Platinum- and Palladium-Based Single-Atom Alloy Catalysts for Selective Hydrogenation and Dehydrogenation Reactions

A dissertation submitted by

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in partial fulfillment of the requirements for the degree of

Doctor of Philosophy

in

Chemical Engineering

Tufts University

May 2018

Advisor:

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Abstract

Approximately 80% of all chemical, materials and fuel production in the modern industry use heterogeneous catalysts during some phases in the process. Platinum group metals, especially Pt or Pd are often used in heterogeneous catalysts but susceptible to unselective reactivity, deactivation, and poisoning. The industrial Pt and Pd catalysts, although very successful in many cases, are basically “black boxes” without well controlled structural and compositional properties. Different additives are used to modify the metal nanoparticle catalysts on the support to form the multiple metallic catalysts with high precious metal loading. Some toxic poisons, such as Pb or V, are often used to improve the selectivity of these catalysts. Due to these drawbacks, a rational catalyst design approach to make Pt and Pd catalysts more efficient and cleaner is of great interest.

The major challenges in Pt and Pd catalysts are their high price, unselective reactivity, carbon deposition and CO poisoning. This thesis addresses these problems with a novel single-atom alloy approach, where isolated Pt atoms, stabilized in Cu nanoparticle surface, are identified as the active sites for key elementary reactions. The Pt metal was predesigned with minimum atomic ensembles and loaded in the surface of cheaper metal nanoparticles to achieve the best possible selectivity, stability and reactivity and reduced concentration of precious metals. This class of single-atom alloy catalysts were applied in catalyzing industrially important selective hydrogenation and dehydrogenation reactions and thoroughly studied for their surface chemistry and structural and compositional properties.
Pt-Cu single-atom alloy nanoparticle catalysts were prepared with the galvanic replacement method and supported on alumina and silica. With comprehensive characterization, the formation of single-atom alloy nanoparticles was for the first time demonstrated in the field. This class of catalysts showed excellent performance in selective hydrogenation of 1,3-butadiene, selective dehydrogenation of butane and CO tolerant hydrogen activation reactions. By depositing a small amount of Pt single-atoms in the surface of Cu nanoparticles, the reactivity in selective hydrogenation and dehydrogenation reactions is improved more than one order of magnitude. Notably, close to 100% selectivity to desired partial hydrogenation and dehydrogenation products are also achieved. The PtCu single-atom alloy catalysts are stable in reaction conditions without carbon deposition or sintering.

In these processes, the single-atom Pt is efficient in catalyzing key elementary reactions including H₂ dissociation and C-H activation, while the highly selective nature of Cu remains unaffected. The energy barriers of these elementary reaction steps are significantly lowered with the single-atom Pt sites. But single-atom Pt sites bind CO and the product molecules weakly compared to monometallic Pt nanoparticles, which results in CO tolerance and high selectivity.

The suitability of the single-atom approach in PdAu catalysts was also studied. The PdAu single-atom alloy nanoparticle catalysts were successfully synthesized with sequential reduction methods and demonstrated to be highly selective, stable and reactive in partial hydrogenation of 1-hexyne in the liquid phase.
This thesis work uses an integrated approach, including catalyst preparation, advanced characterization, catalyst evaluation under realistic conditions. Collaborative surface science information and DFT calculations complement the catalyst. This provides the platform materials that bridge the materials gap between catalysis and surface science studies. Moreover, it highlights a rational catalyst design approach to optimize the catalytic performance at the single-atom limit.
Acknowledgements

“A journey of a thousand miles begins with a single step.”

-Tao Te Ching, 500 B.C.

My graduate study at Tufts would not have been so enjoyable and productive without the mentors, teachers, co-workers, collaborators, friends and family I have had the pleasure of working or living with. I am very grateful to all of them.

First, I would like to express my deep appreciation to my advisor, Prof. Maria Flytzani-Stephanopoulos, for offering me the opportunity to engage in this lab and for being an outstanding mentor and teacher. The reason I came to Tufts University was to work with her. Six years ago, I knocked on her door with no experience on heterogeneous catalysis but just the passion to explore. She accepted me to the lab and gave me the time to learn. Professor Flytzani-Stephanopoulos inspired me with brilliant ideas and taught me to incorporate creative thinking in my research. Her perspective on identifying and addressing important problems is critical to my PhD study. I highly cherish her guidance and advice that would always serve as inspiration for the personal and professional development in my life. It has been my great privilege and pleasure working with a P.I. with so much passion and enthusiasm for research and holding such a high standard for work ethic.

Professor Charles Sykes is a great mentor and collaborator who played an important role in my PhD research, for which I am truly grateful. I gained the critical fundamental chemistry insight through collaborating with him. I benefited a lot from his
unbelievable understanding of chemical and physical phenomena on surfaces in the last five years.

I would also like to thank Professor Terry Haas. The collaboration with him on XAS and XRD data analysis has been essential. I owe a great deal to my committee for providing critical insight into my work. Professor Charles Sykes, Professor Terry Haas, Professor Chia-Kuang (Frank) Tsung and Professor Matthew Panzer helped me a lot with their expertise in chemical engineering, chemistry and heterogeneous catalysis, and have given me valuable feedbacks. I want to thank Professor Howard Saltsburg for his valuable critique and comments on my early work.

I have been lucky to work with many exceptional graduate students during my PhD. I would like to thank Dr. Ioannis Valsamakis and Dr. Matthew Boucher for teaching and training me with the catalyst techniques when I joined the group. Dr. Ming Yang has been the model of hard working and critical thinking that has become a strong motivating force. Dr. Felicia Lucci, Dr. Matthew Marcinkowski, Dr. Andrew Therrien, Matthew Uhlman, Matthew Darby and Dr. Matthew Montemore were key to the collaboration with Sykes, Stamatakis and Kaxiras labs. I had the privilege of working with many great lab mates: Dr. Branko Zugic, Dr. Chongyang Wang, Dr. Junjun Shan, Mengwei Li, Sufeng Cao, Georgios Giannakakis, Dr. Antonios Trimpalis, Joshua Wimble, George Cladaras, Matthew Rutter, Dr. Nan Yi, Mengyao Ouyang, Yuan Wang. Without them, NCEL would not have been be such as a pleasant place to work in.

Many thanks to my undergraduate and high school students, Tingyi Dong, Nicholas Cicchetti, Ahmad Sukkar, Karen Li and Wesley Viola. Great appreciation to Beth
Frasso, Margaret DeChiaro and Emily Edward, whose tireless behind the scenes effort kept my research going smoothly.

Last, but foremost I want to thank my wife, Jin and my little daughter, my parents and in-laws. Thank you for being my support through the process, being the cheerleader when I felt difficult to carry on and being the most enthusiastic audiences for every little progress. I love you all.
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Chapter 1. Introduction

1.1 Heterogeneous catalysis

Catalysts are the substances that increase the rate of chemical reactions and maintain their physical and chemical properties in reaction conditions. Approximately 90% of all chemical, materials and fuel production in the modern industry use catalysts during some phases in the process. The majority of these processes (∼ 85%) use a heterogeneous catalyst, where the catalysts are in the solid phase and the reactants are in the liquid or gas phases. The heterogeneous catalysts are complex in their makeup and the specific role of each component in the reaction conditions. Understanding the structure-function relationships have been the focus of a large portion of the ongoing research in heterogeneous catalysis. The multidisciplinary collaboration of ambient/above ambient pressure catalysis studies with surface science and DFT techniques is a powerful approach to fill this gap.

The major motivation of understanding the structure-function relationships in heterogeneous catalysis is to guide the rational design of the catalysts. Catalysts with high activity, selectivity and durability are always desired. Catalysis is called to address the new challenges caused by the rapid development of human society. For example, the fast growth of the middle-class population in the world make the chemical processes that are more efficient, cleaner and environmentally benign more and more attractive, which requires the catalysts to be highly selective to target products and can be used in not too severe conditions. Additionally, the emerging “hydrogen economy” requires efficient fuel cells, which makes the
development of high performance and CO-tolerant fuel cell catalysts crucial. Moreover, the recent “shale gas boom” has put catalysts for coke resistant C-H activation and conversion into the center of the stage. For these cases, the selectivity in forming desired products is very important, if not the most important, as pointed out by Somorjai and McCrea 16 years ago:\(^5\)

“In the past, the focus of research in catalysis science was activity to optimize turnover rates. In the future, selectivity to form the desired product without the formation of byproducts will be the major research challenge.”

1.2 Selective hydrogenation of alkynes and dienes to alkenes

Selective hydrogenations lie in the heart of chemical industry. They are the key reactions in the production of bulk feed stocks for the polymer industry and the production of fine and specialty chemicals including food additives, pharmaceuticals, fragrances and agrochemicals. Specifically, the selective hydrogenation of alkyne and dienes to alkenes is important for the polymer industry. The alkene feedstock produced from cracking processes usually consists 1-8% highly unsaturated hydrocarbons, such as alkynes and dienes. However, the alkyne and diene concentration should be reduced to ppm level to avoid poisoning the polymerization catalysts. One attractive approach is to selectively hydrogenate these molecules to alkenes without further converting alkenes to alkanes.
1.2.1 Selective hydrogenation catalysts

Selective hydrogenations of both double and triple carbon-carbon bonds are important for a wide range of industrial processes. Pd, Pt, Ni, and Ru based systems are commonly employed for hydrogenation of alkenes and alkynes due to their high catalytic activity. However, they are not always selective to the desired products. Among these metals, Pd and Pt are the most selective, but typically exhibit a decrease in alkene selectivity with alkyne conversion. Therefore, Pd and Pt catalysts are commonly used with promoters to improve alkene selectivity; transition metals such as Ag, Rh, Au, Cu, Zn, Cr, and V have been reported to be effective selectivity promoters.

Industrial selective hydrogenation reactions are carried out on Pd-based catalysts with additives. For gas phase selective hydrogenation, Pd is alloyed with Au, Cu or Ag to improve the selectivity and durability and is supported on low-acidic support to avoid side reactions. For liquid phase selective hydrogenation, Pd is modified with Pb or Bi to form the Lindlar catalyst. To further improve the selectivity, CO or quinoline are mixed with the reactant in many processes. However, the wide use of Pd in the catalyst significantly increases the cost and the lead additives draw environmental concerns. New designs of Pt and Pd catalysts with high catalytic performance and minimal usage of precious Pt group metals are highly desired.

Pd and Pt have been extensively studied looking at reaction mechanisms, active intermediates, effect of nanoparticle size, effect of carbon deposits, effect of forming alloys with another metal, effect of additives, etc. In recent years, group IB metals Au and Cu, especially Au, have received attention for selective hydrogenation and were found to be very selective. However, they are much less active than group VIII metals in hydrogenation, lacking
in the activation of hydrogen.\textsuperscript{11–14} Even though Pd and Pt show high selectivity in hydrogenation, their selectivity is suppressed in the presence of excess alkenes, which are present under industrial hydrogenation conditions. Take butadiene hydrogenation for instance, in order to improve the selectivity, people have been investigating the hydrogenation activity and selectivity of bimetallic group VIII – IB metals, including Pd-Cu, Pd-Ag, Pd-Au, Pt-Cu, Pt-Au, etc.\textsuperscript{15–18} The change of the hydrogenation chemistry was attributed to the limited hydrogen chemisorption capability of the bimetallic catalysts, the change in adsorption strength of reactants and intermediates caused by electronic effects, and ensemble effects.\textsuperscript{15–20}

1.2.2 Hydrogenation reactions with Pd and Pt catalysts

The general reaction pathway of alkynes and diene hydrogenation is shown in scheme 1.1. The selectivity to alkenes is attributed to their relatively weak adsorption on the catalyst surface ($k_1/k_1 \gg k_3/k_3$). The well-known Horiuti-Polanyi mechanism assumes the hydrogenation of di-σ-bonded alkenes takes place in two steps through an adsorbed half-hydrogenated alkyl intermediate with atomic hydrogen.\textsuperscript{21} Years later, Cremer and Somorjai suggested that π-bonded and di-σ-bonded species co-exist on Pt (111) and the hydrogenation through π-bonded alkenes was dominant.\textsuperscript{22,23} With development of \textit{in situ} and \textit{in operando} techniques including IR, \textsuperscript{14}C labeling and NMR, evidence emerged to suggest a hydrocarbon fragment layer is formed on the Pt surface after hydrocarbon exposure.\textsuperscript{24–26} The previous model of a molecule on clean Pt surface is too simple compared to the real case. The experimental results suggested that adsorbed hydrocarbons were involved in the
hydrogenation through transferring hydrogen to the adsorbed alkenes.\textsuperscript{27} In recent years, the role of hydrocarbon fragments has been debated.\textsuperscript{28} It is generally believed now that adsorbed hydrocarbons do not directly participate in the hydrogenation reactions but act as site blockers.\textsuperscript{28–30} However, the adsorbed hydrocarbons have a significant effect on the hydrogenation reactions. It is found that the hydrogenation activity can be significantly enhanced when the alkylidyne layer is removed.\textsuperscript{31} The type of hydrocarbon molecules is also a factor that can influence the hydrogenation activity. Testing the hydrogenation of ethylene with different adsorbed hydrocarbon species, Zaera and coworkers identified the trend of hydrogenation activity as alkylidyne > benzyl > C\textsubscript{n}H > graphite.\textsuperscript{32} Under ethylene hydrogenation condition, the alkylidyne surface coverage is in the sub-saturation regime.\textsuperscript{33} The reaction is through a stepwise hydrogen addition pathway with adsorbed ethylene. Utilizing deuterium and hydrogen in ethylene hydrogenation reactions, Zaera and coworkers identified a second reaction pathway which might involve a “reverse” Eley–Rideal step by which gas-phase ethylene reacts with two deuterium atoms adsorbed on adjacent sites of the platinum surface.\textsuperscript{34}

Scheme 1.1.
For the selective hydrogenation of butadiene, the butadiene molecules bind onto the catalytic surface more strongly than alkenes. The kinetics of hydrogenation of butadiene on Pd show negative reaction order with respect to butadiene. Butadiene hydrogenation can occur through 1, 2 or 1, 4 additions to yield 1-butene or 2-butenes, respectively. Joice and coworkers proposed a butadiene hydrogenation mechanism through π-allyl, as shown in scheme 1.2. They associated the variance in the yield of 1-butene and 2-butenes with relative stability of 1-methyl-2-allyl intermediates, as cis-isomer formed from intermediate $a$ and trans-isomer formed from intermediate $b$. 

Scheme 1.2.
In order to explain the direct formation of butane and its correlation with 2-butenes, Boitiaux and coworkers proposed another butadiene hydrogenation mechanism that involves carbene formation.\textsuperscript{38} They suggested that metals that do not favor carbene formation will not allow direct hydrogenation of butadiene. Moreover, Nishimura and coworkers proposed a hydrogenation mechanism with only one double bond associated with the surface to account for the low yield to 2-butenes in hydrogenation of butadiene on Cu.\textsuperscript{36}

Pd and Pt catalysts are poisoned by the adsorbed hydrocarbons at the reaction conditions. When the Pd and Pt surfaces are exposed to hydrocarbon and hydrogen molecules, the dissociative adsorption and polymerization of the unsaturated hydrocarbons lead to a carbonaceous layer on the metal surface.\textsuperscript{39–41} Moreover, surface carbides are formed with the dissolution of carbon in the metal lattices.\textsuperscript{42} The metal carbides formed in the reaction condition result in higher selectivity and lower reactivity through poisoning. Moreover, hydrogen atoms can dissolve in Pd and Pt metal to form hydrides. Hydrogen activation on these metals is facile under ambient conditions. The hydrides, especially the beta hydrides in Pd, contribute to the full hydrogenation of alkynes to alkanes. In the low hydrogen partial pressure regime, the hydride formation is avoided but the carbon deposition becomes a problem.

Ni catalysts are also known for their high catalytic activity in hydrogenation reactions and relatively low cost. Raney nickel is widely used in the industrial hydrogenation processes. However, additives such as Cr are needed to improve its selectivity in partial hydrogenation
reactions. Recently, NiZn has been identified as a promising selective acetylene hydrogenation catalyst, which will be discussed in this chapter.

1.2.3 Pt and Pd based bimetallic catalysts for selective hydrogenation

The modification of Pd and Pt with another metal has been examined extensively for the selective hydrogenation of alkynes and dienes. Through electronic and ensemble effects, the bimetallic alloys of Pt and Pd modify the hydrogen activation and molecule adsorption to reduce the undesired reactions. The additive metals studied include Ag, Au, Bi, Cu, Ga, Sn and Zn.\(^ {43}\) Pd-Ag and Pd-Pb catalysts are applied in the commercial selective hydrogenation processes. In these systems, Pt and Pd are still the bulk phases. The ensemble sizes have been reduced in alloys but they are typically small domains greater than single atoms.

One approach to improve the selectivity of Pt and Pd catalysts is through the partial poisoning of the active sites. The Lindlar catalyst is a notable example, in which Pd is modified with Pb additives and supported on calcium carbonates. It shows high selectivity to the partial hydrogenation products and cis-isomers of alkenes. However, the high use of Pd in the catalyst significantly increases the cost and the Pb additives draw environmental concerns. Pd-Bi and Pd-Sn catalysts were investigated as non-lead alternatives to Lindlar catalysts.\(^ {44,45}\) Bi and Pd can form stable alloy phase as PdBi or PdBi\(_2\).\(^ {46}\) Bi was believed to partially poison the most active low-coordinated Pd atoms in the catalysts and disperse the Pd ensembles.\(^ {44,45,47}\) So the Pd-Bi catalysts showed high selectivity to alkenes and low alkane formation. However, Anderson and coworkers found that Pd-Bi was less selective in 2-hexyne hydrogenation compared to 1-hexyne...
Similarly, the Pd-Sn is more selective than monometallic Pd catalysts in 1,3-butadiene hydrogenation to butene. Through the addition of Sn, Pd domains were broken by Sn ensembles which reduces the beta Pd-H formation.\textsuperscript{48–50} The dilution of Pd also likely leads to lower hydrogen and hydrocarbon adsorption strength on the catalytic surfaces.\textsuperscript{51} Overall, the Pd-Sn showed lower reactivity and higher selectivity compared to Pd catalysts.

Another class of Pd and Pt alloys is with additives such as Cu, Ag, Au, Ga and Zn. In these alloys, significant electronic effects induce decreased binding energy of alkynes and alkenes and lessen the formation of beta Pd-H. Pd-Ag alloy is the active phase in industrial acetylene selective hydrogenation catalysts\textsuperscript{52,53}, so the Pd-Ag system is one of the most studied catalysts for selective hydrogenation\textsuperscript{54–58}. Unlike monometallic Pd, beta Pd-H and subsurface carbon are not formed in Pd-Ag catalysts.\textsuperscript{55} On bare Pd surface, adsorbed ethylidyne partially poisons the catalysts and leads to higher selectivity. The Ag ensembles have similar effects. They reduce the availability of hydrogen and weaken the adsorption of alkenes, leading to high selectivity.\textsuperscript{56,59}

The Pd-Ga and Pd-Zn catalysts have also been studied for their improved selectivity to partial hydrogenation products. Ga and Zn break the Pd domains and induce changes in the electronic properties of Pd, both of which favor the selective hydrogenation over the over hydrogenation.\textsuperscript{60} Moreover, intermetallic Pd-Zn nanoparticles (NPs) were formulated and used in selective hydrogenation reactions.\textsuperscript{61,62} DFT calculations suggest the Pd-Zn intermetallic surface binds ethylene through pi-bonding on the single Pd sites.\textsuperscript{62}
Cu based alloys

Cu is relatively inert compared to Pt or Pd for hydrogenation reactions. This is because of its higher hydrogen activation barrier. However, Cu is intrinsically very selective in the partial hydrogenation reactions. Pt-Cu and Pd-Cu alloys were used to improve the selectivity of Pd and Pt catalysts in selective hydrogenation reactions.

A variety of hydrogenation reactions catalyzed by Pd-Cu have been studied. By adding 30% Cu to the Pd/Al₂O₃ catalysts, Leviness et al. found Pd-Cu catalysts had higher selectivity to ethylene but lower activity in hydrogenation of acetylene. This was attributed to the ensemble effects that lower the adsorption strength of ethylene molecules and decrease the formation of Pd-H. Guzzi et al. investigated the hydrogenation of phenylacetylene and 1-butene with Pd-Cu catalysts supported on pumice. They correlated the high selectivity of Pd-Cu in partial hydrogenation of phenylacetylene to the electronic effect with the presence of CuOx, while the fully reduced Pd-Cu catalyze the isomerization of 1-butene.

