unit cell structure of Mn$_2$O$_3$. Two methods based on the Bond Valence Model and averaging the Mn sites based on their site occupancies were used to construct Mn$_2$O$_3$ theoretical EXAFS signal. Fitting the results showed that the Bond Valence Model and the averaging method converged on the same fitting parameters. In fact, the Bond Valence Model was considered as the general method to model the EXAFS signal of complex crystals such as Mn$_2$O$_3$. Nevertheless, the Bond Valence Model proved the validity of the averaging method, simplifying the structure of Mn$_2$O$_3$ into one single Mn sites. Activity tests showed that the catalysts with lower Mn loading and lower Mn oxidation state had higher rate of toluene oxidation. It was suggested that lower oxidation states of Mn were more favorable in the reaction. This observation was related to the fact that Mn atoms at lower oxidation states were more capable in transferring electrons to ozone, accelerating the rate of ozone decomposition which eventually enhanced the rate of toluene oxidation.

Promotional effect of the noble metals on activity of MnO$_x$/$\gamma$-alumina (10%) was studied in Chapter 6. Two noble metals with loading of 1% were added to manganese oxides to decrease the oxidation temperature of toluene. XANES and EXAFS were applied in order to elucidate
inter-atomic interactions between the noble metals and manganese oxides. Activity tests indicated that the addition of Pt increased the activity of MnOₓ/γ-alumina (10%) by lowering the temperature of maximum toluene conversion to 70 °C. Pd did not improve the catalyst activity, resulting in maximum toluene conversion at 80 °C. The characterization of the catalysts showed that Pt is mostly in metallic state on alumina. The oxidation state of Pt increased after the addition of Pt to Mn, showing that an atomic interaction occurred between Pt and Mn. On the other hand, Pd was present as PdO in the presence and absence of Mn. Results of EXAFS also showed the changes in oxygen and Pt coordination numbers after the addition of Pt to Mn. No changes in EXAFS of Pd were observed after the addition of Pd to Mn, showing that most probably there is no interaction between Pd and Mn atoms. It was concluded that Pt was attached to manganese oxides through the surface oxygen atoms of Mn₂O₃ clusters, forming Pt-O-Mn bonds. It was suggested that the higher activity of Pt-Mn bimetallic catalyst was related to the electronic effect of Pt on Mn by increasing Mn 3d electron occupancy. This was in agreement with the effect of Mn loading, showing that lower oxidation states of Mn were more favorable in the oxidation of toluene. Activity of the reduced catalysts with hydrogen supported the idea that Mn atoms with less degree of oxidation were more effective in the oxidation of toluene.

The kinetics and mechanism of the catalytic oxidation of toluene by ozone over manganese oxides were studied in Chapter 7. Initial reaction rates were obtained in order to fit different kinetic models to the experimental data. Power law model determined the apparent activation energy of the reaction to be 31.0 kJ mol⁻¹, explaining the low temperature activity of the catalyst. Reaction orders with respect to toluene and ozone were -1 and 2, respectively. This suggested that toluene was oxidized faster at lower toluene partial pressures while reaction rates increased by the increase of ozone partial pressure. A Langmuir-Hinshelwood mechanism (LH1)
was considered in order to fit the experimental data. This model was originally developed by Oyama and co-workers in order to explain the mechanism of acetone oxidation by ozone over silica supported manganese oxides. LH1 showed that most likely peroxide species were not involved in the reaction mechanism due to the low surface coverage of peroxide species. It was also observed that 2 oxygen atoms were necessary to start the oxidation reaction. However, the entropy of toluene adsorption turned out to be positive by LH1 mechanism. The positive value of the adsorption entropy is not recommended in catalytic reactions, indicating that LH1 mechanism could not completely explain the mechanism of the reaction. Therefore a second Langmuir-Hinshelwood mechanism (LH2) was proposed as an alternative to LH1 mechanism. LH2 mechanism was built based on the abstraction of one hydrogen atom from the methyl group by the adsorption of toluene on one pair of active sites. It was assumed that toluene molecule was dehydrogenated first by atomic oxygen to produce a hydrogen free carbon structure. The carbon structure further reacted with atomic oxygen to produce CO and CO$_2$. It was shown that LH2 mechanism resulted in a power law model with the reaction orders of -1 and 2 with respect to toluene and ozone, respectively. The reaction orders were the same as the power law model obtained from the experimental data, explaining the relationship of the actual reaction rates and LH2 mechanism. Nevertheless, the goal of Chapter 7 was to look at the mechanism of the reaction from a new angle, introducing alternative models which might justify the reaction mechanism. More evidence was obviously needed to fully understand the mechanism of this reaction which could be the subject of future works.