The electron transfer between Cu and Pd (or Pt) largely affects the hydrogenation chemistry of Pd (or Pt). Through theoretical calculation, Fernandez-Garcia et al. identified the charge transfer between Pd and Cu in Pd₈Cu₉₂ (111) and Pd₄₀Cu₆₀(111) surfaces and found surface Pd atoms are negatively charged. This has been debated in the literature. Skoda et al. suggest there is no significant charge transfer based on the CO-FTIR results. However, other researchers have used Auger spectroscopy, CO chemisorption and XPS to show there is charge transfer on both Cu and Pd, which affects the binding strength of molecules.
Pt-Cu has been synthesized and studied for the selective hydrogenation of alkynes, hydrogenolysis of pentane, hydrogenation of acrylonitrile and hydrodechlorination of 1,2-dichloroethane.\textsuperscript{73–77} Chen and coworkers have studied the alkene and alkyne hydrogenation on Pt-Cu, Pt-Fe, Pt-Co and Pt-Ni catalysts with ambient pressure catalysis studies, surface science techniques and DFT calculations.\textsuperscript{78–82} Pt-Ni, Pt-Co and Pt-Fe show better activity compared to monometallic Pt, which is due to the weaker binding of H atoms and hydrocarbon molecules on the bimetallic surfaces compared to Pt.\textsuperscript{80} This is mainly an electronic effect as subsurface Pt-Ni-Pt catalysts showed even higher hydrogenation activity.\textsuperscript{80,81} Pt-Cu also binds H and hydrocarbon molecules more weakly than Pt, but shows lower activity than Pt-Ni, Pt-Co and Pt-Fe.\textsuperscript{78,79,82} A flow reactor study of 1,3-butadiene hydrogenation was performed with these catalysts, finding Pt-Cu has the lowest reactivity but highest selectivity to butenes.\textsuperscript{82} This suggests Pt-Cu might be a candidate for selective hydrogenation reactions. However, more understanding of Pt-Cu catalysts is needed for rational catalyst design.

The ensemble effects in bimetallic catalysts can have considerable influence on their catalytic behavior and efforts have been made to study the surface structure of Pt-Cu alloys. Lucci et al. examined the Pt-Cu surfaces with sub-monolayer Pt coverage with variable temperature STM and identified a few metastable surface structures at different Pt coverages.\textsuperscript{83} At low Pt coverage (0.01 and 0.03 monolayer), isolated Pt atoms are uniformly distributed on the Cu(111) terraces and populated with higher density in the vicinity of step edges. When more Pt (0.1ML) is deposited into the Cu(111) surface, finger-like protrusions were observed near step edges.\textsuperscript{83} And at even higher Pt loadings, 3D islands were formed on the Cu surface.\textsuperscript{84}
Au based alloys

Gold, like copper, is a group I metal, and is well known for its selective hydrogenation activity.\textsuperscript{85-89} High selectivity to ethylene at 100% acetylene conversion were reported for Au/Al\textsubscript{2}O\textsubscript{3}\textsuperscript{85} and Au/TiO\textsubscript{2}\textsuperscript{89} catalysts, while the catalytic activity was correlated with Au particle size. Lopez-Sanchez and co-workers reported that the Au/TiO\textsubscript{2} catalysts were totally selective to propene in propyne hydrogenation with progressive deactivation.\textsuperscript{13} More than 95% selectivity to butene was also demonstrated for the hydrogenation of 2-butyne catalyzed by Au/Al\textsubscript{2}O\textsubscript{3} and Au/SiO\textsubscript{2}.\textsuperscript{87} The results of DFT simulations suggested that the highly selective character of Au NPs catalysts was related to the preferential adsorption of carbon-carbon triple bonds on the edge of Au NPs, while the binding energy of carbon-carbon double bonds on Au was lower which leads to lower surface coverage.\textsuperscript{90} However, the hydrogen dissociation ability of Au is limited, due to its full d-band.\textsuperscript{91}

Bimetallic Pd-Au catalysts were employed to improve the selectivity of Pd in selective hydrogenation reactions. Louis and coworkers have prepared Pd-Au catalysts with different methods including co-deposition-precipitation, co-impregnation in excess of solution, colloids stabilized by polyvinyl alcohol (PVA) and colloids stabilized by tetrakis(hydroxypropyl)phosphonium chloride and studied their performance in hydrogenation of 1,3-butadiene to butenes.\textsuperscript{92,93} They found the selectivity of the catalysts was correlated with the distribution of Pd in the Au surface and the Pd/Au ratio. With CO-IR techniques, they demonstrated that Pd atoms can be mostly atomically dispersed in Au surface when the Pd/Au ratio is smaller than
1/20. At this condition, the selectivity of the catalysts is optimum. In other work on PdAu catalysts for selective hydrogenation reactions, Zhang et al., Szumelda et al., and Pei et al. drew similar conclusion, namely, that the best possible selectivity is achieved when the size of Pd ensembles is the smallest. Varied approaches were used to obtain different Pd distributions; Szumelda et al. and Pei et al. changed the Pd/Au ratio in catalysts supported on carbon and silica, while Zhang et al. used different pretreatment methods. Recently, Xu et al. studied the structural rearrangement of Pd-Au nanoparticles in different atmospheres with DFT calculations. They found Pd atoms could be stabilized in the surface of PdAu when hydrogen was present.

Bimetallic PdAu materials of various compositions have been studied extensively for a number of reactions, including selective oxidation, selective hydrogenation, the Ullmann reaction, vinyl acetate synthesis, formic acid decomposition, and the direct synthesis of hydrogen peroxide. Generally, it is difficult to elucidate the roles of Pd and Au in the catalytic cycles and their alloying effects due to the presence of many Pd and Au ensembles on the catalyst surfaces. In a few reports, the effect of preparing isolated Pd or Au atoms in the bimetallic PdAu catalysts has been attributed to providing exceptional activity or selectivity because of the unique chemistry of single-atom sites, which also allows for more conclusive mechanistic studies. Pioneering work from the Goodman and Toshima groups has identified Pd or Au single-atom active sites in vinyl acetate synthesis and glucose oxidation, respectively, on catalytic surfaces with well-defined single Pd or Au atom structures. The Mullins group has demonstrated that the PdAu alloy surfaces catalyze the
dehydrogenation of formic acid while the extended Pd surfaces catalyze the dehydration of formic acid.111

**Single-atom Alloys**

Single-atom alloy (SAA) is a class of alloys where the active metal is atomically dispersed and isolated in the surface of the host (inert) metal. The early work on SAA was focused on the Pd-Cu system. The Sykes group demonstrated that fully isolated Pd atoms are stable in the surface of Cu(111)119. These isolated Pd atoms can dissociate H₂ and facilitate spillover of H to the Cu surface, and the weakly adsorbed H on the Cu can participate in the selective hydrogenation reactions. As the hydrogen activation is the rate limiting step for selective hydrogenation reactions with Cu catalysts, SAA becomes an attractive approach to activate Cu and preserve its high selectivity. Boucher et al. prepare a nanoparticle analog of the model SAA catalysts by putting a small amount of Pd in the surface of Cu nanoparticles120. This Pd-Cu catalyst shows good selectivity and activity compared to its monometallic counterparts. Ma et al. studied the PdCu(111) SAA surface for selective hydrogenation of acetylene with DFT calculations. They reported that PdCu(111) SAA has moderate H₂ activation capability and low desorption barrier of ethylene, which results in its high selectivity.121

The hydrogen activation and possible spillover has been studied with Pd-Au, Ni-Cu and Co-Ag, as well. Pd, Ni and Co exist as isolated atoms in the host metals in these systems. In many of these studies, the term of “single-atom alloy” was not used.122–127 Other catalytic reactions including dehydrogenation, oxidation, hydrogenolysis, and Uhlman coupling have been investigated for the SAA catalysts with DFT and experimental approaches.128–134
1.2.4 Selective hydrogenation on non Pd or Pt catalysts

With the consideration of the cost of Pd and Pt, development of non Pd or Pt catalysts has been investigated. One of the promising non-precious metal catalysts is bimetallic Ni-Zn. Norskov and coworkers screened different bimetallic and monometallic catalysts in terms of their methyl heat of adsorption and identified NiZn as the potential non-precious metal selective acetylene hydrogenation catalyst. Furthermore, oxide catalysts have drawn attention in recent years. Pérez-Ramírez and coworkers have demonstrated ceria catalysts with...
high acetylene and propyne hydrogenation selectivity. However, relative high H$_2$/C$_3$H$_4$ ratio and high temperature are required to achieve the desired performance.$^{136,137}$

1.3 CO tolerant Pt catalysts

Pt metal is highly active for H$_2$ dissociation, which is the most important reason that Pt is widely used for fuel cell anodes and hydrogenation. However, Pt is very susceptible to carbon monoxide (CO) poisoning.$^{138}$ Due to the strong binding of CO onto Pt, even trace amounts of CO impurity, which is always present in H$_2$ -rich fuel gas produced from fuel reforming, can diminish H$_2$ activation and the overall reactivity.$^{138,139}$ It is reported that 20 ppm of CO at temperatures less than 100 °C can cause a significant activity drop of fuel cells.$^{140}$ Current research has been focused on the development of CO- tolerant Pt electrocatalysts.

The mechanism of H$_2$ activation in the presence of CO at the fuel cell anode with Pt catalysts can be expressed as follows:$^{141}$

\[ H_2 + 2Pt \rightarrow 2Pt - H \]
\[ CO + Pt \rightarrow Pt - CO \]
\[ 2CO + 2Pt - H \rightarrow 2Pt - CO + H_2 \]
\[ 2Pt - H \rightarrow 2Pt + 2H^+ + 2e^- \]

The dissociative adsorption of hydrogen is the rate limiting step in this process. Thus, the adsorption of CO on the Pt sites largely affects the anode reactions on Pt catalysts. Forming an alloy of Pt is the most-studied approach in improving its CO tolerance.$^{142-147}$ Pt-Ru and Pt-Sn anode catalysts have shown enhanced CO tolerance at fuel cell operating conditions. The effect
of Ru on Pt is twofold: it facilitates the CO electro-oxidation and weakens the Pt-CO bonds. Watanabe et al.\textsuperscript{146} and Venkataraman et al.\textsuperscript{142} demonstrated by \textit{in situ} experiments that the Ru sites lowered the CO oxidation barrier and improved the efficiency of CO conversion. It was also found that the Pt-CO binding strength is decreased with the addition of Ru.\textsuperscript{143,145,147} Moreover, Pt-Au, Pd-Au and Ni-Cu weakly adsorb CO due to electronic effects by the alloy formation, which can be potentially used in fuel cells.\textsuperscript{122,144,148,149} Ensemble effects have been shown to increase CO binding strength at extended catalytic metal sites, therefore, from a fundamental perspective, metal alloys which alter the adsorption geometry of CO should exhibit weaker binding and enhanced CO tolerance.

The strong CO adsorption also hinders the fast conversion of CO to CO\textsubscript{2} at low temperatures (< 200 °C), and becomes a technical challenge for efficient emission control and the water-gas shift (WGS) reaction for hydrogen upgrading. Recently, atomically dispersed Pt\textsubscript{1} (or Pd\textsubscript{1}) -O\textsubscript{x} -stabilized on a variety of supports, such as silica\textsuperscript{150}, titania\textsuperscript{151}, KLTL-Zeolite\textsuperscript{151,152}, MCM-41\textsuperscript{151}, alumina\textsuperscript{153} and iron oxide\textsuperscript{154}, were found to be highly active in WGS and CO oxidation reactions. The single platinum atom sites have a cationic nature that results in weak CO adsorption\textsuperscript{150}, thus becoming the exclusive sites for the low temperature CO conversion. Although single-atom catalytic sites in Pt\textsubscript{1} (or Pd\textsubscript{1})-O\textsubscript{x}-oxides possess the desired property of weaker CO adsorption, they may not be stable enough under hydrogenation conditions. Single site catalysts that can activate hydrogen molecules and are electronically conductive are desired for CO-tolerant fuel cell catalysts.
1.4 Selective dehydrogenation of light alkanes

1.4.1 Alkane to alkene processes

Light alkenes including ethylene, propylene and butene are important feedstocks for the chemical industry to produce a wide variety of chemicals. The major processes to produce light alkenes are steam cracking and fluid catalytic cracking, which make lighter alkenes from naphtha and other feedstocks. However, the high demand of propylene in polymer industry has exceeded the production of propylene from these processes. This has created so-called “propylene gap”.

The recent “shale gas boom” in North America provides low price and abundant light alkanes, which makes direct dehydrogenation of light alkanes to alkenes an economically reasonable process. There are several commercial plants in the world operating with dehydrogenation techniques to produce light alkenes. The high selectivity and lower energy demand makes the direct dehydrogenation more and more favorable for the production of light alkenes, especially with low shale gas price. One of the major technologies is the Olefex (UOP) process using Pt based catalysts. However, coke deposition is a big challenge for the commercial dehydrogenation catalysts and special designs of the reactor and regeneration process are needed to reduce coke deposition and to remove the coke. Indeed, in the industrial dehydrogenation processes, a significant portion of the catalysts are in the regeneration stage and the dehydrogenation process is often dominated by the need to control the coke formation.155,156

The industrial light alkane dehydrogenation processes typically run between 450 and 650 °C. At lower temperatures, the selective dehydrogenation reactions are largely limited by
thermodynamics, making the equilibrium alkene concentration low. Figure 1.2 shows the equilibrium conversion of alkanes at different temperatures. Effective alkene production at lower temperature is possible with lower alkane partial pressure.

![Equilibrium conversion of C2–C4 paraffins to olefins as a function of temperature at 1 bar (left) and pressure dependence of the dehydrogenation of propane as a function of temperature (right).](image)

Figure 1.2. Equilibrium conversion of C2–C4 paraffins to olefins as a function of temperature at 1 bar (left) and pressure dependence of the dehydrogenation of propane as a function of temperature (right). Adapted with permission. Copyright (2014) American Chemical Society.

1.4.2 Catalytic selective dehydrogenation reactions

Pt group metals are known for their reactivity in light alkane dehydrogenation and have been extensively studied. As shown in Scheme 1.3, a Langmuir-Hinshelwood mechanism was proposed for selective dehydrogenation of propane on Pt catalysts. The C-H bond scission happens consecutively followed by the desorption of butene and hydrogen molecules. The rate limiting step is the dissociative adsorption of propane molecule. 157,158
Thus far, Pt is the most often used metal catalyst for light alkane dehydrogenation, which is due to its high activity in C-H activation and relatively low activity in C-C cleavage. It is widely believed that the selective dehydrogenation of light alkanes is structurally insensitive. Thus, higher dehydrogenation activity can be reached with smaller particle size. However, the side reactions in this process are structurally sensitive. For most of the side reactions, especially coke formation, undercoordinated sites and extended Pt surface are the active sites.

Understanding the deactivation of the catalysts is crucial for the design of alkane dehydrogenation catalysts. Carbon deposition is the most important deactivation pathway. To tackle this challenge, both the Pt metal and the catalyst support must be optimized. Another crucial factor in deactivation is the sintering of metal nanoparticles. In the current industrial processes, catalysts are regenerated with oxidation and metal redispersion steps. Coke is burned in oxygen and Pt particles can be fragmented and redispersed with a small amount of oxygen and chlorine at a temperature around 500 °C.
1.4.3 Pt alloy catalysts for selective dehydrogenation

One of the most commonly applied approaches to reduce the carbon deposition in dehydrogenation reaction is to add a second metal to the Pt catalysts. These modifiers can improve the selectivity and control the coking through ensemble and electronic effects.

Pt-Sn, Pt-Zn, Pt-Ga and Pt-In alloys have been studied as catalysts for selective alkane dehydrogenation. Sn is the most studied and industrially widely used secondary metal for alkane dehydrogenation reactions. The addition of Sn reduces the side reactions including hydrogenolysis and isomerization, and decreases the acidity of the support. Hydrogenolysis and isomerization are the precursors of coke formation, while the surface acid sites can catalyze polymerization leading to carbon deposition. Sn is also capable of impeding the sintering of Pt at the reaction conditions.\textsuperscript{166,167} Moreover, the segregation of Sn in the regeneration process is believed to reduce the sintering of Pt.\textsuperscript{166} In the alkane dehydrogenation condition, where H\textsubscript{2} is present, Sn is in metallic state and alloyed with Pt. Both geometric and electronic effects contribute to the catalytic performance of Pt-Sn catalysts.

The geometric effects of alloying Pt with Sn are essentially the decoration of the low coordination Pt sites and the reduction of the size of Pt ensembles. FTIR as well as TPR and STEM-EDS studies have been conducted by Vironvskaia et al.\textsuperscript{168} and Nagaraja et al.\textsuperscript{167}, respectively, to study the structural evolution of Pt-Sn catalysts in alkane dehydrogenation reaction conditions. They found Sn alloyed with Pt with diffusion of Sn species taking place under reaction conditions. The coke formation on Pt-Sn catalysts was considerably reduced due to the blocking of the low coordination sites of Pt nanoparticles, which are responsible for cracking and hydrogenolysis, by Sn species. Moreover, Sn breaks the Pt ensembles. The
ensemble effect on alkane dehydrogenation is minimal as this reaction is structurally insensitive. However, the ensemble effect on the side reactions including hydrogenolysis and cracking is significant.\textsuperscript{169–171} Furthermore, Lieske et al. suggested Sn sites facilitated the diffusion of coke precursors to the oxide support, resulting in reduction of the coke formation.\textsuperscript{170,172}

Sn also alters the electronic properties of Pt upon alloying. Electrons can be transferred to Pt 5d band leading to the weakening of the adsorption of alkene molecules.\textsuperscript{158,173} Through FTIR and adsorption microcalorimetry studies, Dumesic and coworkers found the adsorption of ethylene and isobutene on Pt-Sn is weaker compared to Pt.\textsuperscript{174–176} Moreover, the formation of alkylidyne is reduced with the formation of Pt-Sn alloys. Alkylidyne is believed to be the precursor for undesired coke and hydrogenolysis. DFT studies indicated the dehydrogenation barrier of alkane on Pt-Sn is higher than on Pt, while the alkene desorption barrier on Pt-Sn is lower. Thus, the addition of Sn to Pt decreases the activity of Pt but increases the selectivity.\textsuperscript{157,158,177}

Pt-Ga, Pt-In and Pt-Ge have been investigated for alkane dehydrogenation as well. Knowing that acidity is one of the major causes of coke formation\textsuperscript{178}, the addition of Ga and In can reduce the Bronsted acidity of the catalysts, which in turn reduces the coke formation\textsuperscript{179,180}. Ge has similar effect as Sn, but its electron donation effect is smaller than Sn.\textsuperscript{181} Moreover, Li, Na and K can poison the acid sites on the metal oxide support, decreasing the coke formation.\textsuperscript{182,183} Zn and Mg were doped into the support to induce the phase change, leading to lower acidity support materials.\textsuperscript{184,185}
**Single-atom catalysts for selective dehydrogenation reactions**

Recently, Luo and co-workers proposed a single Pd atom doped in a Cu$_{55}$ cluster and conducted DFT calculations for propane dehydrogenation to propylene.$^{128}$ They found single Pd atom at the edge site of Cu$_{55}$ cluster can lower the energy barrier for the first C-H cleavage, and single Pd atom favors the C-H bond scission over the C-C one. Wasserscheid and coworkers reported a supported liquid phase Pd-Ga catalyst that is highly stable in the butane dehydrogenation reaction at 445 °C. This is due to the Ga diluting the Pd domains to make Pd ensembles smaller or even down to single atoms and the liquid nature of the active phases.$^{186}$

Non-Pt catalysts have been studied for alkane dehydrogenation as well. Single site Zn(II) stabilized with three -O-Si on the amorphous silica surface is stable at 550 °C and catalyzes the heterolytic cleavage of C-H bonds, but is unfavorable for C-C bond cleavage.$^{187}$ Gong and coworkers showed the effect of TiO$_2$ addition on the stability of Pt/Al$_2$O$_3$ catalysts and demonstrated a 10% optimal TiO$_2$ loading.$^{188}$

1.5 Thesis objectives:

In previous work from our lab, single-atom catalysts of Au and Pt on metal oxides and carbon supports have been designed and synthesized. The Au$_1$-O$_x$-OH$_y$ and Pt$_1$-O$_x$-OH$_y$ have been shown to be the active sites for various reactions. However, the single atoms in metal hosts were not investigated. In 2012, Sykes, Flytzani-Stephanopoulos and coworkers demonstrated isolated Pd atoms can be stabilized in the surface of Cu(111) in ultra-high vacuum condition to form “Single-atom Alloy (SAA)”. Sykes et al. postulated that SAAs are
potentially good catalysts for selective hydrogenation reactions. Inspired by the surface science studies on SAAs, Boucher et al. prepared bimetallic Pd$_{0.18}$Cu$_{15}$ nanoparticles as an analog of PdCu(111) SAA and showed Pd$_{0.18}$Cu$_{15}$ NPs were highly selective in hydrogenation of phenylacetylene to styrene. Lacking direct evidence to confirm its structural properties at the atomic level, this catalyst still cannot be categorized as SAA.

Following the inspirational work from the Flytzani-Stephanopoulos and Sykes groups, this thesis aimed to investigate the feasibility of designing catalytically active and stable SAA NPs and apply the SAA strategy in the rational design of practical catalysts. Through combined studies by catalytic and surface science techniques, the latter by the Sykes group at Tufts, as well as collaborative DFT calculations, this thesis also aimed to extend the understanding of structure-function relationship in SAA catalysis. A few industrially important probe reactions/processes were studied: selective hydrogenation and selective dehydrogenation reactions, and the CO tolerance of Pt catalysts. Moreover, the fundamentally important key elementary reactions involved in these reactions/processes, including hydrogen activation, C-H activation and adsorption/desorption of CO, were investigated.

1.5.1 Specific aim 1

The first part of the specific aim 1 was to investigate approaches to form PtCu SAA NP catalysts and to confirm the isolated atomic dispersion of Pt in the host metal. To date, most of the bimetallic catalysts reported in the literature are prepared with co-reduction or co-precipitation methods containing comparable compositions of the two elements. These
methods cannot avoid the formation of monometallic NPs. And the high Pt or Pd loading can result in the formation of Pt or Pd nanoclusters in the catalysts. Recent studies by Boucher et al. showed the effectiveness of galvanic replacement in the preparation of PdCu bimetallic catalysts. In other work, sequential reduction methods have been applied in the formation of core-shell structured bimetallic catalysts with very well-defined structures. Herein, both galvanic replacement and sequential reduction methods were used in the preparation of SAA NPs. Moreover, it is challenging to characterize the single-atom sites in the surface of another metal. In this thesis, in situ IR, in situ X-ray absorption spectroscopy and STEM characterization were employed to investigate the structural and chemical properties of SAA catalysts combined with other characterization techniques.

The close collaboration with surface science groups and DFT groups have also contributed to the rational catalyst design in this thesis work. The feasibility of formation of SAAs and their surface chemistry were studied by the collaborators in UHV conditions experimentally and with DFT calculations. The results are compared among all investigators to extend the understanding of the SAA catalysts.

The second part of the specific aim 1 was to design highly selective catalysts for hydrogenation of butadiene to butene and to extend the understanding of PtCu SAA catalysts in selective hydrogenation reactions. Pt alloys with the smallest possible Pt ensemble, single atoms, has not been studied in ambient pressure selective hydrogenation reactions. The SAA configuration is intrinsically selective in partial hydrogenation reactions as the side reactions that require more than one Pt atoms are avoided. The electronic effects in PtCu alloys also contribute to high selectivity. In this thesis, the selective hydrogenation of 1,3-butadiene in the
presence of excess propylene is studied with supported PtCu SAA catalysts. This is an industrially important reaction and the performance of SAA catalysts is of great interest. This thesis aimed at extending the understanding of selective hydrogenation with SAA catalysts and answer one important fundamental question: are isolated Pt single atoms enough to activate Cu for selective hydrogenation reactions?

1.5.2 Specific aim 2

The specific aim 2 of this thesis was to gain an improved understanding of the nature of active sites of PtCu SAA with CO as probe molecule and to investigate an approach to improve the CO tolerance of Pt catalysts. CO is an important probe molecule to characterize the catalytic surfaces. Its interaction with the surface-active sites can effectively reflect the surface chemistry of the catalysts. For bimetallic alloys, the electronic and ensemble effects can influence the binding strength of CO and the vibrational frequency of adsorbed CO molecules. Thus, studying the interaction of CO with SAA catalysts is of fundamental interest. Moreover, CO poisoning is one of the major practical challenges for Pt catalysts, especially the Pt fuel cell catalysts. Based on the literature of other Pt alloys, the small Pt ensembles and electronic effects in Pt-Cu are beneficial for the CO tolerance of Pt catalysts. Thus, we investigated PtCu SAA as a promising CO tolerant catalyst. Its high performance in hydrogen activation would also contribute to its further application in various reactions.
1.5.3 Specific aim 3

C-H activation is the key elementary step in many important chemical reactions, including the conversion of light alkanes and alcohols. The light alkanes conversion to chemicals is more and more attractive with the recent “shale gas boom”. Pt is one of the most important catalytic metals for alkane dehydrogenation reactions, however, it suffers from instability due to coking. The specific aim 3 of this thesis was to investigate an approach to overcome the challenges of coking in alkane dehydrogenation reactions. Specifically, the use of PtCu SAA catalysts compared to monometallic Pt catalysts was studied. And to investigate the C-H activation reactions with SAA catalysts. C-H activation barrier on Cu is relatively high and the coke formation is unlikely. This thesis determines what is the smallest possible Pt ensemble in Cu that can catalyze the C-H activation reaction. The selective dehydrogenation catalyst is designed with this smallest possible Pt ensemble to avoid the coke formation.

1.5.4 Specific aim 4

The last specific aim of this thesis was to explore the potential of SAAs for the design of selective hydrogenation catalysts with PdAu for hydrogenation reactions in the liquid phase, and to extend the understanding of selective hydrogenation reactions with PdAu SAA catalysts. To date, Cu based SAA catalysts, especially PtCu, have been extensively studied for various selective hydrogenation and dehydrogenation reactions, which are reported in this thesis and in others’ work. Beyond that, this thesis aimed to explore the feasibility of applying the SAA strategy to Au based catalysts, especially PdAu, an important bimetallic catalyst with surface chemistry different from Cu in many ways. The performance of SAA catalysts in liquid
phase selective hydrogenation is also of great interest. Moreover, a sequential reduction preparation approach, adapted from the literature was developed as an effective method to form SAA NPS with Au.\textsuperscript{189} By investigating PdAu SAA systems, this thesis shows that the SAA approach is not limited to a specific class of bimetallic catalysts; indeed it can be a general design approach for many bimetallic catalysts for different reactions.

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Chapter 2. Experimental methods and procedures

2.1 Catalyst preparation

2.1.1 Materials and Chemicals

For the preparation of nanoparticles, metal precursors chloroplatinic acid hydrate ($\text{H}_2\text{PtCl}_6 \cdot x \text{H}_2\text{O}$, ~38% Pt basis, Sigma-Aldrich), and copper nitrate trihydrate ($\text{Cu(NO}_3\text{)}_2 \cdot 3 \text{H}_2\text{O}$, Johnson Matthey) were used. Aluminum oxide (Y-$\text{Al}_2\text{O}_3$, ultra-pure grade 99.99%, surface area 70-100 m$^2$/g, Inframat Advanced Materials) was used as catalyst support. Poly(vinylpyrrolidone) (PVP, MW= 58,000, Alfa Aesar) was used as stabilizing agent for colloidal nanoparticles. L-ascorbic acid (Sigma-Aldrich) was used as anti-oxidizing agent. The copper precursor was reduced with sodium borohydride (NaBH$_4$, Sigma-Aldrich) to form the nanoparticles (NPs).

2.1.2 Pt-Cu catalysts

Pt-Cu bimetallic samples were synthesized by galvanic replacement (GR) reaction as described below. Briefly, an aqueous solution of Cu(NO$_3$)$_2$ and PVP (200:1 molar ratio of Cu to PVP, 1:1 wt/wt) was degassed in vacuum and in nitrogen gas sequentially. A 0.1 M solution of ascorbic acid was added to the Cu/PVP solution, followed by the addition of NaBH$_4$ solution (0.1 M) dropwise. The solution turned to an opaque brown suspension after the addition of NaBH$_4$. Aluminum oxide (2 g) was suspended in 100 ml of DI water under magnetic stirring and added to the Cu NPs colloidal solution dropwise. The solution was kept under flowing nitrogen and constant stirring for 30 min before it was filtered and washed with DI water 3 times. The
resulting materials were dried in vacuum for 24 h, calcined in air to 350 °C at 2 °C/min and held for 4 h.

GR reaction took place in aqueous solution under nitrogen protection with constant stirring and refluxing at 100 °C. The desired amount of Pt precursor (H₂PtCl₆·x H₂O, Sigma - Aldrich) was added to the suspension of Cu NPs (pre-reduced in 100% H₂ at 300 °C) containing HCl (2mM). The Pt precursor was reduced by Cu host via the displacement reaction shown in Scheme 1. After 20 min, the resulting material was filtered, washed and dried in vacuum.

\[
\text{PtCl}_6^{2-} + 4 e^- \rightarrow \text{Pt} + 6 \text{Cl}^- \quad (\Delta E^o = 0.74 \text{ V})
\]
\[
\text{Cu} \rightarrow \text{Cu}^{2+} + 2 e^- \quad (\Delta E^o = -0.34 \text{ V})
\]

**Scheme 1.** Galvanic replacement reaction.

2.1.3 Pd-Au catalysts

PdAu-SAA NPs were prepared using sequential reduction methods as reported in Tedsree et al. with some modifications. First, Au NPs were prepared as follows: 0.3 g HAuCl₄·3H₂O and 1.2 g poly(vinylpyrrolidinone) (PVP, MW= 58,000) were mixed in 50 mL ethylene glycol (EG) in a round bottom flask with vigorous stirring. The solution was purged with N₂ for one hour before 0.3 g NaHCO₃ was added. The solution was then heated to 90 °C at 5 °C/min with N₂ flowing through the round bottom flask. The solution was held at 90 °C for half an hour before cooling to ambient temperature. The Au NPs were thus obtained. To prepare PdAu-SAA NPs, the desired amount of Pd(NO₃)₂·x H₂O was dissolved in 1 mL EG and added to
the round bottom flask containing the EG-dispersed Au NPs. Under N\textsubscript{2} purge and vigorous stirring, the solution was heated to 90 °C at 5 °C/min and held for 8 h. After the solution was cooled to ambient temperature, 300 mL of acetone was added in order to precipitate the NPs. The acetone-EG solution was centrifuged at 3000 rpm for 10 minutes to separate the NPs. The retrieved NPs were then washed with ethanol and hexane thrice and dried in vacuum overnight. To obtain silica supported Au and PdAu NPs, fumed silica (heat treated in air at 650 °C for 3 h) was introduced into the NP gold and PdAu-SAA solutions in water and stirred overnight. The slurry carrying the supported NPs was centrifuged and dried in vacuum for 24 h. The supported NPs were calcined in air at 300 °C for 1 hour followed by reduction in H\textsubscript{2} at 250 °C for 1 hour. The unsupported Pd-NPs were prepared by the same method with Pd(NO\textsubscript{3})\textsubscript{2} precursor.

2.2 Catalyst characterization and activity measurements

2.2.1 Catalyst characterization

2.2.1.1 Surface Area

The BET surface areas of the samples were measured by single-point N\textsubscript{2} adsorption/desorption cycles in a Micromeritics AutoChem II 2920 apparatus. Each sample was degassed in He at 250-350°C for 60 min before the adsorption. A 30 % N\textsubscript{2}-He gas mixture was used in the BET measurement.
2.2.1.2 High-resolution transmission electron microscopy (HR-TEM)

HR-TEM was conducted on a JEOL 2010 electron microscope with 200 kV and 107 μA beam emission. The specimens were obtained by drying one drop of colloidal solution of NPs in ethanol onto a carbon film with nickel or copper covered microgrid.

2.2.1.3 Materials compositions

The elemental composition of the catalysts was measured by inductively coupled plasma atomic emission spectrometry (ICP-AES). The samples were dissolved in HCl-H₂O₂ or HCl-HNO₃ mixture overnight. The solution was further diluted before the ICP tests for all elemental concentrations to be within the 1-100 ppm range.

The Pt and Cu loading on each sample was measured by ICP-AES on a Leeman Labs PS1000 instrument. The solid samples were digested in concentrated HCl and H₂O₂ prior to the testing. No Pt or Cu was detected by XPS on the solid alumina support after digestion. The collected wash solution from GR and Cu NP preparation was used without additional treatments.

2.2.1.4 Ultraviolet and visible light (UV-Vis) absorption spectroscopy

UV-Vis absorption spectra were collected in the range of 200 - 800 nm using an Evolution™ 300 UV-Vis Spectrophotometer (Thermo Scientific). The diluted solution of the samples was placed in a quartz cuvette. Standard solution was prepared with HPtCl₆ and Cu(NO₃)₂.
2.2.1.5 X-ray photoelectron spectroscopy (XPS)

XPS data were obtained by using a Thermo Scientific K-Alpha system equipped with an Al source and a double focusing hemispherical analyzer with a 128-channel detector at a pass energy of 50-100 eV. To prepare the XPS samples, the powders were deposited on the copper tapes or silver plates. The copper tapes and silver plates were fixed on a glass slide with the carbon tapes before being transferred into the XPS chamber.

2.2.1.6 X-ray powder diffraction (XRD)

XRD analysis was performed on a PANalytical X'Pert Pro instrument. Cu Kα radiation was used with a power setting of 40 mA, 45 kV. Data was collected for 2θ between 10° and 80°. The catalysts were reduced in 100% H₂ at 350 °C (Pt-Cu samples) or 300 °C (Cu monometallic samples) prior to the XRD measurements.

2.2.1.7 X-ray absorption spectroscopy (XAS)

XAS was performed at beamline X18B at Brookhaven National Laboratory and beamline 12-BM at Argonne National Laboratory. The Pt-Cu and Pt NPs were reduced in 100% H₂ at 350 °C (Pt-Cu bimetallic samples) or 400 °C (Pt monometallic samples) in situ prior to the tests. PdAu samples were treated in H₂ at ambient temperature to 150 °C. The X-ray absorption near edge structure (XANES) spectra were collected in H₂ or He atmospheres. Extended X-ray
absorption fine structure (EXAFS) was measured in H₂ or He. The spectra were recorded in the fluorescence mode at room temperature.

XAS data were processed and analyzed using Athena and Artemis.² The XANES spectra were background corrected and normalized. And the EXAFS data were fitted in r-space with the models based on metallic Pt, PtCu₃, and PtO.³

2.2.1.8 Scanning transmission electron microscopy (STEM)

Aberration-corrected high-angle annular dark-field (HAADF) STEM images of Pt-Cu NPs were obtained at a nominal resolution of 0.07 nm using a JEOL 2200FS-AC STEM/TEM equipped with a hexapole corrector (CEOS GmbH, Heidelberg, Germany) at Oak Ridge National Laboratory. Energy-dispersive X-ray spectroscopy (EDS) imaging was conducted with a Bruker-AXS 30 mm² silicon-drift detector system and Pt, Cu, Al and O elemental maps were collected. The instrument was operated at 200 kV for all imaging and EDS work.

HAADF-STEM images of Pd/Au NPs were obtained at a nominal resolution of 0.078 nm using a JEOL ARM200 STEM/TEM at center for nanoscale systems at Harvard University. EDS imaging was conducted, and elemental maps were collected. The instrument was operated at 200 kV for all imaging and EDS work.

2.2.1.9 Fourier-transformed infrared spectroscopy (FTIR)

FTIR spectra were collected using a Nicolet Nexus 470 spectrometer equipped with a MCT-B detector in the single beam absorbance mode at a resolution of 2 cm⁻¹. Prior to the IR
experiments, the samples were reduced in pure H$_2$ at 300 (Cu-NP), 350 (Pt$_{0.008}$Cu-SAA and Pt$_{0.39}$Cu-bimetallic) or 400 (Pt-NP) °C in a quartz microreactor and sealed in the glass vials. The powder sample of 0.02 g was made into a thin pellet (~1cm in diameter). The sample was then loaded into a flow cell sealed with NaCl windows and allows the CO flow through the catalyst pellet. In a typical CO adsorption IR experiment, the sample was first reduced in 50 mL/min 5 % H$_2$-He for 1 hr at 200 °C followed with the He purge to cool down to 25 °C. Baseline was collected at 25 °C under He purge after the catalyst has been swept for at least 30 min. The sample was then exposed to 1 % CO-He flow at the same temperature for 45 min and was purged again with He for at least another 30 min until the gas phase CO signal was fully diminished (rather quickly within 5 min of purge) and the catalyst surface adsorption spectra has minimal difference between the IR scans. The IR spectra shown in the main text were collected afterwards at 25, 40, 60, 80, and 100 °C. A ramping rate of 5 °C/min was used. The corresponding spectra at each temperature were collected after the sample was stabilized at the target temperature for 10 min.

2.2.2 Catalytic activity measurements

2.2.2.1 Flow reactor studies

The selective hydrogenation activity of the catalysts was tested in a quartz-bed flow reactor for 1,3-butadiene hydrogenation with 400 mg of catalyst diluted by 1.5 g of quartz particles. The as synthesized Pt-Cu catalysts were reduced in H$_2$ at 350 °C for 4 h prior to the reaction, the Cl-residues were removed in H$_2$ as well. A gas mixture of 1.25 % 1,3-butadiene, 20 % H$_2$ and balance He (flow rate=20 mL/min, GHSV=1,200 h$^{-1}$) was introduced at 120 °C,
followed by descending temperature testing. Gas chromatograph (GC) injections were done at each temperature after the temperature was stabilized for at least 10 min. The exit gas stream was analyzed by a HP-6890 GC equipped with a 30 ft. column (1/8 in., filled with Sebaconitrile 20 % Chromosorb Paw 80/100).

The hydrogenation activity of the catalysts was tested in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) with 100 mg of catalyst diluted by 0.5 g of quartz particles. The samples were pre-reduced in 100% H₂ at 25 mL/min (300 °C for Cu₁₅/Al₂O₃, 350 °C for Pt-Cu samples and 400 °C for Pt₀.₁/Al₂O₃) with a heating rate of 2 °C/min, and then cooled to 30 °C before the reaction gas mixture (2 % 1,3-butadiene, 20 % H₂ and balance He. GHSV=12,000 h⁻¹) was introduced. The tests were started at 30 °C and the temperature was raised to higher values (60, 90, 120, 150 and 170 °C) at 5 °C/min and held at each temperature for 30 min. After reaching 170 °C, the samples were cooled back to RT, collecting data at selected intermediate temperatures to check for (de)activation. Gas chromatograph (GC) injections were done by sampling the gas at various temperatures after stabilizing the temperature for 10 min. Moreover, long-time stability tests were conducted isothermally at 160 °C for 46 h following the activity tests (2 % 1,3-butadiene, 20 % H₂ and balance He. GHSV=12,000 h⁻¹). The exit gas stream was analyzed by a HP6890 GC equipped with a flame ionization detector (FID). A 30 ft. column (1/8 in. dia.) filled with Sebaconitrile 20 % Chromosorb Paw 80/100 supplied by Sigma-Aldrich was used to separate the gas species.

The activity tests with added propylene were conducted in a similar fashion with a gas mixture of 2 % 1,3-butadiene, 20 % propylene, 16 % H₂ and balance He. The total flow rate was 50 mL/min, and the catalyst load was 100mg (GHSV=12,000 h⁻¹). The steady-state stability tests
were conducted isothermally at 160 °C and 145 °C, for 12 h at each temperature. The exit gas stream composition was determined by GC as described above.

Cyclic hydrogenation tests were performed to probe the stability of the catalysts. The first cycle was performed as described for the activity tests. Then the sample was treated in 100 % H₂ at 350 °C for 1 h. The reaction gas mixture (2 % 1,3-butadiene, 20 % H₂ and balance He. GHSV=12,000 h⁻¹) was introduced at 170 °C followed by cooling down to other temperatures. GC injections were done at specified temperatures after the temperature was held stable for at least 10 min.

H₂-D₂ exchange was conducted in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) with 80-90 mg of catalyst diluted by 0.5 g of quartz particles. The samples were pre-reduced in 100% H₂ (at 350 °C for Pt₀.₀₀₈Cu-SAA and 400 °C for Pt-NP) and then cooled to desired temperature before the reaction gas mixture (33% H₂, 33% D₂, 3.3% CO and balance Ar. Flow rate=50mL/min) was introduced. For the isothermal H₂-D₂ exchange, CO was introduced to the gas phase and turned off at the certain time points while H₂ and D₂ were flowing. To do temperature programmed H₂-D₂ exchange, a gas mixture of 33% H₂, 33% D₂, 3.3% (or 667ppm) CO and balance Ar (flow rate=50 mL/min) was introduced at 120 °C and temperature was increased to 200 °C at 5 °C/min. The gas effluent from the reactor was analyzed by mass spectrometry (SRS RGA 200). Acetylene hydrogenation was performed in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) at 70 and 80 °C isothermally (20% H₂, 2.2% C₂H₂, 200 ppm CO, 10% Argon, balance He, flow rate=24-48 mL/min). 50-500 mg catalysts were used. The exit gas stream was analyzed by mass spectrometry.
Temperature-programmed oxidation (TPO) was conducted in a Micromeritics AutoChem II 2920 instrument equipped with a mass spectrometer. The spent catalyst (33 mg) used in the long-time stability testing with or without addition of propylene was loaded to the microreactor for the TPO test. The sample was degassed with He, flowing at a rate of 50 mL/min, with a heating rate of 10 °C/min up to 100 °C, and cooled to 25 °C in He. Next, 20% O₂ in He at the same flowrate was introduced. The sample was stabilized at 25 °C for 5 min followed by raising the temperature to 800 °C at 3 °C/min. The CO₂ in the gas stream was analyzed online by a mass spectrometer (Pfeiffer, TCP270). Temperature-programmed desorption in He (He-TPD) was performed with a similar setup. The spent catalyst (33 mg) from the long-time stability testing with or without addition of propylene was loaded to the microreactor for the He-TPD test. After degassing in pure He flow (50 mL/min) for 20 min at ambient temperature, the sample was heated in the He gas (50 mL/min) to 800 °C at 3 °C/min. The exit gas stream was analyzed online by a mass spectrometer in histogram-spectrum mode scanning mass-to-charge ratio (m/z) between 1 and 120.