8.2. Conclusions

The followings are the general conclusions of the thesis based on the objectives defined in Chapter 2:
• The active phase of the catalyst (MnO\textsubscript{x}/\gamma-alumina) is made of MnO\textsubscript{2} and Mn\textsubscript{2}O\textsubscript{3} effective in the oxidation of toluene below 100 °C. The main by-products of the catalytic ozonation of toluene are CO and CO\textsubscript{2}. Catalyst deactivation occurs at room temperature due to the accumulation of carbonaceous species on the catalyst. At least, 65 °C is required for the stable operation of the catalyst.

• EXAFS signal of Mn\textsubscript{2}O\textsubscript{3} can be modeled using the Bond Valence Model, proposing a general approach for EXAFS analysis of transition metal oxides with more than one absorbing atom in their unit cell structure. A direct relationship exists between the oxidation state and the loading of manganese on alumina. Mn\textsubscript{2}O\textsubscript{3} is the dominant phase at Mn loadings lower than 10% while more MnO\textsubscript{2} is formed at higher loadings. Lower Mn loadings are more favorable in the oxidation of toluene by ozone.

• EXAFS and XANES show that Pt is attached to manganese oxides through surface oxygen atoms of Mn\textsubscript{2}O\textsubscript{3} clusters, changing the atomic environment around Mn. The net effect is the oxidation of Pt atoms and a slight increase in Mn coordination number, enhancing the oxidation rate of toluene by ozone. No atomic interaction is found between Pd and Mn, explaining why the addition of Pd does not promote the activity of Mn\textsubscript{2}O\textsubscript{3} in the oxidation of toluene.

• It is concluded form the activity of different loadings of Mn and the addition of Pt to Mn that the catalytic oxidation of toluene can be promoted by decreasing the oxidation state of Mn. The bottleneck of the oxidation reaction seems to be the rate of ozone decomposition, controlling the overall rate of toluene oxidation.

• The kinetic study of the catalytic ozonation of toluene shows that the activation energy of toluene oxidation can be greatly reduced using ozone and MnO\textsubscript{x}/\gamma-alumina.
This explains the low temperature activity of the catalyst comparing to that of oxygen which proceeds at around 200 °C. It is also proposed that the oxidation of toluene starts with the abstraction of hydrogen atoms and production of water. The mechanism is followed by the oxidation of the carbon skeleton of toluene, leading to the production of CO and CO₂.

8.3. **Recommendations**

Even though the result of this thesis and other works reported in the literature prove the effectiveness of catalytic ozonation, there are still open questions related to this method. Deactivation of the catalysts at room temperature is in fact a limiting factor for operating the catalyst at ambient temperature. It has been reported that the deactivation of manganese oxides in the oxidation of formaldehyde and benzene can be prevented in the presence of water vapor [60, 61]. The support type also plays an important role in the long term activity of catalysts. Einaga et al. showed that ultra stable Y (USY) zeolite was stable for the oxidation of benzene in the presence of water vapor at room temperature [22]. Therefore, it is recommended to investigate the effect of water vapor using different supports in order to obtain the room temperature activity of manganese oxides. Understanding the mechanism of catalyst deactivation at room temperature can also provide fundamental knowledge about the interaction of reaction by-products and the catalyst surface. This can explain the accumulation of carbonaceous species such as alcohols and carboxylic acids produced as the result of incomplete oxidation of toluene.