2.2.2.2 Batch reactor studies

A stainless-steel Parr reactor was used for the hydrogenation of 1-hexyne. The desired amount of sample was added to 20-80 mL ethanol. The mixture was stirred for half an hour and sonicated for 15 minutes before being loaded into the reactor. Pure hydrogen was bubbled through the solution while stirring for 1 h. 1-hexyne was injected into the solution to reach 1% vol/vol concentration in ethanol. The reactor was then pressurized with pure hydrogen to the
desired pressure (5 bar) under stirring at 600 rpm. Liquid-phase specimens were taken from the reactor at different time points.

Leaching test: the catalyst sample was put in the typical hydrogenation reaction condition (5 bar H₂, 600 rpm, 25 °C) for 6 hours before centrifugation to separate the solid sample from the liquid solution. The liquid solution was purged with pure H₂ for one hour before adding fresh 1-hexyne with 1/100 (vol/vol) ratio. The solution was transferred into the reactor and held under reaction condition for 60 min. Liquid-phase specimens were taken from the reactor at different time points.

2.3 References


Chapter 3. PtCu single-atom alloy nanoparticle catalysts for selective hydrogenation of 1,3-butanediene

3.1 Introduction

Pt is one of the most widely used transition metal catalysts due to its superior catalytic performance for both oxidation and hydrogenation reactions. It is used extensively in electrochemical and heterogeneous catalysts with applications in fuel cells, automotive catalytic converters, and industrial hydrocarbon cracking processes.\(^1,2\)

This chapter reports the very low concentrations of individual, isolated Pt atoms in a Cu surface catalyze the industrially important reaction of butadiene hydrogenation with high selectivity to butene. Butadiene poisons polymerization catalysts even at low concentrations (\(<\ 10\ \text{ppm}\)) in industrial alkene streams.\(^3\) Of particular interest is the butadiene impurity in propene feedstocks used to produce 42.3 million tons of polypropylene annually.\(^4\) The selective hydrogenation of butadiene to butene serves to increase the purity of alkene feedstocks without reducing their overall concentration. Therefore, catalysts that selectively hydrogenate butadiene to butene and prevent the hydrogenation of butene to butane are of great interest. It has been proposed that the observed product distribution is controlled by the adsorption energy of butadiene to the catalytic surface and weaker bonding is known to direct the product distribution in favor of butene formation.\(^5,6\) We designed and prepared highly diluted Pt-Cu nanoparticles using the single-atom alloy (SAA) principle.\(^7,8\) Nominal amounts of Pt (2 at. %) in
Cu nanoparticle catalysts were found to exhibit high activity and selectivity for butadiene hydrogenation to butene under relatively mild conditions.

3.2 Experimental methods

Pt-Cu nanoparticles (NPs) with different Pt-loadings were synthesized using the galvanic replacement (GR) method on the pre-reduced Cu NPs as described by Boucher et al. The Cu NPs were pre-formed and supported on ϒ-Al₂O₃, followed by calcination in air at 350 °C. GR took place in aqueous solution under nitrogen protection with constant stirring and refluxing at 100 °C. Desired amounts of Pt precursor (H₂PtCl₆·xH₂O, Sigma - Aldrich) were added to a suspension of Cu NPs in an aqueous solution containing HCl (2mM). After 20 min, the resulting material was filtered, washed and dried in vacuum. The sample information is listed in Table 3.1.
**Table 3.1. Sample composition**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Composition$^a$</th>
<th>Preparation Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu$_{15}$/Al$_2$O$_3$</td>
<td>14.9 at.% Cu</td>
<td>Colloidal</td>
</tr>
<tr>
<td>Pt$_{0.1}$/Al$_2$O$_3$</td>
<td>0.12 at.% Pt</td>
<td>IWI$^b$</td>
</tr>
<tr>
<td>Pt$<em>{0.1}$Cu$</em>{14}$/Al$_2$O$_3$</td>
<td>0.11 at.% Pt, 13.7 at.% Cu</td>
<td>GR</td>
</tr>
<tr>
<td>Pt$<em>{0.2}$Cu$</em>{14}$/Al$_2$O$_3$</td>
<td>0.25 at.% Pt, 12.3 at.% Cu</td>
<td>GR</td>
</tr>
<tr>
<td>Pt$_2$Cu$_6$/Al$_2$O$_3$</td>
<td>2.2 at.% Pt, 5.6 at.% Cu</td>
<td>GR</td>
</tr>
</tbody>
</table>

[a] Determined by ICP-AES.

[b] Incipient wetness impregnation (IWI).

HAADF-STEM imaging and EDS elemental mapping were conducted as described in chapter 2. XAS measurements at the Pt L$_{III}$-edge were made at Argonne National Laboratory and Brookhaven National Laboratory in fluorescence mode at room temperature. As described in chapter 2, all samples were reduced in H$_2$ *in situ* prior to the measurements. XAS data were processed and analyzed using Athena and Artemis. The XANES spectra were background corrected and normalized. And the EXAFS data were fitted in r-space with the models based on metallic Pt, PtCu$_3$, and PtO.

The selective hydrogenation activity of the catalysts was tested in a quartz-bed flow reactor for 1,3-butadiene hydrogenation with 400 mg of catalyst diluted by 1.5 g of quartz particles. The as synthesized Pt-Cu catalysts were reduced in H$_2$ at 350 °C for 4 h prior to the
reaction, the Cl-residues were removed in H₂ as well. A gas mixture of 1.25 % 1,3-butadiene, 20 % H₂ and balance He (flow rate=20 mL/min, GHSV=1,200 h⁻¹) was introduced at 120 °C, followed by descending temperature testing. Gas chromatograph (GC) injections were done at each temperature after the temperature was stabilized for at least 10 min. The exit gas stream was analyzed by a HP6890 GC equipped with a 30 ft. column (1/8 in., filled with Sebaconitride 20 % Chromosorb Paw 80/100).

The activity tests with added propylene were conducted with a gas mixture of 2 % 1,3-butadiene, 20 % propylene, 16 % H₂ and balance He at a flow rate of 50 mL/min (100 mg catalyst, GHSV=12,000 h⁻¹). After the activity tests with ascending temperature up to 170 °C, the stability tests were performed isothermally at 160 °C and 145 °C for 12 h at each temperature.

3.3 Results and Discussion

3.3.1 Characterization of PtCu NP catalysts

Similar to Boucher et al.,¹¹ the Cu NPs were prepared without a support. The average particle size of colloidal Cu NPs is 4.24 ± 1.00 nm with a narrow particle size distribution (Figure 3.1). The Al₂O₃ powder were subsequently introduced to the Cu@PVP colloidal solution to effect Cu@PVP deposition on the support. The alumina supported Cu@PVP were calcined in air to 350 °C to burn off the PVP surfactant. A typical TEM image of the calcined Cu₃Al₂O₃ is shown in Figure 3.1B. Figure 3.2 shows the high resolution TEM (HRTEM) images of Cu₃Al₂O₃. The lattice spacing of Cu facets is consistent with that of metallic Cu (Figure 3.2B). Additionally,
we found the average crystalline size of Cu NPs increased to 13 nm upon the heat treatments, as determined from the X-ray diffractograms in Figure 3.8 by Scherrer equation. Boucher et al. reported the sintering of Cu NP during the similar heat treatment in air, and they showed the particle size of Cu NP was more than double as a result of this process.¹¹

Figure 3.1. (A) TEM images of unsupported Cu NPs, inset is particle size distribution (scale bar=50 nm). (B) TEM image of Y-Al₂O₃ supported Cu NPs (calcined).
Figure 3.2. (A) HR-TEM images of supported Cu NPS. (B) An enlarged region from (A).

Complete Pt uptake was verified by UV-Vis spectroscopy of the GR filtrates after replacement of Cu with Pt, as shown in Figure 3.3. In contrast to PtCl$_6^{2-}$ standard, the GR wash solution did not show any absorption band at 260 nm. The broader band in the range of 600 –
800 nm indicates the presence of Cu$^{2+}$ in the GR wash solution. The PtCl$_6^{2-}$ absorption peaks at 260 nm of the standard solutions are consistent with those reported in the literature. ICP-AES was used to quantify Pt in the wash solution and no Pt was detected. The observed concentration of Cu$^{2+}$ is (around 4 times) more than the Cu$^{2+}$ produced in GR process. The extra Cu$^{2+}$ is a result of some dissolution of surface CuO in acidic solution (CuO +2H$^+$ → Cu$^{2+}$ + H$_2$O). The Cu NPs were shortly exposed to the air when transferred from the reduction furnace (100% H$_2$, 300 °C) to GR solution. The surface CuO layer that would otherwise impede the GR process was removed by the acid in the solution.

**Figure 3.3.** UV-Vis spectra of PtCl$_6^{2-}$ standards compared to the filtrate from GR preparation of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$
Three different Pt-Cu NPs were prepared by the galvanic replacement (GR) method, in which a controlled amount of Pt was exchanged with Cu on pre-formed Cu NPs supported on γ-Al₂O₃. Low concentrations of Pt in solution form SAA Pt₀.₁Cu₁₄/Al₂O₃ and Pt₀.₂Cu₁₂/Al₂O₃ NPs, while higher concentrations of Pt form Pt₂Cu₆/Al₂O₃.

Elemental mapping by energy dispersive X-ray spectroscopy (EDS) indicates that the Pt-Cu NPs are all bimetallic. Pt, Cu, Al, and O elemental mapping by EDS shows that Pt is distributed over the Cu NPs and is not deposited on the γ-Al₂O₃ support (Figure 3.4). In situ EXAFS performed at room temperature at the Pt-L₃ edge reveals the coordination of Pt in Pt-Cu alloys (Figure 3.5 and Table 3.1). No Pt-Pt bonds are detected in Pt₀.₁Cu₁₄/Al₂O₃ and Pt₀.₂Cu₁₂/Al₂O₃, which provides direct evidence for individual, isolated Pt atoms in the Pt-Cu bimetallic NPs. The Pt-Cu first shell interaction distance is 2.61 Å which is between Pt-Pt (2.77 Å) and Cu-Cu bond lengths (2.56 Å), further supporting SAA formation. The finding of Pt-Pt bonds in Pt₂Cu₆/Al₂O₃ suggests that Pt islands and/or clusters are formed in the NPs with higher Pt loading.
Figure 3.4. (A) STEM image and elemental mapping (EDS) of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$. (B) Al, (C) O, (D) Cu and (E) Pt maps obtained from the same region on the sample.

The *in situ* EXAFS of Pt-Cu bimetallic samples and Pt foil are plotted in k- and Fourier transform spaces (Figure 3.5 and 3.6E). EXFAS of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ and Pt$_{0.2}$Cu$_{12}$/Al$_2$O$_3$ are significantly distinct from that of Pt foil and Pt$_2$Cu$_6$/Al$_2$O$_3$. Indeed, no Pt-Pt bonds were formed in Pt$_{0.2}$Cu$_{12}$/Al$_2$O$_3$ or Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$. Also, no Pt-O bonds were detected in the Pt-Cu NPs samples (Table 3.1).
Figure 3.5. *In situ* Pt L\(\text{III}\) EXAFS of selected samples and Pt foil plotted in k space.

**Table 3.1.** EXAFS model fitting

<table>
<thead>
<tr>
<th>Sample</th>
<th>Shell</th>
<th>CN(^{[a]})</th>
<th>R(^{[b]}) (Å)</th>
<th>(\sigma^2) (Å(^2))</th>
<th>R-factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt foil</td>
<td>Pt-Pt</td>
<td>12</td>
<td>2.77</td>
<td>0.0052</td>
<td>0.015</td>
</tr>
<tr>
<td>Pt(<em>{0.1})Cu(</em>{14})/Al(_2)O(_3)</td>
<td>Pt-Pt</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pt-Cu</td>
<td>11.2 ± 1.93</td>
<td>2.61</td>
<td>0.0066</td>
<td>0.006</td>
</tr>
<tr>
<td>Pt(<em>{0.2})Cu(</em>{12})/Al(_2)O(_3)</td>
<td>Pt-Pt</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pt-Cu</td>
<td>11.77 ± 2.31</td>
<td>2.64</td>
<td>0.0078</td>
<td>0.016</td>
</tr>
<tr>
<td>Pt(_2)Cu(_6)/Al(_2)O(_3)</td>
<td>Pt-Pt</td>
<td>4.30 ± 1.44</td>
<td>2.70</td>
<td>0.0050</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pt-Cu</td>
<td>7.95 ± 1.51</td>
<td>2.61</td>
<td>0.0087</td>
<td>0.009</td>
</tr>
</tbody>
</table>

\(^{[a]}\) CN, coordination number; \(^{[b]}\) R, distance between absorber and backscattered atoms. R-factor, closeness of the fit. \(\sigma^2\) is the disorder in the neighbor distance.
Figure 3.6 shows ac-HAADF-STEM images of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$. Aberration-corrected STEM can distinguish isolated atoms due to differences in Z-contrast.$^{16,17}$ In this work, we observed a number of bright, atom-sized features within the Cu lattice (Figure 3.6). Our EXAFS results indicate that these features are isolated Pt atoms in Cu. Additionally, the lattice spacing of Pt-Cu is comparable to the pure Cu lattice spacing which also supports dilute dispersion of Pt atoms (Figure 3.6D). Variations in the background structure of the Cu NPs make imaging of isolated atoms difficult, but in Figure 3.6D the single Pt atoms are more apparent due to the uniform background.$^{17}$ STEM and EXAFS analysis demonstrate the formation Pt-Cu SAA NPs, which provides a new catalytic system to study selective hydrogenation reactions.

The Pt-Cu SAA NPs in Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ is imaged by the ac-HAADF-STEM after reduction in H$_2$ at 350 °C. The isolated Pt atoms are brighter than the Cu areas in the dark field STEM images. Thus, we are able to identify the isolated Pt atoms from its higher brightness comparing to its surrounding area. As shown in Figure 3.7, the isolated Pt atoms are dispersed on the Cu NPs. The white circles highlight some of the isolated Pt atoms. The background from Cu NPs in the ac-HAADF-STEM images is not uniform. Assuming a spherical shape of the NPs, the center region appears brighter due to the greater contribution from the Cu as highlighted by black circles. However, within localized regions, the background remains constant and we observe the bright Pt atoms surrounded by dark Cu regions, which is the evidence for the formation of isolated Pt atoms. Besides that, we do not observe any single atoms or fine clusters of Pt atoms on the alumina.
Figure 3.6. Characterization of Pt-Cu SAA NPs. (A-D) HAADF-STEM images with (C) colored intensity map from selected region, and (E) EXAFS $k^3$ – weighted Fourier transforms. (A, D) typical regions of the sample $\text{Pt}_{0.1}\text{Cu}_{14}/\text{Al}_2\text{O}_3$, showing Cu metal particles with isolated Pt atoms. Isolated Pt atoms are highlighted by red arrows. The lattice spacing of Cu(111) is 0.21 nm. (B)
Enlarged image and (c) colorized intensity map of highlighted region showing isolated Pt atoms.

(E) EXAFS data was collected in situ at Pt-L_{III} edge at room temperature from Pt foil, Pt_{0.1}Cu_{14}/Al_{2}O_{3}, Pt_{0.2}Cu_{12}/Al_{2}O_{3} and Pt_{2}Cu_{6}/Al_{2}O_{3}.

![Typical HAADF-STEM images of Pt_{0.1}Cu_{14}/Al_{2}O_{3}.](image)

**Figure 3.7.** Typical HAADF-STEM images of Pt_{0.1}Cu_{14}/Al_{2}O_{3}. The samples were pre-reduced in H$_{2}$ at 350 °C. Isolated Pt atoms are highlighted by white circles.

XRD patterns of selected samples are shown in Figure 3.8. Fresh Pt_{0.2}Cu_{12}/Al_{2}O_{3} and Pt_{0.2}Cu_{12}/Al_{2}O_{3} were used in the hydrogenation of butadiene have similar XRD patterns as Cu_{15}/Al_{2}O_{3}. Only the metallic Cu phase was detected in Pt_{0.2}Cu_{12}/Al_{2}O_{3} and used Pt_{0.2}Cu_{12}/Al_{2}O_{3}. Any CuO skin formed by exposure of the samples to air upon transfer for analysis was too little to show by XRD. No shift in the Cu-peaks was observed between Pt_{0.2}Cu_{12}/Al_{2}O_{3} and used Pt_{0.2}Cu_{12}/Al_{2}O_{3} or Cu_{15}/Al_{2}O_{3}. This suggests the addition of small amounts of Pt did not change the lattice structure of Cu, which is consistent with the lattice spacing determined by STEM
No CuO or Pt phases were observed by XRD. However, Pt-Cu alloy phases were present in Pt$_2$Cu$_6$/Al$_2$O$_3$ and the Cu phase diminishes, which indicates the formation of bulk Pt-Cu alloys. Furthermore, no significant change in the XRD spectra was found between fresh and used catalysts. The average crystallite size of Cu$_{15}$/Al$_2$O$_3$ was estimated to be 13 nm based on FWHM of Cu peaks as determined by the Scherrer equation.

**Figure 3.8.** XRD analysis of selected samples including a fresh and Pt$_{0.2}$Cu$_{12}$/Al$_2$O$_3$ used for butadiene hydrogenation. All catalysts, except the sample used in the hydrogenation reaction, were reduced in H$_2$ prior to XRD. (Cu JCPDS No. 85-1326, Cu$_3$Pt JCPDS No. 65-3247, CuPt JCPDS No. 48-1549)

To gain insight into the electronic states of the Pt species, XANES spectra were collected for Pt monometallic and Pt-Cu bimetallic samples at Pt-L$_{III}$ edge (Figure 3.9). The Pt-Cu
bimetallic and Pt monometallic samples were reduced in 100% H₂ *in situ* prior to the XANES scans. The white line intensity of Pt-Cu samples is significantly lower than that of Pt⁴⁺, Pt²⁺, Pt foil and Pt NPs indicating that the electronic states of Pt are altered in the Pt-Cu bimetallic NPs. Lee and co-workers have reported the decrease in Pt-L₃ white line intensity of CuPt alloys compared to pure Pt. The change was attributed to changes in Pt d₅/₂ density of state upon alloying.¹⁸ Since Pt-L₃ edge characterizes the Pt d states, the decrease in white line intensity of Pt-Cu samples compared to Pt₀.₁/Al₂O₃ and Pt foil indicates the Pt d states were affected in the Pt-Cu alloy samples.¹⁹

![Normalized in situ XANES spectra of selected samples collected at room temperature in H₂ (except where otherwise noted). The Pt-Cu samples were pre-reduced in H₂ at 350 °C in situ, while the Pt₀.₁/Al₂O₃ was pre-reduced in H₂ at 400 °C in situ.](image)

**Figure 3.9.** Normalized *in situ* XANES spectra of selected samples collected at room temperature in H₂ (except where otherwise noted). The Pt-Cu samples were pre-reduced in H₂ at 350 °C *in situ*, while the Pt₀.₁/Al₂O₃ was pre-reduced in H₂ at 400 °C *in situ*. 
3.3.2 Selective hydrogenation of 1,3-butadiene

Our catalytic data reveals that adding a trace amount of Pt to Cu NPs significantly enhances hydrogenation. The butadiene hydrogenation activity and selectivity as a function of temperature on Pt\textsubscript{0.1}Cu\textsubscript{14}/Al\textsubscript{2}O\textsubscript{3}, Pt\textsubscript{0.2}Cu\textsubscript{12}/Al\textsubscript{2}O\textsubscript{3}, and Cu\textsubscript{15}/Al\textsubscript{2}O\textsubscript{3} NPs is shown in Figure 3.10. Under the conditions employed, Pt\textsubscript{0.1}Cu\textsubscript{14}/Al\textsubscript{2}O\textsubscript{3} has a hydrogenation reaction onset at 40 °C, which is 40 °C lower than that of the monometallic Cu catalyst. The reaction rate over the Pt\textsubscript{0.1}Cu\textsubscript{14}/Al\textsubscript{2}O\textsubscript{3} at 60 °C is an order of magnitude higher than the monometallic Cu catalyst. Pt\textsubscript{0.2}Cu\textsubscript{12}/Al\textsubscript{2}O\textsubscript{3} exhibits greater hydrogenation activity than Pt\textsubscript{0.1}Cu\textsubscript{14}/Al\textsubscript{2}O\textsubscript{3} due to a higher Pt atom surface amount due to the increased Pt loading. However, the selectivity remains unaffected because the Pt atoms exist as isolated species in Pt\textsubscript{0.2}Cu\textsubscript{12}/Al\textsubscript{2}O\textsubscript{3} as indicated by EXAFS model fitting. The monometallic Pt\textsubscript{0.1}/Al\textsubscript{2}O\textsubscript{3} catalyst is reactive for 1,3-butadiene hydrogenation at ambient temperature. At 60 °C, the activity of Pt\textsubscript{0.1}/Al\textsubscript{2}O\textsubscript{3} is around 3 times higher than that of the PtCu SAA catalysts, while the selectivity to butenes is significantly lower than that of the PtCu SAA catalysts.
Figure 3.10. Selective hydrogenation of butadiene as a function of temperature over Cu$_{15}$/Al$_2$O$_3$, Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$, and Pt$_{0.2}$Cu$_{12}$/Al$_2$O$_3$ NPs (1,3-butadiene (1.25 %), H$_2$ (20 %) and He (balance), GHSV=1,200 h$^{-1}$).