Studying the kinetics and mechanism of the reaction is another important research area which can be explored in more details. It was shown in Chapter 7 that different molecules most likely follow different elementary steps in the catalytic oxidation of VOCs by ozone. An alternative mechanism (LH2) was introduced to the widely used mechanism proposed by Oyama.
and co-workers (LH1). LH2 mechanism was basically developed based on the activation of the adsorbed toluene molecule by dehydrogenation, following the oxidation of the carbon structure. In order to show the validity of LH2 mechanism over LH1 mechanism, it is possible to track the reaction rates at wider toluene partial pressures at a constant ozone partial pressure. Fig. 8.1 shows the trend of toluene oxidation rate predicted by LH1 and LH2 mechanisms as a function of toluene partial pressure.

It can be seen in Fig. 8.1 (A) that LH1 mechanism predicts zero reaction rates at toluene concentration close to zero. By the increase of toluene partial pressure, the reaction rates pass through a maximum and approach zero at high toluene partial pressures. On the other hand, Fig. 8.1 (B) shows the reaction rates predicted by LH2 mechanism. It can be seen that the reaction rate increases by the decrease of toluene partial pressure and reaches high values at toluene concentrations close to zero. Therefore, it is possible to check the validity of LH1 and LH2 mechanisms by tracking the reaction rates at low toluene partial pressures. This requires accurate analytical devices such as an on-line FT-IR gas analyzer which might become available in the future to test the validity of these mechanisms. In addition to this approach, it is also possible to study the validity of LH2 mechanism by other techniques such as isotopic exchange, density functional theory (DFT) calculations and in-situ spectroscopy to elucidate the elementary steps involved in the catalytic oxidation of VOCs by ozone.
Finally, technology development and reactor scale up can be considered if there is an industrial interest in this method. Catalytic ozonation can be integrated with HVAC systems as standalone units for the removal of VOCs from indoor environment. There is still need to explore
the effectiveness of this method in a real indoor air composition, consisting of a mixture of VOCs. Fine-tuning the reactions condition and the catalyst seems to be necessary for the treatment of actual air.
LIST OF REFERENCES


70. E. Park, S. Chin, J. Kim, G.-N. Bae, J. Jurng, Preparation of MnO_s/TiO_2 ultra fine nanocomposite with large surface area and its enhanced toluene oxidation at low temperature, Powder Technol. 208 (2011) 740-743.


APPENDIX A: GC-MS CALIBRATION DATA

Tables A.1 to A.3 contain the TIC peak area at each toluene, CO, and CO₂ concentrations, respectively. Measurements were repeated three times. Figures A.1 to A.3 show the GC-MS calibration curves of toluene, CO, and CO₂, respectively.

Table A.1. Toluene TIC peak area

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<th>Standard Deviation</th>
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Table A.3. CO\textsubscript{2} TIC peak area

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Fig. A.1. Toluene calibration curve; error bars are standard deviations.

Fig. A.2. CO calibration curve; error bars are standard deviations.
Fig. A.3. CO$_2$ calibration curve; error bars are standard deviations.
APPENDIX B: EFFECT OF PARTICLE SIZE ON THE CATALYST ACTIVITY

0.2 g of MnOx/γ-alumina (10%) with different particle sizes (dp) of dp<80 μm, 80 μm< dp<208 μm, 208 μm< dp<355 μm and 355 μm< dp<417 μm were tested in the temperature range of 22-100 °C to select the appropriate particle size which did not impose internal mass transfer limitation on the reaction rate. The flow rate of toluene/nitrogen and ozone/oxygen streams were 400 and 600 ml min⁻¹, respectively which resulted in a weight hour space velocity (WHSV) of 300 l hr⁻¹ g⁻¹. This combination gave toluene and ozone concentration of 120 and 1050 ppm, respectively. The mole percent of oxygen and nitrogen were 59.94 and 39.94%, respectively.