Notably, the SAA catalysts maintain the high selectivity to butenes exhibited by Cu. At full conversion, there is over 90% selectivity towards butene isomers. The selectivity of SAA NPs (Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ and Pt$_{0.2}$Cu$_{12}$/Al$_2$O$_3$) was comparable to Cu$_{15}$/Al$_2$O$_3$ whereas Pt monometallic catalysts fully converted butadiene to butane under these conditions. Monometallic Pt is known for over-hydrogenating dienes and alkynes.$^{20,21}$ Therefore, by combining the hydrogen activation ability of Pt with the weak binding of butadiene on Cu and the latter’s selectivity to butenes, Pt-Cu SAA catalysts exhibit superior performance for this important industrial reaction. Various other alloy surfaces, including Pd-Au and Sn-Pt, have been shown to improve partial
hydrogenation reactions.\textsuperscript{22,23} However, this is the first report that single Pt atoms can enhance selective hydrogenation reactions on a less active metal, such as copper.

The yields of different products are shown in Figure 3.11. Under the conditions employed, the yield to butane over Pt\textsubscript{0.1}/Al\textsubscript{2}O\textsubscript{3} and Pt\textsubscript{2}Cu\textsubscript{6}/Al\textsubscript{2}O\textsubscript{3} was close to zero.

![Figure 3.11. Conversion and product yields in the selective hydrogenation of butadiene corresponding to Figure 6. (0.4 g catalyst, flow rate = 20 mL/min; 1, 3-butadiene (1.25 %), H\textsubscript{2} (20 %) and He (balance), GHSV=1,200 h\textsuperscript{-1}.)](image)

The stability of Pt-Cu SAA catalysts in butadiene hydrogenation reaction is demonstrated at 160 °C. As shown in Figure 3.12, the production rates of butene isomers (the desired products) and the selectivity to butene are stable for at least 46 hours. The space velocity here is greater (12,000 h\textsuperscript{-1} versus 1,200 h\textsuperscript{-1}) than that of Figure 3.10; thus an accelerated deactivation rate was expected. The spent catalyst used for the stability test was
tested in He-TPD and TPO whereby no desorption of hydrocarbons took place and a negligible amount of CO$_2$ was produced (see below, Figure 3.16), respectively.

**Figure 3.12.** Long-time hydrogenation steady-state activity test over Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ at 160 °C. (~0.1 g catalyst, flow rate = 50 mL/min. 2% 1, 3-butadiene, 20% H$_2$ and balance He. GHSV=12,000 h$^{-1}$. Conversion of butadiene in one hour is around 98%).

Under the same reaction conditions, the PtCu SAAs samples were tested in consecutive ascending and descending temperature cycles. Figure 3.13A shows the conversion of butadiene in a cycle over Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$. The comparison of conversion at 150 °C and 120 °C between the increasing and decreasing temperature ramps shows that the hydrogenation activity was recovered during the decreasing temperature ramp. But at near ambient temperatures, the activity in the cooling part of the cycle is lower. The same was true with all the Pt-Cu alloyed
samples. The results suggest the hydrogenation activity was reduced at lower temperatures but recovered at higher temperatures. Considering that oligomer formation adversely affects the metal-based catalysts in the hydrogenation of hydrocarbons at mild temperatures,\textsuperscript{24} we hypothesize the oligomers partially blocked the surface sites at lower temperature and desorbed with increasing temperature.

To check this, after the first temperature heating-cooling cycle (Figure 3.13A), the catalyst was treated at 350 °C in H\textsubscript{2} to fully desorb the weakly adsorbed hydrocarbons on the surface. We found the selective hydrogenation activity was fully recovered after the high temperature treatment for at least 4 cycles, as shown in Figure 3.13B. This result suggests the active sites were maintained in the cyclic reaction tests.
Figure 3.13. Selective hydrogenation of butadiene as a function of temperature over Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ (0.1 g catalyst, flow rate = 50 mL/min. 2 % 1,3-butadiene, 20 % H$_2$ and balance He. GHSV=12,000 h$^{-1}$) (A) Conversion of butadiene and selectivity to butene with increasing and decreasing temperature ramps. (B) Conversion of butadiene in cyclic hydrogenation experiments, the catalysts were treated in H$_2$ at 350 °C between each cycle. The conversion of the first cycle is shown in (A).

To demonstrate the ability of Pt-Cu SAAs to selectively hydrogenate alkadiene impurities in alkene feedstocks,$^{25}$ we tested the selective hydrogenation activity of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ catalysts in the presence of excess propylene and found that butadiene is preferentially hydrogenated on the Pt-Cu SAA NPs (Figure 3.15). Below 120 °C, conversion of propylene was not observed. At
100% conversion of butadiene, less than 1 % of propylene was converted to propane. Comparing these results to the hydrogenation activity of Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ in propylene-free condition, we found the propylene has no effect on the activity and selectivity of Pt-Cu SAAs for the hydrogenation of butadiene. At 160 °C and 145 °C, butadiene is fully converted and >95% converted, respectively, over Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$ without significant propylene hydrogenation (1.2 % at 160 °C and 0.5 % at 145 °C) for >12 h on-stream at each temperature (Figure 3.14).

Figure 3.14. Butadiene conversion in the presence of excess propylene. Conversion and selectivity in long-time steady state selective hydrogenation of butadiene at 160 and 145 °C (~ 0.1 g catalyst Pt$_{0.1}$Cu$_{14}$/Al$_2$O$_3$, flow rate = 50 mL/min. 2 % 1, 3-butadiene, 20 % propylene, 16 % H$_2$ and balance He. GHSV=12,000 h$^{-1}$).
Figure 3.15. Conversion and selectivity in the selective hydrogenation of butadiene over 
$\text{Pt}_{0.1}\text{Cu}_{14}/\text{Al}_2\text{O}_3$ as a function of temperature. (~ 0.1 g catalyst, flow rate = 50 mL/min. 2 % 1, 3-
butadiene, 20 % propylene, 16 % $\text{H}_2$ and balance He. GHSV=12,000 h$^{-1}$).

Although the catalytic activity of Pt-Cu SAA catalyst is stable at 145 °C, deactivation of the 
catalysts was observed at the lower temperatures. The hysteresis phenomenon in Figure 3.15 
indicates some weakly adsorbed species poison the catalysts at the temperatures lower than 
90 °C but desorbed at the higher temperatures. These weakly adsorbed molecules are very likely 
to be the oligomer of 1,3-butadiene formed through the side reactions.
Our experiments reveal that Pt-Cu SAAs exhibit high stability under realistic hydrogenation conditions. The Pt-Cu SAAs maintain stable butadiene conversion for more than 46 h at 160 °C (Figure 3.12). Temperature-programmed oxidation (TPO) studies of the catalysts after hydrogenation show negligible CO$_2$ formation (Figure 3.16) demonstrating the robustness of SAA catalysts to decomposition, oligomerization, and coke formation. Based on our model catalyst studies, the increase in the selectivity of SAAs is due to the inhibition of hydrocarbon decomposition commonly observed with Pt catalysts because SAAs do not offer extended Pt ensembles where these unfavorable reactions occur. On the other hand, by running at higher temperatures, there are no issues of instability. It is well known that Cu or Pd catalysts for diene and alkyne hydrogenation are affected by oligomer formation at mild temperatures, which leads to instability in their hydrogenation activity.\textsuperscript{24} We found that after reaction at near ambient temperatures, the hydrogenation activity of Pt-Cu SAAs can be fully recovered by treatments at higher temperature, effectively desorbing the adsorbed hydrocarbons (Figure 3.13). Running at the higher temperatures shown Figure 3.12 preserved the catalyst activity without affecting its selectivity.

TPO and He-TPD were performed on both of the spent catalysts that have been used in 48 h hydrogenation of butadiene (Figure 3.12) and 30 h hydrogenation in the presence of propylene (Figure 3.14). In TPO test, CO$_2$ produced by the combustion of carbon species on the catalyst was monitored by mass spectrometry (m/z=44), as shown in Figure 3.16. We found that carbon deposition on the catalyst was negligible. The CO$_2$ concentration was below 0.02 % with TPO performed up to 800 °C. Moreover, for the catalysts used in the hydrogenation of butadiene, the total CO$_2$ production corresponds to less than 1/26,000 of the amount of
butadiene converted over the catalysts. For another spent catalyst, the CO₂ production in TPO corresponds to less than 1/90,000 of the amount of hydrocarbon flowed through the catalysts during the hydrogenation of butadiene in the presence of propylene. Similarly, He-TPD shows no desorption of hydrocarbons up to 800 °C as indicated by the lack of hydrocarbon fragments in the range of m/z=1-120 for both spent catalysts.

**Figure 3.16.** TPO profile of spent Pt₀.₁Cu₁₄/Al₂O₃ catalyst used for the long-time activity test in (a) 2 % 1, 3-butadiene, 20 % propylene, 16 % H₂ and balanced He for around 48 h; (b) 2 % 1, 3-butadiene, 20 % H₂ and balanced He for around 30 h. (TPO condition: 20 % O₂/He, 50 mL/min, 3 °C/min) Scale bar corresponds to 0.05 % CO₂ in the gas stream.
3.4 Collaboration: surface science studies on Pt-Cu(111) SAAs

Surface science studies were conducted by Lucci et al. as one part of the collaborative effort in investigating PtCu SAA for selective hydrogenation of 1,3-butadiene.\textsuperscript{26,27} At low loadings, Pt exists as individual, isolated atoms substituted into the Cu surface. These single Pt atoms activate the dissociation and spillover of H to Cu. The weak binding of butadiene to Cu allows for the highly selective hydrogenation to butenes. Thus, a bifunctional mechanism for hydrogenation of butadiene is confirmed. No decomposition or poisoning of these alloys was observed which can be attributed to the lack of extended Pt surface sites. At higher Pt content in Pt-Cu alloy surfaces, we directly visualized extended Pt surface sites responsible for the reduced selectivity.

3.6 Summary

This chapter reports a novel SAA catalyst design for selective hydrogenation reactions. This is the first such kind of work in the field. A careful galvanic replacement reaction was conducted to prepare PtCu SAA NPs. With comprehensive characterization approaches including EDS, STEM, EXAFS, XANES, XRD, TEM and UV-Vis, the compositional and structural properties of the NP catalysts were thoroughly investigated.

In selective hydrogenation of 1,3-butadiene, PtCu SAA catalysts show activity at near-to-ambient temperature. By adding a small amount of Pt single atoms to the surface of Cu NPs, the onset of hydrogenation activity is lowered for 40 °C. This is due to the single-atom Pt lower the hydrogenation activation barrier on the Cu surface. PtCu SAA catalysts are highly selective
to partial hydrogenation products compared to monometallic Pt. With the presence of excess propylene, butadiene was converted to butenes at more than 95% selectivity with less than 1% conversion of propylene. The activity of SAA is highly stable under the hydrogenation condition. The work reveals the highly selective nature of the SAA catalysts in hydrogenation reactions and demonstrates the single-atom Pt sites can activate the otherwise inert Cu for selective hydrogenation. Furthermore, in addition to their promising selective hydrogenation properties, single-atom alloys provide the ultimate limit for the efficient use of costly catalytic elements like Pt.
3.7 References


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Chapter 4. Tackling CO poisoning with single-atom alloy catalysts

4.1 Introduction

Pt metals are highly active for H\textsubscript{2} dissociation, but are very susceptible to carbon monoxide (CO) poisoning.\textsuperscript{1} Due to its strong binding to Pt, even trace amounts of CO impurity, which is always present in H\textsubscript{2} gas produced from fuel reforming, can diminish H\textsubscript{2} activation and reactivity.\textsuperscript{2}

As the natural expansion of the SAA concept, we tackled the CO poisoning issue of platinum catalysts with Pt-based SAAs. The situation for isolated Pt atoms may be different than traditional bimetallic alloys, where ensemble effects have been shown to increase the CO binding strength at extended catalytic metal sites\textsuperscript{3-5}. From a fundamental perspective, metal alloys that alter the adsorption geometry of CO can exhibit weaker CO binding and enhanced CO tolerance. Isolated Pt atoms enable exclusively atop adsorption configuration of CO, which we show is weaker than CO adsorption on extended Pt sites. In this chapter, we investigated the interaction between CO and Pt sites on a Pt-Cu SAA and demonstrate improved CO tolerance of Pt-Cu SAA catalysts under realistic working conditions for H\textsubscript{2} activation and acetylene hydrogenation reaction.
4.2 Experimental methods

Pt-Cu SAA NPs were prepared with the same galvanic replacement method described in the chapter 3. XAS studies were performed at beamline 12-BM at Argonne National Laboratory. The experiments and data analysis were conducted as described in chapter 3.

FTIR spectra were collected using a Nicolet Nexus 470 spectrometer equipped with a MCT-B detector in the single beam absorbance mode at a resolution of 2 cm\(^{-1}\). Prior to the IR experiments, the samples were reduced in pure H\(_2\) at 300 (Cu-NP), 350 (Pt\(_{0.008}\)Cu-SAA and Pt\(_{0.39}\)Cu-bimetallic) or 400 (Pt-NP) °C in a quartz microreactor and sealed in the glass vials. The powder sample of 0.02 g was made into a thin pellet (~ 1cm in diameter). The sample was then loaded into a flow cell sealed with NaCl windows and allows the CO flow through the catalyst pellet. In a typical CO adsorption IR experiment, the sample was first reduced in 50 mL/min 5 % H\(_2\)-He for 1 hr at 200 °C followed with the He purge to cool down to 25 °C. Baseline was collected at 25 °C under He purge after the catalyst has been swept for at least 30 min. The sample was then exposed to 1 % CO-He flow at the same temperature for 45 min, and was purged again with He for at least another 30 min until the gas phase CO signal was fully diminished (rather quickly within 5 min of purge) and the catalyst surface adsorption spectra has minimal difference between the IR scans. The IR spectra shown in the main text were collected afterwards at 25, 40, 60, 80, and 100 °C. A ramping rate of 5 °C/min was used. The corresponding spectra at each temperature were collected after the sample was stabilized at the target temperature for 10 min.

H\(_2\)-D\(_2\) exchange was conducted in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) with 80-90 mg of catalyst diluted by 0.5 g of quartz particles. The samples were pre-
reduced in 100% H₂ (at 350 °C for Pt₀.₀₀₈Cu-SAA and 400 °C for Pt-NP) and then cooled to desired temperature before the reaction gas mixture (33% H₂, 33% D₂, 3.3% CO and balance Ar. Flow rate=50mL/min) was introduced. For the isothermal H₂-D₂ exchange, CO was introduced to the gas phase and turned off at the certain time points while H₂ and D₂ are flowing. To do temperature programmed H₂-D₂ exchange, a gas mixture of 33% H₂, 33% D₂, 3.3% (or 667ppm) CO and balance Ar (flow rate=50mL/min) was introduced at 120 °C and temperature was increased to 200 °C at 5 °C/min. The gas effluent from the reactor was analyzed by a mass-spectrometer. Acetylene hydrogenation was performed in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) at 70 and 80 °C isothermally (20% H₂, 2.2% C₂H₂, 200 ppm CO, 10% Argon, balance He, flow rate=24-48 mL/min). 50-500 mg catalysts were used. The exit gas stream was analyzed by a mass-spectrometer.

4.3 Results and Discussion

PtCu alloy nanoparticles (NPs) were prepared by galvanic replacement (GR) of Pt for Cu on preformed Cu NPs (Table 4.1)⁶,⁷. Pt₀.₀₀₈Cu supported on γ-Al₂O₃ is a SAA as confirmed by aberration-corrected high angle annular dark field scanning transmission electron microscopy and extend X-ray absorption fine structure (EXAFS) studies⁷. Importantly, with CO in the gas phase, no Pt-Pt interactions were found by EXAFS in Pt₀.₀₀₈Cu-SAA (Table 4.2, Figure 4.1), which demonstrates the stability of the isolated Pt atoms in the presence of CO.
Table 4.1. Sample information.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Composition^[a]</th>
<th>Preparation</th>
<th>Pt Species Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu-NP</td>
<td>14.9 at.% Cu on Al(_2)O(_3)</td>
<td>Colloidal</td>
<td>---</td>
</tr>
<tr>
<td>Pt-NP</td>
<td>0.12 at.% Pt on Al(_2)O(_3)</td>
<td>IWI^[b]</td>
<td>Single atom, clusters and NPs</td>
</tr>
<tr>
<td>Pt(_{0.008})Cu-SAA</td>
<td>0.11at.% Pt, 13.7 at.%</td>
<td>GR</td>
<td>Single atoms in Cu surface^[c]</td>
</tr>
<tr>
<td>Cu on Al(_2)O(_3)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt(_{0.39})Cu-bimetallic</td>
<td>2.2 at.% Pt, 5.6 at.%</td>
<td>GR</td>
<td>Single atom, clusters and NPs ^[d]</td>
</tr>
<tr>
<td>Cu on Al(_2)O(_3)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\^[a] Determined by ICP-AES.

\^[b] Incipient wetness impregnation (IWI).

\^[c] Determined by STEM, EXAFS and CO-IR.

\^[d] Determined by EXAFS and CO-IR.
**XAS studies**

The EXAFS of Pt$_{0.008}$Cu-SAA is plotted in Figure 4.1 and the EXAFS Model fitting results are summarized in Table 4.2. The EXAFS of Pt$_{0.008}$Cu-SAA with CO in the Fourier transform space is different from that of Pt foil. The peak position is at a significantly lower R value compared to that of Pt. This peak is similar to that of PtCu-SAA without CO in the gas phase. Indeed, no Pt-Pt bonds were formed with CO in the gas phase, which demonstrated the stability of the single Pt atoms. The Pt-Cu coordination number (CN) is 10.54 for Pt$_{0.008}$Cu-SAA. The Pt-Cu interaction distance is 2.601 Å, which is smaller than that of Pt-Pt but greater than that of Cu-Cu.$^{7,8}$

![Figure 4.1](image.png)

**Figure 4.1.** Fourier transform of $k^3$-weighted Pt L$_{III}$ EXAFS of Pt foil and Pt$_{0.008}$Cu-SAA in CO and H$_2$ gases mixture plotted in R-space.
### Table 4.2. EXAFS model fitting.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Shell</th>
<th>CN(^{[a]})</th>
<th>R(^{[b]}) (Å)</th>
<th>σ(^2) (Å(^2))</th>
<th>R-factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt foil</td>
<td>Pt-Pt</td>
<td>12</td>
<td>2.764±0.002</td>
<td>0.004</td>
<td>0.001</td>
</tr>
<tr>
<td>Pt(_{0.008})Cu-SAA</td>
<td>Pt-Pt</td>
<td>0</td>
<td>10.54±2.28</td>
<td>2.601±0.006</td>
<td>0.015</td>
</tr>
<tr>
<td>H(_2)+CO</td>
<td>Pt-Cu</td>
<td>10.54±2.28</td>
<td>2.601±0.006</td>
<td>0.007</td>
<td></td>
</tr>
</tbody>
</table>

\(^{[a]}\) CN, coordination number. \(^{[b]}\) R, distance between absorber and backscattered atoms. R-factor, closeness of the fit, if < 0.05, consistent with broadly correct models.

**FTIR studies on adsorbed CO**

The adsorption of CO on Pt-Cu alloys and their monometallic counterparts in NP forms was investigated at ambient pressure by IR spectroscopy. CO-FTIR for Pt-NP is shown in Figure 4.2. The absorption peak centered at around 2033-2020 cm\(^{-1}\) is assigned to linearly CO adsorbed on Pt, while the broad band centered at around 1808 cm\(^{-1}\) is attributed to the bridge CO on Pt. This is in agreement with the typical CO-IR spectra for supported Pt nanoparticles.\(^9,^{10}\)
Figure 4.2. IR spectra in the carbonyl range of pre-reduced Pt-NP.

Figure 4.3 reveals the CO-FTIR of Pt$_{0.39}$Cu-bimetallic. The peaks centered at 2110 cm$^{-1}$ are ascribed to linearly adsorbed CO on Cu sites, and the peak at 2043-2029 cm$^{-1}$ is assigned to linearly adsorbed CO on Pt sites in the bimetallic catalysts.$^{11-15}$ The CO-Pt peak is slightly blue shifted compared to Pt-NP, which might be due to the CO surface coverage effect or the alloying effect.$^{16}$ The FWHM of the atop CO peak on Pt$_{0.008}$Cu-SAA is around 25 cm$^{-1}$, which indicates the relatively non-uniform distribution of the Pt sites.$^{17,18}$ The bridge CO peak appears at around 1800 cm$^{-1}$.$^{19,20}$ The existence of bridge adsorbed CO indicates there are CO clusters or extended Pt ensembles in Pt$_{0.39}$Cu-bimetallic, which is in agreement with the aforementioned EXAFS data fitting results.
Figure 4.3. IR spectra in the carbonyl range of pre-reduced Pt$_{0.39}$Cu-bimetallic.