Activity of the catalyst with different particle size ranges are shown in Fig. B.1. Results show that by the increase of particle size to higher than 208 μm, the activity of the catalyst in toluene oxidation decreases. Therefore, it is decided to use a particle size with diameter less than 208 μm for the reaction rate studies.
Fig. B.1. Effect of particle size ($d_p$) on activity of MnO$_x$/γ-alumina (10%), (1) $d_p<80$ μm, (2) 80 μm $< d_p<208$ μm, (3) 208 μm $< d_p<355$ μm, (4) 355 μm $< d_p<417$ μm.
Mear’s equation can be expressed as [124]:

\[
- \frac{r_A \rho_b R_n}{k_c C_A} < 0.15
\]  \hspace{1cm} (C.1)

in which,

\( n \) = reaction order
\( R \) = radius of the catalyst particle (m)
\( \rho_b \) = catalyst bed bulk density (kg m\(^{-3}\))
\( C_A \) = concentration of A in the gas phase (kg mol m\(^{-3}\))
\( r_A \) = reaction rate of A (kg mol kg\(^{-1}\) s\(^{-1}\))
\( k_c \) = external mass transfer coefficient (m s\(^{-1}\))

\( k_c \) around a spherical particle can be estimated by [124]:

\[
Sh = 2 + 0.6 \text{Re}^{1/2} \text{Sc}^{1/3}
\]  \hspace{1cm} (C.2)

Sherwood number (Sh), Schmidt number (Sc), and Reynolds number (Re) are expressed by equations (C.3), (C.4), and (C.5), respectively:

\[
Sh = \frac{k_c d_p}{D_{AB}}
\]  \hspace{1cm} (C.3)

\[
Sc = \frac{\mu_g}{\rho_g D_{AB}}
\]  \hspace{1cm} (C.4)

\[
Re = \frac{U_d d_p}{\mu_g}
\]  \hspace{1cm} (C.5)
\( d_p \) (m) and \( D_{AB} \) (m\(^2\) s\(^{-1}\)) are the particle diameter and the diffusivity coefficient of A in B, respectively. \( \mu_g \) is the viscosity of the gas phase in kg m\(^{-1}\) s\(^{-1}\). U is superficial velocity in m s\(^{-1}\). \( \rho_g \) represents the gas phase density in kg m\(^{-3}\).

The following information is used as the typical conditions of the reactor to examine the effect of the external mass transfer:

Gas composition: 40% nitrogen and 60% oxygen

Total gas flow rate: 1000 ml min\(^{-1}\)

Reactor internal diameter: 1/4 in

Temperature: 60 °C

Gas viscosity (\( \mu_g \)): 2.1 \( \times \) 10\(^{-5}\) kg m\(^{-1}\) s\(^{-1}\)

Gas density (\( \rho_g \)): 1.1 kg m\(^{-3}\)

Catalyst particle diameter (\( d_p \)): 208 \( \times \) 10\(^{-6}\) m

Toluene diffusivity coefficient (\( D_{AB} \)): 9.9 \( \times \) 10\(^{-6}\) m\(^2\) s\(^{-1}\)

Using the above information along with Eq. (C.2) to (C.5), one can show that the external mass transfer coefficient can be found as 0.18 m\(^2\) s\(^{-1}\).

The conversion of toluene over 0.2 g of the catalyst at 60 °C is about 80% for the inlet toluene concentration of 120 ppm. Therefore, it is reasonable to assume an average toluene concentration of 70 ppm (2.6 \( \times \) 10\(^{-6}\) kg mol m\(^{-3}\)) at 60 °C inside the reactor. For 80% toluene conversion with the inlet toluene concentration of 120 ppm, the average rate of the reaction is 3.3 \( \times \) 10\(^{-7}\) kg mol kg\(^{-1}\) s\(^{-1}\). By considering the reaction order of one with respect to toluene and bulk density of 500 kg m\(^{-3}\) for the catalyst bed, the Mear's criterion is satisfied as 0.037<0.15, showing that external mass transfer does not control the rate of toluene oxidation.
APPENDIX D: RAW DATA OF THE CATALYTIC OXIDATION OF TOLUENE BY OZONE

Toluene was oxidized over 0.2 g of MnO$_x$/γ-alumina (10%) and the experiment was repeated in order to evaluate the repeatability of the data. Inlet concentration of toluene and ozone were 120 and 1050 ppm, respectively. Reaction conditions have been presented in section 3.3. Tables D.1 and D.2 represent the concentrations and conversions of toluene and ozone at room temperature, respectively. Concentrations and conversions of toluene and ozone at higher temperature are included in Tables D.3 and D.4, respectively.

Table D.1. Toluene concentration and conversion at room temperature

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>6</th>
<th>41</th>
<th>76</th>
<th>118</th>
<th>150</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outlet toluene concentration (ppm)</td>
<td>Run #1</td>
<td>1</td>
<td>14</td>
<td>37</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>Run #2</td>
<td>1</td>
<td>17</td>
<td>35</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>Average (ppm)</td>
<td>1</td>
<td>16</td>
<td>36</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>Standard deviation (ppm)</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Toluene conversion (%)</td>
<td>Run #1</td>
<td>99</td>
<td>88</td>
<td>69</td>
<td>56</td>
</tr>
<tr>
<td></td>
<td>Run #2</td>
<td>99</td>
<td>86</td>
<td>71</td>
<td>58</td>
</tr>
<tr>
<td></td>
<td>Average (%)</td>
<td>99</td>
<td>87</td>
<td>70</td>
<td>57</td>
</tr>
<tr>
<td></td>
<td>Standard deviation (%)</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>2</td>
</tr>
</tbody>
</table>
Table D.2. Ozone concentration at conversion at room temperature

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>6</th>
<th>41</th>
<th>76</th>
<th>118</th>
<th>150</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run #1</td>
<td>0</td>
<td>285</td>
<td>553</td>
<td>700</td>
<td>800</td>
</tr>
<tr>
<td>Run #2</td>
<td>30</td>
<td>300</td>
<td>515</td>
<td>670</td>
<td>770</td>
</tr>
<tr>
<td>Average (ppm)</td>
<td>15</td>
<td>293</td>
<td>534</td>
<td>685</td>
<td>785</td>
</tr>
<tr>
<td>Standard deviation (ppm)</td>
<td>2</td>
<td>11</td>
<td>27</td>
<td>21</td>
<td>21</td>
</tr>
</tbody>
</table>

Table D.3. Toluene concentration and conversions at higher temperatures

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>40</th>
<th>60</th>
<th>80</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run #1</td>
<td>59</td>
<td>18</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Run #2</td>
<td>54</td>
<td>23</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>Average (ppm)</td>
<td>56</td>
<td>20</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Standard deviation (ppm)</td>
<td>4</td>
<td>4</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Run #1</td>
<td>51</td>
<td>85</td>
<td>92</td>
<td>92</td>
</tr>
<tr>
<td>Run #2</td>
<td>55</td>
<td>81</td>
<td>94</td>
<td>93</td>
</tr>
<tr>
<td>Average (%)</td>
<td>53</td>
<td>83</td>
<td>93</td>
<td>93</td>
</tr>
<tr>
<td>Standard deviation (%)</td>
<td>3</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>
Table D.4. Ozone concentration and conversion at higher temperatures

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>40</th>
<th>60</th>
<th>80</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outlet ozone concentration (ppm)</td>
<td>Run #1</td>
<td>685</td>
<td>239</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Run #2</td>
<td>695</td>
<td>259</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>Average (ppm)</td>
<td>690</td>
<td>249</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>Standard deviation (ppm)</td>
<td>7</td>
<td>14</td>
<td>37</td>
</tr>
<tr>
<td>Ozone conversion (%)</td>
<td>Run #1</td>
<td>35</td>
<td>77</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Run #2</td>
<td>34</td>
<td>75</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>Average (%)</td>
<td>34</td>
<td>76</td>
<td>98</td>
</tr>
<tr>
<td></td>
<td>Standard deviation (%)</td>
<td>1</td>
<td>1</td>
<td>4</td>
</tr>
</tbody>
</table>
APPENDIX E: UNCERTAINTIES OF BET, ICP-MS, AND CHNS ANALYSES

MnOx/γ-alumina (10%) was selected to show the uncertainties of BET, ICP-MS, and CHNS analyses. Carbon content of 0.2 g of the same catalyst was measured after 150 minutes oxidation of toluene at room temperature by ozone. Inlet toluene and ozone concentrations were 120 and 1050 ppm, respectively. BET and ICP-MS data is presented in Table E.1 while Table E.2 shows the CHNS data.