In CO-IR of Pt$_{0.008}$Cu-SAA (Figure 4.4), we do not see the bridge CO peaks between 1900 and 1700 cm$^{-1}$, as observed on the Pt-NP and Pt$_{0.39}$Cu-bimetallic. This indicates the highly dispersed Pt atoms in the surface of Cu, which is consistent with the fact that exclusively isolated Pt atoms are in the surface of Pt$_{0.008}$Cu-SAA, unlike the existence of a mixed phase of Pt atoms, clusters and particles on the Pt-NP and Pt$_{0.39}$Cu-bimetallic. Overlapping peaks between 2150 and 2060 cm$^{-1}$ are seen for Pt$_{0.008}$Cu-SAA, as indicated by the asymmetric absorption band (Figure 4.5 B). The unique absorption band centered at ca. 2088 cm$^{-1}$ is distinct from atop CO on the Cu or Pt NPs, and is assigned to the linearly adsorbed CO on the isolated single Pt atoms. The absorption band of linearly adsorbed CO is at ca. 2020 cm$^{-1}$ on Pt-NP, 2031 cm$^{-1}$ on the Pt ensembles on Pt$_{0.39}$Cu-bimetallic and 2093 cm$^{-1}$ on Cu-NP (Figure 4.3, 4.4 and 4.5). As for the
different Pt species characterized by CO-IR, their formal charges and coordination numbers are essentially the same, and dipole-dipole coupling has little effect on Pt$_{0.008}$Cu-SAA due to the weak CO-metal bonds on isolated Pt atoms and Cu surfaces$^{21}$. Hence, the vibrational frequency of adsorbed CO indicates the electronic properties of the Pt atoms in the surface of catalysts. The change of electronic states of Pt atoms is also observed by aforementioned XANES studies. The change in electronic states will affect the binding strength to CO. In principle, the higher $\nu$(CO) indicates less back donation from Pt to CO molecule. Moreover, the ensemble effect of isolated Pt atoms can lower the CO adsorption strength by eliminating the more strongly adsorbed bridge CO. So PtCu SAA is a promising catalyst with low CO binding strength. Indeed, it has been shown by DFT calculation that the adsorption strength of CO on the Cu$_3$Pt(111) alloy is lower than Pt(111)$^{22}$. 
Figure 4.4. IR spectra in the carbonyl range of pre-reduced Pt$_{0.008}$Cu-SAA.

Figure 4.5. IR spectra in the carbonyl range of pre-reduced (A) Pt$_{0.008}$Cu-SAA and (B) Cu-NP.
The temperature effects on the adsorption of CO on Pt$_{0.008}$Cu-SAA, Pt$_{0.39}$Cu-bimetallic and Pt-NP catalysts were investigated. Figure 4.6 summarizes the IR absorption intensity of atop CO at Pt sites for different samples was compared as temperature is increased from 20 to 60 °C. The intensities of the CO adsorption peaks decreased with increased temperature. Pt$_{0.008}$Cu-SAA shows most significant temperature effect as the CO peak intensity decreases to half of the initial value with temperature increases from 20 to 60 °C. While the CO peak intensity on Pt$_{0.39}$Cu-bimetallic decreases to 80%. The CO peak decrease to a further less value on Pt-NP. These results indicate that the CO-Pt bond is the weakest on the Pt$_{0.008}$Cu-SAA compared to Pt-NP and Pt$_{0.39}$Cu-bimetallic.

![Figure 4.6](image)

**Figure 4.6.** Temperature effects on IR intensity of the linearly adsorbed CO on Pt sites of (A) Pt$_{0.008}$Cu-SAA, (B) Pt$_{0.39}$Cu-bimetallic and (C) Pt-NP. The vertical axis is the ratio of the peak absorbance at the given temperature to the peak absorbance at 25 °C. The peak vibrational frequency of each carbonyl is (A) 2088 cm$^{-1}$, (B) 2045-2031 cm$^{-1}$ and (C) 2033-2018 cm$^{-1}$. 
Table 4.3. CO-IR peak assignments

<table>
<thead>
<tr>
<th>Peak position (cm(^{-1}))</th>
<th>Assignments</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>2092-2110</td>
<td>Atop CO on Cu species (Cu-NP, Pt(<em>{0.39})Cu-bimetallic, Pt(</em>{0.008})Cu-SAA)</td>
<td>11,12,16,23</td>
</tr>
<tr>
<td>ca. 2088</td>
<td>Atop CO on isolated Pt atoms in the surface of Cu NPs (Pt(_{0.008})Cu-SAA)</td>
<td></td>
</tr>
<tr>
<td>2018-2043</td>
<td>Atop CO on extended Pt species (Pt-NP, Pt(_{0.39})Cu-bimetallic)</td>
<td>14,16,19,20,24</td>
</tr>
<tr>
<td>1788-1812</td>
<td>Bridging CO on two adjacent Pt sites (Pt-NP, Pt(_{0.39})Cu-bimetallic)</td>
<td>19,20</td>
</tr>
</tbody>
</table>

\(H_2\)-\(D_2\) exchange in the presence of CO

\(H_2\)-\(D_2\) exchange was employed to characterize the \(H_2\) dissociation activity on the corresponding nanocatalysts at ambient pressure in a flow reactor setup.\(^{25}\) As described in chapter 3, the \(H_2\) activation barrier on Cu surface is significantly higher than that of on extended Pt surface or Pt single atoms. So the \(H_2\) dissociation sites on PtCu-SAA is Pt single atoms. At the meantime, the \(H_2\) desorption barrier on single Pt atoms is lower than that of Cu surface. So the single-atom Pt sites also act as the exit of the \(H_2\) molecule. Thus, the \(H_2\)-\(D_2\) exchange characterizes
the activity and availability of the Pt atoms in the surface of PtCu-SAA under practical reaction conditions.

As shown in Figure 4.7, HD is initially produced when H₂ and D₂ are co-fed into the reactor at a certain temperature. Once CO (3.3%) is introduced, the HD production decreases, which indicates the CO poisoning of the Pt sites. With the same CO concentration in the gas phase, Pt₀.₀₀₈Cu-SAA yields 12 times more HD than Pt-NP at 150 °C. Since the CO coverage is a function of binding strength, and we have shown that CO binds to Pt-Cu SAAs more weakly compared to pure Pt, the reduced binding strength of CO to Pt₀.₀₀₈Cu-SAA yields more CO-free sites that are available for H₂ activation and hence HD production. In other words, Pt₀.₀₀₈Cu-SAA is one order of magnitude more CO tolerant compared to monometallic Pt catalysts under the reaction conditions employed.

The temperature effects on the CO tolerance of Pt catalysts are also investigated. As shown in Figure 4.7, the H₂-D₂ exchange in the presence of CO was studied at 120, 150 and 200 °C. At higher temperatures, the CO effect on HD production decreases. Moreover, Pt₀.₀₀₈Cu-SAA shows significantly higher HD production at different temperatures with the presence of 3.3% CO. This further confirms the CO tolerance of Pt₀.₀₀₈Cu-SAA catalysts at different operating conditions.
Figure 4.7. Isothermal $\text{H}_2$-$\text{D}_2$ exchange over Pt-NP (A, B, C) and $\text{Pt}_{0.008}\text{Cu}$-SAA (D, E, F) at 120, 150 and 200 $^\circ$C. CO was turned on and off during the experiments. Gas composition: 33% $\text{H}_2$, 33% $\text{D}_2$, 3.3% CO, balance Argon. Total flow rate=50 mL/min. 90 mg catalysts.
The CO tolerance of Pt\textsubscript{0.008}Cu-SAA was studied in temperature-programed H\textsubscript{2}-D\textsubscript{2} experiments, over a wide temperature range (120 to 200 °C) with CO concentrations from 667 ppm to 3.3% (Figure 4.8). At this temperature range, Pt\textsubscript{0.008}Cu-SAA shows higher HD production compared to that of Pt-NP. Moreover, Pt\textsubscript{0.008}Cu-SAA outperforms Pt-NPs at lower CO concentration (667 ppm) as well, which demonstrates the broad operation window of Pt-Cu SAA as a CO-tolerant catalyst.

**Figure 4.8.** H\textsubscript{2}-D\textsubscript{2} exchange as a function of temperature. Gas composition: 33% H\textsubscript{2}, 33% D\textsubscript{2}, 667ppm CO (A), or 3.3% CO (B) balance Argon. Total flow rate=50 mL/min. 80 mg catalyst loading.

H\textsubscript{2} activation is the rate limiting step for hydrogenation reactions and the key elementary step the anode reaction of H\textsubscript{2} fuel cells. The H\textsubscript{2}-D\textsubscript{2} exchange with and without CO indicates Pt-Cu
SAA as a class of promising CO-tolerant catalysts for fuel cells and other applications in the chemical industry.

To investigate the CO-tolerance of Pt-Cu SAA under realistic catalytic reaction conditions, we conducted the selective hydrogenation of acetylene over Pt$_{0.008}$Cu-SAA and Pt-NP with and without CO in a flow reactor. Figure 4.9 plots the ratio between reaction rate with 200 ppm CO in the gas phase and rate without CO. The reactions were performed at temperatures lower than 80 °C and with 200 ppm CO, which are very close to the operating conditions of the hydrogen fuel cells. We found about half of the reaction rate was retained for Pt$_{0.008}$Cu-SAA. However, the hydrogenation activity of Pt-NP decreased 15-fold with 200 ppm CO. The H$_2$-D$_2$ exchange and acetylene hydrogenation activity of Cu-NP was at least one order of magnitude lower than that of Pt$_{0.008}$Cu-SAA at these temperatures. Our results demonstrate that Pt-Cu SAAs offer a new approach to design CO-tolerant materials for industrial applications.
Figure 4.9. Ratio of retained reaction rate of acetylene hydrogenation in CO over Pt$_{0.008}$Cu-SAA and Pt-NP at different temperatures. Rate retained = reaction rate with CO/rate without CO in the reaction gas. Gas composition: 20% H$_2$, 2.2% C$_2$H$_2$, 200 ppm CO, 10% Argon, balance He. Flow rate = 24 mL/min.

4.3 Collaboration: CO-TPD and STM on Pt-Cu(111) SAA

In agreement with the Pt surface structure observed by scanning tunneling microscopy (STM)$^{26,27}$, the CO desorption profiles reveal the development of 3 distinct Pt-Cu alloyed structures with increasing Pt coverage. With atomic resolution, the Pt atoms image ~20 pm taller than the surrounding Cu lattice and appear as brighter protrusions. At low Pt coverages, CO desorbs from isolated Pt atoms (peak at 350 K) and at intermediate coverages, CO desorbs from extended ensembles of Pt atoms (peak at 480 K). At 5 ML Pt, the surface layer is composed primarily of continuous ordered Pt atoms and CO desorbs at 410 K. At low CO
coverages, CO desorption from Pt(111) is reported at 450 K and with increasing CO coverage desorption is observed at 400 K due to the repulsion between co-adsorbed CO molecules\textsuperscript{28}. In Pt-Cu alloys, CO adsorbs weakest on the Pt-Cu SAA and strongest on the extended Pt clusters.

The mobility of H atoms on Pt-Cu SAAs was demonstrated in UHV by co-adsorption of H\textsubscript{2} and D\textsubscript{2} onto SAA Pt-Cu(111). We observed desorption of HD (m/z = 3) which provides evidence of the dissociation of H\textsubscript{2} and migration of H atoms away from the dissociation site. When equal amounts of H and D atoms are adsorbed on the surface, we observe complete scrambling of H and D with ratio of H\textsubscript{2}:HD:D\textsubscript{2} of 1:2:1 due to H atom spillover and its facile diffusion on Cu.

4.4 Summary

In this chapter, we investigated the fundamentals of CO adsorption on Pt-Cu alloys using a variety of catalysis techniques. We found that CO binds weaker to single Pt atoms in Cu compared to larger Pt ensembles or monometallic Pt. Thus, when CO is present in the gas phase, more CO-free Pt sites are available on the SAA than the monometallic Pt, yielding higher hydrogenation activity under realistic conditions. We postulate that this SAA approach in other metal combinations can be used for the design of novel CO-tolerant catalysts for other industrially important reactions such as hydrocarbon and alcohol oxidation reactions, and for the present generation, low-temperature hydrogen fuel cells where the platinum electrocatalyst is prone to CO-poisoning.

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Chapter 5. PtCu single-atom alloy catalysts for coke resistant C-H activation and selective dehydrogenation of n-butane

5.1 Introduction

The production of shale gas from hydraulic fracturing has boosted the supply of light alkanes such as ethane and propane in recent years.\textsuperscript{1,2} However, there are few low-carbon footprint methods that efficiently convert these gases to more valuable chemicals due to their relative inertness.\textsuperscript{3} While steam cracking is the traditional industrial process, it is energy-demanding and less than 60\% efficient.\textsuperscript{2} Catalytic processes are sought after to potentially improve the energy balance sheet. Facile activation of C-H bonds in alkanes would open new routes to synthesize commodity and fine chemicals.\textsuperscript{3-6} There are numerous ways to functionalize alkanes including alkane halogenation/oxyhalogenation, oxidative coupling, and oxidative dehydrogenation.\textsuperscript{7-10} This chapter focused on the dehydrogenation of alkanes to produce alkenes, which are precursors to industrially relevant polymers.\textsuperscript{1,6,11,12} Preventing coking is indeed an active, yet challenging area of current research.\textsuperscript{13} Pt catalysts also suffer from coking and Pt’s high price prohibits widespread use.\textsuperscript{14-16} Cu catalysts are typically not considered viable due to a high C-H activation barrier on Cu surfaces, but are resistant to coking.\textsuperscript{17}

Alloys often exhibit unique properties compared to their constituent metals.\textsuperscript{18,19} For example, using a combination of surface science, theory, and catalysis, Besenbacher et al. showed that small amounts of Au dispersed in Ni can suppress carbon deposition in methane
steam reforming by both raising the barrier to C-H activation and decreasing the binding strength of carbon to the surface.\textsuperscript{18} We took the opposite approach in this work, using the smallest amount of a catalytic metal (Pt) in the form of single atoms in the surface layer of a more inert host metal (Cu) to facilitate C-H activation while avoiding coking that typically occurs on larger ensembles.\textsuperscript{19} These single-atom alloys (SAAs)\textsuperscript{20–22} are analogous to single-atom catalysts\textsuperscript{23,24}, but unlike many single-atom catalysts, reaction at the active site can be understood with atomistic detail on SAAs.

PtCu alloys activate C-H bonds with significantly improved activity over Cu, while avoiding coking that occurs on catalysts with extended Pt ensembles due to Cu’s ability to facilitate C-C coupling chemistry. These PtCu alloy nanoparticle catalysts exhibit reaction temperatures significantly lower than Cu under realistic operating conditions. The Pt-Cu SAAs were demonstrated to be highly stable and coke-resistance in butane dehydrogenation reactions.

5.2 Experimental methods

PtCu NP catalysts were prepared as described in chapter 2.\textsuperscript{21,25,26} Cu NPs prepared as colloids by reducing Cu(NO$_3$)$_2$ in aqueous solution with NaBH$_4$ as reducing agent and polyvinylpyrrolidone as capping agent. The resulting copper NPs were deposited onto a silica support (fumed silica, 0.2-0.3 μm average aggregate particle size, surface area 200 m$^2$/g)
followed by calcination in air at 300 °C.\textsuperscript{21,26,27} \textit{Pt}_{0.03}\textit{Cu}-SAA and \textit{Pt}_{0.01}\textit{Cu}-SAA were prepared by the galvanic replacement method on the copper NPs.\textsuperscript{21,26}

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) and X-ray absorption spectroscopy (XAS) were conducted using the same protocols as described in chapter 3 and 4. DRIFTS was conducted using a Thermo Nicolet i550 FT-IR equipped with a Harrick DRIFTS cell.\textsuperscript{25} The samples were pre-reduced in H\textsubscript{2} at 400 °C \textit{in situ}. Samples were degassed with helium for 15 minutes before introducing CO. After 10 minutes CO exposure, helium was passed through the sample while the IR spectra were collected.

Temperature-programmed oxidation (TPO) experiments were performed in a packed-bed flow microreactor (L = 22 inch, O.D. = 1/2 inch) following the long-hour butane-D\textsubscript{2} isotope scrambling (B-D scrambling) experiments. The samples were treated in Argon (pure, 50 mL/min) at 300 °C for 2 h prior to the TPO. After cooling to ambient temperature in Argon, 20 % O\textsubscript{2}/He (20 mL/min) was introduced to the reactor and allow the system to stabilize for one hour. To perform TPO, the temperature was increased from ambient temperature to 600 °C at 3 °C/min. The gas effluent from the reactor was analyzed by a mass-spectrometer.

TEM was conducted on a JEOL 2010 electron microscope with 200 kV and 107 μA beam emission. The specimens were obtained by adding one drop of ethanol solution of nanoparticles onto a carbon film with a nickel covered microgrid.

B-D scrambling and butane dehydrogenation experiments were performed in a packed-bed flow microreactor (L=22 inch, O.D.=1/2 inch) with 100 mg of catalyst diluted by 1 g of quartz particles. Prior to the introduction of reaction gas mixture, the samples were reduced in
20% H$_2$/He at 400 °C for 20 minutes and cooled to ambient temperature. After stabilizing in the reaction gas mixture at ambient temperature for 1 hour, the temperature was raised to the desired point. The gas effluent from the reactor was analyzed by a mass-spectrometer. To perform temperature programmed B-D scrambling, the temperature was increased from ambient temperature to 400 °C at 5 °C/min.

For butane dehydrogenation reactions, the samples were reduced in 20% H$_2$/He at 400 °C for 20 minutes and cooled to ambient temperature. After stabilizing in the reaction gas mixture at ambient temperature for 1 hour, the temperature was raised to the desired point. The gas effluent from the reactor was analyzed by a mass-spectrometer. For the apparent activation energy measurements, the catalysts were tested in steady state at highest temperature followed by decreasing the temperature to the desired points.

5.3 Results and discussion

Pt$_{0.03}$Cu-SAA and Pt$_{0.01}$Cu-SAA NPs were prepared by a galvanic replacement reaction as reported in Lucci et al. and Liu et al.$^{21,26}$ and supported on silica (Table 5.1). We conducted extended X-Ray absorption fine structure (EXAFS) and Fourier transform infrared spectroscopy (FTIR) studies to confirm the formation of NPs with isolated Pt atoms.
Table 5.1. Sample Information

<table>
<thead>
<tr>
<th>Sample</th>
<th>Composition (^{[a]})</th>
<th>Preparation Method</th>
<th>Pt Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu-NP</td>
<td>3.6 wt.% Cu on SiO(_2)</td>
<td>Colloidal</td>
<td>---</td>
</tr>
<tr>
<td>Pt-NP</td>
<td>0.5 wt.% Pt on SiO(_2)</td>
<td>IWI(^{[b]})</td>
<td>NPs</td>
</tr>
<tr>
<td>Pt(_{0.03})Cu-SAA</td>
<td>0.25 wt.% Pt, 3 wt.% Cu on SiO(_2)</td>
<td>Galvanic replacement</td>
<td>Single atoms in Cu surface(^{[c]})</td>
</tr>
<tr>
<td>Pt(_{0.01})Cu-SAA</td>
<td>0.1 wt.% Pt, 3.4 wt.% Cu on SiO(_2)</td>
<td>Galvanic replacement</td>
<td>Single atoms in Cu surface(^{[d]})</td>
</tr>
</tbody>
</table>

\(^{[a]}\) Determined by ICP-AES.

\(^{[b]}\) Incipient wetness impregnation (IWI).

\(^{[c,d]}\) Determined by STEM in ref. 9 and EXAFS and CO-IR in this work.

Figure 5.1 and 5.2 show the FTIR spectra for Pt\(_{0.01}\)Cu-SAA and Pt\(_{0.03}\)Cu-SAA with CO chemisorption, respectively. The peaks at 2130 to 2112 cm\(^{-1}\) are assigned to linearly adsorbed CO on Cu and Pt respectively.\(^{26,28-31}\) Peaks corresponding to linearly adsorbed CO and bridge CO on extended Pt surfaces are not observed in the SAA samples, which typically appear at 2018-2043 cm\(^{-1}\) and 1780-1820 cm\(^{-1}\), respectively.\(^{31-35}\) This suggests that Pt atoms are fully isolated in the copper surface. The CO peaks observed are not shifted as the CO coverage is changed, which indicates small dipole-dipole interaction because of the high dispersion of individual Pt atoms and low CO coverage on the Cu surface.
Figure 5.1. FTIR spectra of chemisorbed CO molecule on Pt$_{0.01}$Cu-SAA catalyst. The time values on the graph indicate the time of helium gas purge after CO exposure.
Figure 5.2. FTIR spectra of chemisorbed CO molecule on the Pt\textsubscript{0.03}Cu-SAA catalyst. The time values on the graph indicate the time of helium gas purge after CO exposure.