Table E.1 MnOx/γ-alumina (10%) BET and ICP-MS data

<table>
<thead>
<tr>
<th>BET surface area (m² g⁻¹)</th>
<th>Run #1</th>
<th>171</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Run #2</td>
<td>187</td>
</tr>
<tr>
<td></td>
<td>Run #3</td>
<td>172</td>
</tr>
<tr>
<td>Average (m² g⁻¹)</td>
<td></td>
<td>177</td>
</tr>
<tr>
<td>Standard deviation (m² g⁻¹)</td>
<td></td>
<td>9</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pore volume (cm³ g⁻¹)</th>
<th>Run #1</th>
<th>0.51</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Run #2</td>
<td>0.52</td>
</tr>
<tr>
<td></td>
<td>Run #3</td>
<td>0.49</td>
</tr>
<tr>
<td>Average (cm³ g⁻¹)</td>
<td></td>
<td>0.51</td>
</tr>
<tr>
<td>Standard deviation (cm³ g⁻¹)</td>
<td></td>
<td>0.02</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mn loading (wt%)</th>
<th>Run #1</th>
<th>9.8</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Run #2</td>
<td>9.2</td>
</tr>
<tr>
<td>Average (wt%)</td>
<td></td>
<td>9.5</td>
</tr>
<tr>
<td>Standard deviation (wt%)</td>
<td></td>
<td>0.4</td>
</tr>
</tbody>
</table>
Table E.2. Deposited carbon content of MnO$_x$/$\gamma$-alumina (10\%) at room temperature

<table>
<thead>
<tr>
<th>Carbon content (wt%)</th>
<th>Run #1</th>
<th>11.29</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Run #2</td>
<td>9.79</td>
</tr>
<tr>
<td></td>
<td>Run #3</td>
<td>9.67</td>
</tr>
<tr>
<td>Average (wt%)</td>
<td></td>
<td>10.25</td>
</tr>
<tr>
<td>Standard deviation (wt%)</td>
<td></td>
<td>0.90</td>
</tr>
</tbody>
</table>
APPENDIX F: REDUCTION OF BIMETALLIC CATALYSTS BY HYDROGEN; XRD AND ACTIVITY

Pt-MnOₓ/γ-alumina and Pd-MnOₓ/γ-alumina were reduced by pure hydrogen at 250 °C for 2 hrs. The XRD spectra of the reduced catalysts are shown in Fig. F.1. It can be seen that all Mn₂O₃ and MnO₂ are transformed to MnO after the reduction.

![XRD spectra of reduced catalysts by hydrogen](image)

Fig. F.1. XRD spectra of reduced catalysts by hydrogen, (a) Pd-MnOₓ/γ-alumina, (b) Pt-MnOₓ/γ-alumina, *: γ-alumina, ○: Platinum, ▪: MnO.

The activity of the reduced and reduced catalysts are indicated in Fig. F.2 and Fig. F.3 in terms of toluene and ozone conversions, respectively. It can be seen that the reduced Pd-MnOₓ/γ-alumina show almost the same activity as Pt-MnOₓ/γ-alumina, implying that lower oxidation state of manganese (higher 3d electron occupancy) are more favorable in the oxidation of
toluene. It has to be noted that room temperature activities (22 °C) are reported after 150 min running the reaction.

Fig. F.2. Toluene conversion, (a) Pt-MnO\textsubscript{x}/γ-alumina (unreduced), (b) Pd-MnO\textsubscript{x}/γ-alumina (unreduced), (c) Pd-Mn/γ-alumina (reduced), and (d) Pt-MnO\textsubscript{x}/γ-alumina (reduced).
Fig. F.3. Ozone conversion, (a) Pt-MnO$_x$/γ-alumina (unreduced), (b) Pd-MnO$_x$/γ-alumina (unreduced), (c) Pd-Mn/γ-alumina (reduced), and (d) Pt-MnO$_x$/γ-alumina (reduced).