Figures 5.3 and 5.4 show the EXAFS of Pt\textsubscript{0.01}Cu-SAA, Pt\textsubscript{0.03}Cu-SAA and Pt foil plotted in k-space and Fourier transform space (R-space). The EXAFS Pt\textsubscript{0.01}Cu-SAA and Pt\textsubscript{0.03}Cu-SAA are significant different from that of Pt foil. EXAFS at Pt edge characterize the coordination of Pt atoms, especially in the first shell. Pt\textsubscript{0.01}Cu-SAA and Pt\textsubscript{0.03}Cu-SAA have Pt-Cu interaction peak at lower R value in comparison to that of Pt-Pt interaction peak of Pt foil. EXAFS model fitting suggests there is no Pt-Pt interaction in Pt\textsubscript{0.01}Cu-SAA or Pt\textsubscript{0.03}Cu-SAA in the first coordination shell but only in Pt-Cu interactions (Table 5.2). This further confirms that Pt atoms are fully isolated in the Cu NPs. The first shell bond length (2.59 and 2.61 Å) of SAA NPs is 100% Pt-Cu without Pt-Pt contribution. This is consistent with the formation of SAAs.
Table 5.2. EXAFS model fitting.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Shell</th>
<th>CN(^{[a]})</th>
<th>(R^{[b]}) (Å)</th>
<th>(\sigma^2) (Å(^2))</th>
<th>R-factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt foil</td>
<td>Pt-Pt</td>
<td>12</td>
<td>2.764±0.002</td>
<td>0.004</td>
<td>0.001</td>
</tr>
<tr>
<td>Pt(_{0.01})Cu-SAA</td>
<td>Pt-Pt</td>
<td>0</td>
<td></td>
<td></td>
<td>0.013</td>
</tr>
<tr>
<td></td>
<td>Pt-Cu</td>
<td>10.51±1.62</td>
<td>2.591±0.004</td>
<td>0.005</td>
<td></td>
</tr>
<tr>
<td>Pt(_{0.03})Cu-SAA</td>
<td>Pt-Pt</td>
<td>0</td>
<td></td>
<td></td>
<td>0.016</td>
</tr>
<tr>
<td></td>
<td>Pt-Cu</td>
<td>10.80±1.75</td>
<td>2.609±0.004</td>
<td>0.006</td>
<td></td>
</tr>
</tbody>
</table>

\(^{[a]}\) CN, coordination number. \(^{[b]}\) R, distance between absorber and backscattered atoms. R-factor, closeness of the fit, if < 0.05, consistent with broadly correct models.

Figure 5.3. *In situ* Pt L\(_{III}\) EXAFS of (a) Pt\(_{0.01}\)Cu-SAA, (b) Pt\(_{0.03}\)Cu-SAA and (c) Pt foil plotted in k space.
Figure 5.4. Fourier transform of $k^3$-weighted Pt L$_{III}$ EXAFS of (a) Pt$_{0.01}$Cu-SAA, (b) Pt$_{0.03}$Cu-SAA and (c) Pt foil plotted in R space.

Butane – deuterium scrambling (B-D scrambling)

We compared the performance of Cu NPs and PtCu SAA NP catalysts for B-D scrambling using temperature programmed surface reaction (TPSR) (Figure 5.5). Butane-deuterium isotope scrambling (B-D scrambling) experiments were performed with nanoparticle (NP) catalysts in a flow reactor. The production rate of deuterated butane in B-D scrambling experiments indicates the C-H activation capability of the catalysts as hydrogen (deuterium) dissociation is a facile process under the reaction conditions applied.$^{21,26}$ The reaction was operated within the kinetics controlled regime (conversion < 5%), so that the effluent concentration of B-D scrambling products is proportional to the overall C-H activation rate.
The second cycle data were plotted here as there is no activity difference between the second and third cycle. In Figure 5.5, the ratio between mass 59 and 58 reflects the ratio between deuterated butane (C₄H₉D) and butane (C₄H₁₀). Pt₀.₀₃Cu-SAA and Pt₀.₀₁Cu-SAA start to convert butane (C₄H₁₀) to deuterated butane (C₄H₉D) at around 250 °C but Cu-NP is active only at temperatures above 500 °C (Figure 5.6). This demonstrates that the activity of B-D scrambling i.e. C-H activation, is significantly improved by the addition of a small amount of isolated Pt atoms into the Cu surface. At the same condition, the silica support is not active for B-D scrambling.

![Mass-spec Signal 59/58](image.png)

**Figure 5.5.** Reactor studies of B-D scrambling. TPSR data for the B-D scrambling reaction over Pt₀.₀₃Cu-SAA, Pt₀.₀₁Cu-SAA and Cu-NP catalysts, followed by mass spectrometry. (Temperature ramp 5 °C/min). The 59/58 mass ratio indicates the C₄H₉D level. The data are from the second
reaction cycle. Gas composition: 5% butane, 2% deuterium and balance argon. 50 mL/min, 100 mg catalysts, 1 g quartz beads. Collaborative work with Joshua Wimble.\textsuperscript{36}

Figure 5.6. Temperature programmed surface reaction data for the B-D scrambling reaction over Cu-NP catalyst. Collaborative work with Joshua Wimble.\textsuperscript{36}

Figure 5.7 shows the B-D scrambling TPSR of Pt-NP for the first three cycles up to 400 °C. In the first cycle, Pt-NP begins to catalyze the B-D scrambling around 100 °C. Although the Pt-NP deactivated in the second and third cycles, we can still observe B-D scrambling activity at temperatures higher than 100 °C. Thus, TPSR finds the onset of C-H activation with PtCu-SAA, Cu-NP and Pt-NP at 250, 550, and 100 °C, respectively. This suggests the barriers of C-H
activation on the NP catalysts follow the order: Pt < PtCu-SAA < Cu. This corroborates the trend of C-H activation barriers identified in the model catalyst study\textsuperscript{37}.

![Image of TPSR data](image.png)

**Figure 5.7.** Reactor studies of B-D scrambling. TPSR data for the B-D scrambling reaction over Pt-NP catalysts, followed by mass spectrometry. (Temperature ramp 5 °C/min). The 59/58 mass ratio indicates the C\textsubscript{4}H\textsubscript{9}D level. First, second and third cycles of TPSR are shown. Gas composition: 5% butane, 2% deuterium and balance argon. 50 mL/min, 100 mg catalyst, 1 g quartz beads. Collaborative work with Joshua Wimble.\textsuperscript{36}

Monometallic Pt NP catalysts rapidly deactivate during the reaction (Figure 5.7). In the first cycle of TPSR, the production of C\textsubscript{4}H\textsubscript{9}D ‘lights-off’ at 100 °C and increases with increasing
temperature up to 230 °C. The production rate of C₄H₉D then drops off after 230 °C indicating significant deactivation, overcoming the effect of temperature increase on reaction rate. The second and third cycles of TPSR show much lower activity compared to the first cycle and the activity decreases cycle by cycle. On the other hand, PtCu-SAAs are very stable. Figure 5.8 shows that Pt₀.₀₁Cu-SAA catalyzes the B-D scrambling at 360 °C without any deactivation for at least 12 hours. The deactivation of Pt-based catalysts in alkane C-H activation reactions is mainly due to coke formation and sintering of Pt NPs, while the coke formation is the major cause in most cases. However, Pt₀.₀₁Cu-SAA does not experience any deactivation under the reaction conditions employed.

**Figure 5.8.** Reactor studies of B-D scrambling. 12-hour stability test for the Pt₀.₀₁Cu-SAA catalyst at 360 °C. The catalyst was at room temperature at the beginning and end of the test. Gas composition: 5% butane, 2% deuterium and balance argon. 50 mL/min, 100 mg catalysts, 1 g quartz beads. Collaborative work with Joshua Wimble.³⁶
Figure 5.9. Determination of carbon deposition. Temperature programmed oxidation (TPO) of (a) Pt-NP, (b) Pt$_{0.012}$Cu-SAA and (c) silica. (a) and (b) were treated in butane-D2 exchange reaction condition at 360 °C for 12 hours. (a), (b) and (c) were treated in argon flow at 300 °C for 2 hours before TPO to desorb the hydrocarbon adsorbates. Collaborative work with Joshua Wimble.\textsuperscript{36}

We performed temperature-programmed-oxidation (TPO) on used catalysts to determine whether carbon deposition occurs during B-D scrambling. Pt-NP and Pt$_{0.01}$Cu-SAA were both exposed to B-D scrambling conditions at 360 °C for 12 hours prior to TPO measurements. As shown in Figure 5.9, a broad CO$_2$ evolution from the used Pt-NP is seen, beginning at 130 °C. At least two different types of carbon are shown, the second matching graphitic carbon oxidation\textsuperscript{38}, while any adsorbed hydrocarbon species must have desorbed
during the Ar heat treatment. These features are absent on the Pt$_{0.01}$Cu-SAA. The amount of CO$_2$ produced from this catalyst is essentially the same as that produced from the silica support without any hydrocarbon exposure. Thus, PtCu SAAs are coke-resistant at atmospheric pressure reaction conditions while Pt NPs catalyze carbon formation and deposition, which causes severe deactivation of the catalyst during the B-D scrambling conditions.

In catalysis with exclusively single-atom active sites, a linear relationship between active metal loading and reactivity was reported$^{39,40}$. Here, a linear relationship between the B-D scrambling rate and Pt loading in the catalysts was demonstrated for the PtCu SAA catalysts, as shown in Figure 5.10. This corroborates the fact that Pt atoms are fully isolated in Pt$_{0.01}$Cu-SAA and Pt$_{0.03}$Cu-SAA.

![Figure 5.10](image.png)

**Figure 5.10.** B-D scrambling rate as a function of Pt loading in the catalysts. Gas composition: 5% butane, 2% deuterium and balance argon. 50 mL/min, 100 mg catalyst, 1 g quartz beads.
We increased the Pt/Cu ratio to 0.39 in Pt\textsubscript{0.39}Cu catalyst, which is no longer SAA as indicated by CO-IR and EXAFS model fitting in previous studies\textsuperscript{41} that Pt clusters are formed at this Pt loading level. As shown in Figure 5.11, in the first cycle, the reaction rate starts to drop at 390 °C, even with increasing temperature. The B-D scrambling reactivity of the catalyst decreases with cyclic operation. This indicates the deactivation of PtCu bimetallic catalysts when extended Pt ensembles are present.

![Graph showing mass spec signal vs. temperature for different cycles](image)

**Figure 5.11.** Reactor studies of B-D scrambling. TPSR data for the B-D scrambling reaction over Pt\textsubscript{0.39}Cu followed by mass spectrometry. (5 °C/min) Gas composition: 5% butane, 2% deuterium and balance argon. 50 mL/min, 100 mg sample, 1 g quartz beads.

In model catalyst studies, the Sykes group used methyl groups adsorbed on the single-atom Pt sites in the surface of Cu(111) and demonstrated the C-H activation barrier of them is significantly lower than that on Cu(111).\textsuperscript{37} DFT calculation indicates single-atom Pt binds the hydrocarbon intermediates weaker compared to Pt(111) but stronger compared to Cu(111). The strong adsorption of hydrocarbon intermediates on Pt(111) causes the over-
dehydrogenation as the favorable pathway instead of coupling or desorption. Overall, single-atom Pt sites are active for C-H activation but not active in over-dehydrogenation or C-C bond breaking.

**Nonoxidative dehydrogenation of butane**

Encouraged by the success in the coke resistance B-D scrambling with SAA catalysts, we explored the nonoxidative dehydrogenation of butane with SAA catalysts. At 400 °C, Pt$_{0.01}$Cu-SAA stably converts butane to butene for at least 52 hours without any deactivation (Figure 5.13). An Arrhenius type plot for butane dehydrogenation over Pt$_{0.01}$Cu-SAA and Pt-NP catalysts is shown in Figure 5.12. The apparent activation energy of the reaction over Pt$_{0.01}$Cu-SAA and Pt-NP is 112 and 51 kJ/mol, respectively. The rate limiting step for light alkane dehydrogenation is the first C-H activation. We have found the C-H activation barrier over Pt$_{0.01}$Cu-SAA is higher than that of Pt-NP from the B-D scrambling studies. Thus, the trend we observed in the apparent activation energy of butane dehydrogenation is in agreement with the trend in the C-H activation barrier. The high stability of Pt$_{0.01}$Cu-SAA in butane dehydrogenation conditions is also in agreement with its stability in B-D scrambling conditions.

In the industrial Pt-Sn alloy based catalysts for selective dehydrogenation of light alkanes, Pt is the major active phase and Sn is added to mitigate the coke formation. Due to the breaking of the Pt ensembles and the electronic effect as electrons are transferred to Pt 5d band, the active sites for the cracking and hydrogenolysis are reduced. In other studies, addition of a second metal such as Ga, In, Li, Na and K lead to lower acidity of the catalysts,
which results in lower coke formation. This thesis reports a novel approach for the alkane dehydrogenation catalysis that the single atom Pt active sites are formed in the surface of a relatively inert but highly selective metal (Cu). Thus, no active sites for side reactions exist in the first place. And this catalyst is proved to be stable in selective alkane dehydrogenation conditions. Moreover, the single-atom configuration allows the highest material efficiency of precious Pt metal compared to the nanoparticle configurations.

Figure 5.12. Arrhenius-type plot. The reaction rate of butane dehydrogenation was measured between 450 and 390 °C with conversion below 10%. Reaction condition: 2.5% butane, 5% H₂, bal. He. 50 mL/min, 30-80 mg sample.
Figure 5.13. Long-term butane dehydrogenation reaction over Pt\textsubscript{0.01}Cu-SAA catalyst. Reaction condition: 2.5% butane, 5% H\textsubscript{2}, bal. He. 50 mL/min, 0.1 g catalyst, 400 °C.

The carbon deposition on spent Pt NP and PtCu SAA catalysts after long-hour butane dehydrogenation reactions was followed with TPO experiments, as shown in Figure 5.14. Significant CO\textsubscript{2} peaks were observed on Pt NP, revealing the carbon deposition. There are multiple CO\textsubscript{2} peaks in this TPO profile, at least one of them corresponding to the graphitic carbon. Different from Pt NP, PtCu SAA catalyst only shows minimal carbon deposition, which is essentially the same as that on the bare silica support. These results are in agreement with the stability tests shown in Figure 5.13. PtCu SAA catalyst is highly stable in butane dehydrogenation conditions without coke formation, while monometallic Pt catalyzes the coke formation. The two major mechanisms of the deactivation of Pt catalysts in alkane dehydrogenation reactions are carbon deposition and sintering of Pt nanoparticles. From TEM imaging, we found the particle size of Pt slightly increases from 1.8 to 2.1 nm after being used
in the reaction conditions. Thus, the instability of the monometallic Pt catalysts is mainly due to the carbon deposition.

![Graph showing CO₂ concentration over temperature](image)

**Figure 5.14.** Comparing the degree of coking in Pt NP vs PtCu SAA catalysts. Temperature-programmed-oxidation (TPO) of used (a) Pt-NP, (b) Pt₀.₀₁Cu-SAA and (c) silica. (a) and (b) are spent catalysts after butane dehydrogenation reaction. (a), (b) and (c) were treated in a He flow at 300 °C for 1 h before TPO to desorb the hydrocarbon adsorbates. TPO conditions: 10% O₂/N₂, 50 mL/min, 3 °C/min, 100 mg sample.

To investigate the sintering of PtCu SAA NPs in butane dehydrogenation condition, TEM images were taken for fresh Pt₀.₀₁Cu-SAA samples reduced in H₂ and Pt₀.₀₁Cu-SAA used in butane dehydrogenation at 400 °C for 52 hours. The particle size of Pt₀.₀₁Cu-SAA does not
change, as shown in the TEM images and particle size distribution in Figure 5.15. This confirms the stability of PtCu SAA NPs in butane dehydrogenation conditions.

![Figure 5.15](image)

**Figure 5.15.** TEM images and particle size distribution (insets) of (a) fresh Pt$_{0.01}$Cu-SAA and (b) spent Pt$_{0.01}$Cu-SAA after butane dehydrogenation reaction at 400 °C.

Controlling coke formation is one of the most important considerations C-H bond activation catalysts. A well-known example is coke formation during methane steam reforming over nickel which requires costly regeneration processes to be used at industrial scales. In the same vein, the design of commercial dehydrogenation processes is often dominated by the need to control coke formation on the catalyst surface. The most popular dehydrogenation technology for light alkanes is UOP’s Oleflex process which uses a supported Pt on Al$_2$O$_3$ catalyst. Here, a complex and high-cost continuous catalyst regeneration is used to burn off coke from the particles followed by treatment with chlorine gas to redisperse the Pt particles. Similarly, CB&I’s Catofin process uses a series of fixed bed reactors containing a chromium oxide catalyst. These reactors must be regenerated after only a few minutes on stream in part
due to rapid coke deposition. Clearly, the development of coke resistant C-H bond activation catalysts based on single-atom alloys would have important practical implications for these industries.

Figures 5.16 and 5.17 show the FTIR spectra for used Pt$_{0.01}$Cu-SAA and Pt$_{0.03}$Cu-SAA catalysts, where we cannot find any linearly or bridge adsorbed CO on extended Pt surfaces. This confirms that Pt atoms remain isolated after long term B-D scrambling experiments. These results demonstrate the structural stability of PtCu SAA NPs under atmospheric pressure reaction conditions.

**Figure 5.16.** FTIR spectra of chemisorbed CO molecule on the used Pt$_{0.01}$Cu-SAA. The time values on the graph indicate the time of helium gas purge after CO exposure.
Figure 5.17. FTIR spectra of chemisorbed CO molecule on the used Pt$_{0.03}$Cu-SAA catalyst. The time values on the graph indicate the time of helium gas purge after CO exposure.

5.4 Collaboration: surface science and DFT studies on the C-H activation over PtCu SAA catalysts

The surface science studies on C-H activation over PtCu SAA catalysts were performed in the Sykes lab at Tufts. Methyl iodide (CH$_3$I) provides a simple method for adding methyl groups to the alloy surfaces and TPR studies of subsequent methane evolution allows us to examine C-H activation energetics. On Cu(111), methane and the larger hydrocarbons desorb at ~450 K. It has been previously demonstrated that the rate limiting step to form these products is the activation of C-H bonds in methyl groups to produce methylene and H$_2$.\textsuperscript{50,51} On the 0.01ML PtCu(111) SAA surface, methane and carbon coupling products desorb at ~350 K, 100 K cooler than on the pure Cu(111) surface. Because the rate limiting step in methane evolution is C-H
activation, this 100 K temperature shift reveals that single Pt atoms in the Cu surface significantly lower the barrier to C-H activation in CH\textsubscript{3}. The formation of ethene, ethane and propene implies the Pt-Cu SAA maintains the ability of Cu to avoid coking via C-C coupling. On 1ML Pt-Cu(111), the majority of methane desorbs at \textasciitilde250 K, and hydrogen from methylene decomposition desorbs at high temperature.

DFT studies were performed at Stamatakis’s Lab at University College London. DFT calculations find the strong binding of CH\textsubscript{x} intermediates to the Pt(111) surface compared to Cu(111) which explains the tendency for coke to form on Pt and the lack thereof on Cu. Interestingly, the Pt-Cu(111) SAA exhibits intermediate barrier heights compared to the two pure metals. Dehydrogenation of CH\textsubscript{x} species on Pt-Cu(111) SAA and Cu(111) are endothermic, however the net pathway on Pt(111) is exothermic. These findings explain our experimental observations that the Pt-Cu(111) SAA may readily activate C-H bonds in adsorbed CH\textsubscript{3}* while resisting coke formation.

5.5 Summary

The development of coke resistant C-H activation catalysts is an important but difficult task for the field of heterogeneous catalysis. Here we introduced a novel single-atom alloy (SAA) approach in designing Pt- based C-H activation catalysts. In this chapter, C-H activation was examined on PtCu SAAs using supported nanoparticle catalysts in a fixed-bed flow reactor at atmospheric pressure. Our catalytic studies with supported PtCu, Pt, and Cu NPs show that
PtCu-SAAs activate C-H bonds using the B-D scrambling reaction in butane. We find that the PtCu SAAs catalyze the exchange reaction at 250 °C compared to 550 °C on Cu NPs and are stable during the reaction. Pt NPs activate the reaction at temperatures as low as 100 °C, but quickly deactivate due to coking. Moreover, the PtCu SAA catalysts showed high stability and coke resistance in butane dehydrogenation reaction conditions. The PtCu SAA catalysts can stably convert butane to butene at 400 °C for at least 52 hours without any carbon deposition or sintering. The single-atom Pt sites are stable in the reaction condition as indicated by CO-IR and EXAFS results.

We find the C-H activation barrier in butane molecule follows the order of: Pt < PtCu SAA < Cu, which is in agreement with the surface science and DFT results of our collaborators. The apparent activation energy measured for the butane dehydrogenation reaction corroborates this trend. These findings illustrate that PtCu SAAs are promising catalysts for C-H bond activation in small alkane molecules at moderate temperatures and can be further developed as robust catalysts resistant to coking that plagues traditional Pt, Pd and Ni catalysts.

5.6 References


34. Ferri D, Bürgi T, Baiker A. Pt and Pt/Al2O3 Thin Films for Investigation of Catalytic Solid–Liquid Interfaces by ATR-IR Spectroscopy: CO Adsorption, H2-Induced


Chapter 6. PdAu single-atom alloy catalysts for selective hydrogenation of 1-hexyne to 1-hexene

6.1 Introduction

Selective hydrogenations of both double and triple carbon-carbon bonds are important for a wide range of industrial processes. Pd, Pt, Ni, and Ru based systems are commonly employed for hydrogenation of alkenes and alkynes due to their high catalytic activity. While these catalysts are highly active, they may not always be selective to the desired products. Among these metals, Pd is the most selective but typically exhibits a decrease in alkene selectivity with alkyne conversion. Therefore, Pd catalysts are commonly used with promoters to improve alkene selectivity; transition metals such as Ag, Rh, Au, Cu, Zn, Cr, and V have been reported to improve selectivity.\(^1\) Another approach to combat this problem is to design catalysts that contain only the Pd ensembles necessary to perform or to assist the reaction of interest.

Individual, isolated Pd atoms in the Cu surfaces, termed single-atom alloys (SAAs), act as sites for hydrogen adsorption, dissociation and spillover onto the otherwise inert Cu surface. The weak binding of the H atom intermediates offered by Cu(111) led to the selective hydrogenation of both styrene and acetylene, as demonstrated by a combination of scanning tunneling microscopy (STM) and temperature programmed desorption/reaction (TPD/R) experiments under UHV conditions.\(^2\) Recently the SAA strategy was applied to the preparation of PdCu alloy nanoparticles (NPs) which are
active for the hydrogenation of phenylacetylene to styrene at ambient temperature with 94% selectivity at all conversions.\textsuperscript{3} NP catalysts were prepared by depositing small amounts of Pd into the surface of pre-formed Cu NPs by the galvanic replacement (GR) reaction.

Furthermore, over the last few years, several studies of single Pd atoms stabilized in different matrices for selective hydrogenation reactions have been reported.\textsuperscript{4–6} Liu et al. showed single-atom Pd dispersed on ethylene glycolate-stabilized TiO\textsubscript{2} nanosheets are highly active for the hydrogenation of alkenes and aldehydes.\textsuperscript{4} Yan et al. showed that atomically dispersed Pd on graphene is selective for the partial hydrogenation of 1,3-butadiene.\textsuperscript{5} Vilé et al. reported that single Pd atoms anchored into the cavities of graphitic carbon nitride (g-C\textsubscript{3}N\textsubscript{4}) have high activity and selectivity for the partial hydrogenation of 1-hexyne.\textsuperscript{5,7} Yan et al. attributed the high selectivity to a steric effect induced by the high packing density at the single-atom Pd sites compared to the extended Pd surfaces.\textsuperscript{5} Through DFT calculations, Vilé et al. concluded that the weak adsorption of products and the electron deficiency at Pd sites contribute to the high selectivity.\textsuperscript{6}

In this chapter, single Pd atoms were incorporated in gold surfaces, and the resulting Pd\textsubscript{1}Au SAAs were evaluated for the selective hydrogenation of alkynes, specifically the hydrogenation of 1-hexyne to 1-hexene. Bimetallic PdAu materials of various compositions have been studied extensively for a number of reactions, including selective oxidation\textsuperscript{8–11}, selective hydrogenation\textsuperscript{12–19}, the Ullmann reaction\textsuperscript{20}, vinyl acetate synthesis\textsuperscript{21}, formic acid decomposition\textsuperscript{22}, and the direct synthesis of hydrogen peroxide\textsuperscript{23–25}. Generally, it is difficult to
elucidate the roles of Pd and Au in the catalytic cycles and their alloying effects due to the presence of many Pd and Au ensembles on the catalyst surfaces. In a few reports, the effect of preparing isolated Pd or Au atoms in the bimetallic PdAu catalysts has been attributed to providing exceptional activity or selectivity because of the unique chemistry of single-atom sites, which also allows for more conclusive mechanistic studies.\textsuperscript{8,21,26–28} Pioneering work from the Goodman\textsuperscript{21} and Toshima\textsuperscript{29} groups has identified Pd or Au single-atom active sites in vinyl acetate synthesis and glucose oxidation, respectively, on catalytic surfaces with well-defined single Pd or Au atom structures. The Mullins group has demonstrated that the PdAu alloy surfaces catalyse the dehydrogenation of formic acid while the extended Pd surfaces catalyse the dehydration of formic acid.\textsuperscript{22} Through careful bimetallic nanoparticle synthesis with modified impregnation methodology, Luo et al. prepared Au-Pd random alloy nanoparticles with homogeneous structures and demonstrated the enhancement of hydrogenation activity by the alloying effect.\textsuperscript{30}

The selective hydrogenation reactions of alkynes to alkenes are important processes for the production of high purity alkene feedstocks for the polymer industry. Pd is widely used for hydrogenation due to its low hydrogenation activation barrier.\textsuperscript{1} While Au NP catalysts are highly selective for partial hydrogenation of alkynes, they are rarely used for these reactions due to the low intrinsic activity of Au.\textsuperscript{31} Some studies have used Au as an additive to improve the selectivity of Pd catalysts for partial hydrogenation reactions, while Pd is still the bulk catalyst phase.\textsuperscript{12,13,15,17,18,32} Recent work by the Louis and Zhang groups has evaluated various Pd ensembles from clusters to single atoms and suggested that the selectivity increases with the dispersion of Pd in PdAu alloys.\textsuperscript{33,34} However, these reports fall short of elucidating the
specific roles of Pd and Au in selective hydrogenation of alkadienes\textsuperscript{33} and alkynes\textsuperscript{34} and have only investigated the hydrogenation reactions in the gas phase.

In this work, we demonstrate that PdAu SAAs are both active and highly selective for the partial hydrogenation of 1-hexyne in the liquid phase, and attribute these properties to the unique structure of PdAu SAAs.

6.2 Experimental methods

6.2.1 Catalyst Synthesis

PdAu-SAA NPs were prepared using sequential reduction methods as reported in Tedsree et al. with some modifications\textsuperscript{35}. First, Au NPs were prepared as follows: 0.3 g HAuCl\textsubscript{4}·3H\textsubscript{2}O and 1.2 g poly(vinylpyrrolidinone) (PVP, MW = 58,000) were mixed in 50 mL ethylene glycol (EG) in a round bottom flask with vigorous stirring. The solution was purged with N\textsubscript{2} for one hour before 0.3 g NaHCO\textsubscript{3} was added. The solution was then heated to 90 °C at 5 °C/min with N\textsubscript{2} flowing through the round bottom flask. The solution was held at 90 °C for half an hour before cooling to ambient temperature. The Au NPs were thus obtained. To prepare PdAu-SAA NPs, the desired amount of Pd(NO\textsubscript{3})\textsubscript{2}·xH\textsubscript{2}O was dissolved in 1 mL EG and added to the round bottom flask containing the EG-dispersed Au NPs. Under N\textsubscript{2} purge and vigorous stirring, the solution was heated to 90 °C at 5 °C/min and held for 8 h. After the solution was cooled to ambient temperature, 300 mL of acetone was added in order to precipitate the NPs. The acetone-EG solution was centrifuged at 3000 rpm for 10 minutes to separate the NPs. The retrieved NPs were then washed with ethanol and hexane thrice and dried in vacuum overnight. To obtain silica supported Au and PdAu NPs, fumed silica (heat treated in air at
650 °C for 3 h) was introduced into the NP gold and PdAu-SAA solutions in water and stirred overnight. The slurry carrying the supported NPs was centrifuged and dried in vacuum for 24 h then calcined in air at 300 °C. The unsupported Pd-NPs were prepared by the same method with Pd(NO₃)₂ precursor.

Pd/C is provided by Sigma-Aldrich (St. Louis, MO). It is 10 wt% Pd on activated carbon support (surface area: 933 m²/g, pore size: 150-200 Å, pore volume 0.92 ml/g). Pyridine-IR shows peaks at 1588, 1575, 1446 and 1438 cm⁻¹, which suggests Lewis acid sites³⁶,³⁷. All peaks disappeared after degassing at room temperature for 60 minutes. The Lewis acid sites are relatively weak.

6.2.2 Catalyst characterization

The NPs were digested with concentrated HCl and H₂O₂ to prepare the samples for inductively coupled plasma atomic emission spectroscopy (ICP-AES) with a Leeman Labs PS1000 instrument. High-resolution transmission electron microscopy (HR-TEM) was conducted on a JEOL 2010 electron microscope with 200 kV and 107 μA beam emission. The samples were obtained by drop casting of the ethanol solution of NPs on carbon film over a copper microgrid.

X-ray photoelectron spectra (XPS) were obtained on a Thermo Scientific K-Alpha system equipped with an Al source and a double focusing hemispherical analyser with a 128-channel detector at a pass energy of 50 eV.

X-ray powder diffraction (XRD) tests were performed on a PANalytical X’Pert Pro instrument. Cu Kα radiation was used as the X-ray source. Data was collected for 2θ between 30° and 70°.
Infrared (IR) spectroscopy was performed in both attenuated total reflection (ATR) and diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) modes on a Thermo Nicolet iS50 FT-IR.

6.2.3 H$_2$-D$_2$ exchange

To measure the hydrogen activation activity of the catalysts, H$_2$-D$_2$ exchange reaction was conducted in a packed-bed flow microreactor (L=22inch, O.D.=1/2 inch). The samples were treated in 20% H$_2$ at 150 °C for 5 minutes prior to the H$_2$-D$_2$ exchange experiments. A gas mixture of 10% H$_2$, 10% D$_2$ and bal. Ar was introduced to the reactor at a total flow rate of 50 mL/min at ambient temperature. After flowing the gas mixture for 15 minutes, the temperature was raised to desired temperature at 5 °C/min. The gas composition was analysed by a mass-spectrometer.

6.2.4 Partial hydrogenation of 1-hexyne

A stainless-steel Parr reactor was used for the hydrogenation of 1-hexyne. The desired amount of sample was added to 20-80 mL ethanol. The mixture was stirred for half an hour and sonicated for 15 minutes before being loaded into the reactor. Pure hydrogen was bubbled through the solution while stirring for 1 h. 1-hexyne was injected into the solution to reach 1% vol/vol concentration in ethanol. The reactor was then pressurized with pure hydrogen to the desired pressure (5 bar) under stirring at 600 rpm. Liquid-phase specimens were taken from the reactor at different time points.
6.3 Results and discussion

**NP preparation and characterization**

**Table 6.1. NP sample properties**

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Composition</th>
<th>Pd species</th>
</tr>
</thead>
<tbody>
<tr>
<td>PdAu-SAA</td>
<td>0.4 at.% Pd, 99.6 at.% Au</td>
<td>Single atoms</td>
</tr>
<tr>
<td>PdAu-SAA/SiO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>0.008 wt. % Pd, 3.8 wt. % Au / SiO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Single atoms</td>
</tr>
<tr>
<td>Pd/C</td>
<td>10 wt. % Pd / C</td>
<td>NPs*</td>
</tr>
<tr>
<td>Au/SiO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>3.8 wt. % Au / SiO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>--</td>
</tr>
</tbody>
</table>

* Dispersion of Pd is 26%. (Determined by CO chemisorption)

To study the catalytic performance of PdAu SAAs in ambient pressure reaction conditions, we prepared PdAu SAA NPs and other samples, as listed in table 6.1. We followed the sequential reduction (SR) method in ethylene glycol to prepare the PdAu SAA NPs. As reported in the literature, the SR methods are effective in forming core-shell nanoparticles. In order to prepare PdAu SAAs, a small amount of Pd species was added onto the pre-formed Au NPs. The Pd to Au atomic ratio was 1/250 as determined by ICP-AES. Pd and Au metals are miscible and their negative mixing enthalpy drives the Pd atoms to disperse well in the surface of the Au NPs. Therefore, at low Pd loadings,
Au NPs with isolated Pd atoms can be formed. The PdAu-SAA NPs were prepared in unsupported form, then a portion of the sample was deposited on silica followed by calcination. The silica-supported samples are denoted as PdAu-SAA/SiO$_2$.

The ethanol solution of PdAu-SAA NPs exhibited the typical deep red colour of Au NPs (Figure 6.1a). Figures 6.1 (b, d) show TEM micrographs of the as-synthesized PdAu-SAA NPs. The NPs have well defined shapes indicating crystalline structures. Figure 1c shows the particle size distribution with an average particle size of 7.6±1.9 nm. In the zoom-in TEM image (Figure 6.1d), Au(111) lattice spacing of 0.23nm was found. These results confirm the formation of Au NPs and the dilute distribution of Pd atoms within them.

![Figure 6.1](image_url)

**Figure 6.1.** (a) Photograph of PdAu-SAA NP solution. (b, d) TEM images and (c) particle size distribution of PdAu-SAA NPs.

The XRD results (Figure 6.2) corroborate the TEM observations. Peaks at 2θ of 38, 45 and 65 degrees indicate only Au diffraction patterns, in agreement with the TEM results. The average crystallite size is 7.6 nm as determined by the band broadening of
the Au(111) and (200) peaks which is very close to the NP diameter found in the TEM measurements.

![XRD spectrum](image)

**Figure 6.2.** Powder XRD of PdAu-SAA/SiO₂.

We performed EDS elemental mapping on PdAu SAAs to characterize the distribution of Pd and Au in the catalysts. As shown in Figure 6.3, EDS elemental mapping over a large area of the sample does not show any monometallic Pd nanoparticles. Pd is in the Au area in all cases. This demonstrates that truly bimetallic PdAu NPs are formed.
Figure 6.3. Dark field STEM image and Au and Pd EDS maps of PdAu-SAA.

XRD analysis of samples of different Pd/Au ratios were performed. As shown in Figure 6.4, the peaks of PdAu₃ and Au phases are very close. We did not observe any AuPd₃ or Pd phases in these samples. XRD shows that there are no monometallic Pd nanoparticles, which confirms the effectiveness to form bimetallic nanoparticles by this preparation method. However, XRD alone cannot prove the formation of PdAu alloy phases due to the very close 2θ of PdAu₃ and Au peaks.
EDS and XRD confirm the formation of PdAu bimetallic NPs, which also demonstrate the effectiveness of the SR method in preparation of bimetallic NPs. SR method has been applied in the preparation of core-shell structure NPs\textsuperscript{38,40}, this is the first time it is applied in the preparation of PdAu SAA NPs.

In order to probe the atomic geometry of the Pd ensembles in the surface of the Au NPs we performed CO-IR experiments in the ATR-IR cell with CO in the gas phase.\textsuperscript{41,42} Figure 6.5 shows the IR spectra of the carbonyl region for Pd-NP, PdAu-SAA and Au-NP samples. Atop CO on Au and Pd sites was observed at 2099 and 2015 cm\(^{-1}\), respectively, while the stretch frequency at 1935-1920 cm\(^{-1}\) corresponds to bridged CO on the extended Pd surfaces.\textsuperscript{33,43,44} However, no bridged CO was detected on PdAu-SAAs, which confirms that the Pd atoms are isolated from one another in the surface of the gold NPs. Only
linearly adsorbed CO (atop) on the Au and Pd atoms was found on the SAA samples.\textsuperscript{33,45} These results provide strong evidence that the Pd atoms are fully isolated in the surface of PdAu-SAA NPs.

**Figure 6.5.** ATR-IR spectra of the CO stretch on Pd-NP, Au-NP and PdAu-SAA samples. Spectra were collected at 30 °C with pure CO flowing in the gas phase for Au-NP and PdAu-SAA. The spectra for Pd-NP were collected in H\textsubscript{2}/He (20%) gas after CO adsorption.

**Catalytic reaction studies**

The catalytic performance of PdAu-SAA catalysts for the partial hydrogenation of 1-hexyne was studied in a batch reactor.
Figure 6.6. Leaching test: 1-hexene/1-hexyne ratio as a function of time. The liquid phase was separated from the reaction mixture after 6-hour reaction. Fresh 1-hexyne was injected and the typical hydrogenation reaction conditions used in this work were applied. (5 bar H₂, 600 rpm, 25 °C) The 1-hexene/1-hexyne ratio is plotted here. No conversion of 1-hexyne was observed.

It is clear from Figure 6.6 that the liquid phase of the reaction mixture is not active for hydrogenation of 1-hexyne. In this experiment, the catalyst was filtered out of the solution after 6-hour reaction in the batch reactor. 1-hexyne was re-introduced into the solution but was not converted under the hydrogenation reaction condition, as indicated by the constant 1-hexene/1-hexyne ratio over the course of 60 minutes. Therefore, the PdAu-SAA system is a truly heterogeneous hydrogenation reaction catalyst.
Figure 6.7 shows time-resolved batch reactor data for PdAu-SAA/SiO₂ and Pd/C catalysts at ambient temperature. PdAu-SAA is highly selective in the hydrogenation of 1-hexyne to 1-hexene as the product distribution consists over 85% hexenes (96% 1-hexene and 4% 2-hexene isomers) at full conversion of 1-hexyne. Isomers of 1-hexenes were formed, but less than 4% over the course of the hydrogenation reaction. Similar observations that a small amount of 2-hexene isomers can be formed in the hydrogenation of 1-hexyne have been reported on various supported Pd catalysts.46–48 By comparison, the monometallic Pd catalysts (10 wt. % Pd on carbon) have less than 10% selectivity to hexenes because they convert most of the 1-hexyne to n-hexane at 100% conversion. This is in agreement with previous reports that monometallic Pd catalysts over-hydrogenate alkynes to undesired alkanes, while single Pd atom catalysts have high selectivity for partial hydrogenation reactions.5,33

Figure 6.7. Time-resolved batch reactor data for the hydrogenation of 1-hexyne over (a) 10 wt% Pd/C catalysts and (b) PdAu-SAA/SiO₂ catalysts. (c) selectivity to 1-hexene as a function of
conversion. Reaction condition: 1% hexyne-ethanol solution, 5 bar H₂, 600 rpm, 25 °C. Same 1-hexyne/Pd ratio was used for hydrogenation reaction with 10wt% Pd/C and PdAu-SAA/SiO₂.

We have demonstrated in prior work that PdCu and PtCu SAAs are highly selective for the partial hydrogenation of alkynes and dienes. A bifunctional mechanism was verified for those catalysts in which hydrogen is activated at the single Pt atom sites and hydrogen atoms spill over to the copper surface where they are weakly bound and participate in the selective hydrogenation reactions. Surface science investigations have found that single Pd atoms facilitate molecular hydrogen dissociation on the otherwise inert Au(111) surface, however, hydrogen spillover from Pd to Au is unfavourable due to weak H-Au binding.

The high selectivity of Au catalysts was preserved in PdAu SAA NPs for the partial hydrogenation of alkynes. We propose this is due to the weak adsorption of alkenes on PdAu SAAs as metallic Au is well-known to bind hydrocarbon molecules weakly. However, Pd surfaces strongly bind alkenes and their desorption barrier is higher than their hydrogenation reaction barrier which leads to low selectivity for partial hydrogenation. Alloying Pd with Au leads to modification of the electronic structure of Pd atoms: the s and p electron density moves toward Au while d electron density moves toward Pd. So the d band center is shifted away from the Fermi level and the adsorption strength of adsorbates on Pd is weakened. These electronic effects,
coupled with the ensemble effects that no extended Pd ensembles are present in the surface give exclusively weak adsorption sites.

The reaction rate per surface Pd atom on PdAu-SAA/SiO$_2$ is lower than that of the monometallic Pd catalysts (Figure 6.8). This might be due to the ligand effects by alloying with Au.

**Figure 6.8.** Initial reaction rate of hydrogenation of 1-hexyne normalized by amount of surface Pd. (Batch reactor, 1% 1-hexyne-ethanol solution, 5 Bar H$_2$, 600 rpm, 25 °C)

In batch reactor studies, a reaction rate of $3 \times 10^4$ mol$_{1\text{-hexene}}$/mol$_{Pd}$/h was measured at 70 °C and 5 bar H$_2$ with PdAu-SAA/SiO$_2$, while the high selectivity to 1-hexene was maintained (Figure 6.9). Vilé et al. tested Pd/mpg-C$_3$N$_4$ catalyst under the
same temperature and H$_2$ pressure in a continuous-flow three-phase reactor and found lower normalized reaction rate ($\sim 1.5 \times 10^3$ mol$_{1\text{-hexene}}$/mol$_{\text{Pd}}$/h) but similar selectivity.$^{56}$

**Figure 6.9.** Time-resolved batch reactor data of partial hydrogenation of 1-hexyne to hexene over PdAu-SAA/SiO$_2$ catalysts. 1% hexyne-ethanol solution, 5 bar H$_2$, 600 rpm, 70 oC.

**Stability test**

The activity of the PdAu-SAA/SiO$_2$ catalysts is stable at the reaction conditions applied. As shown in Figure 6.10, no decrease of the hydrogenation rate was observed during 3 cycles of 1-hexyne hydrogenation reactions. The high stability of SAA catalysts has been demonstrated in gas phase hydrogenation and dehydrogenation.
However, this is the first report showing good stability of SAA NPs in liquid phase hydrogenation reactions.

**Figure 6.10.** Stability test for PdAu-SAA/SiO$_2$ catalyst. The reaction rate of hydrogenation of 1-hexyne (bars) and selectivity to 1-hexene at 50% conversion (squares) was compared between the first, second and third cycles. (Batch reactor, 1% Hexyne-ethanol solution, 5 bar H$_2$, 600 rpm, 25 °C. ~3h each cycle)

**Characterization of used catalysts**
We characterized the spent catalysts to examine the stability of the single-atom alloy structure. TEM images of used PdAu-SAA NPs show crystalline samples with an average particle size of 8.3 ± 2.0 nm with narrow size distribution (Figure 6.12). The minor increase of the average gold particle size in the used catalyst indicates the PdAu-SAA NPs are stable in the reaction conditions with respect to sintering.

XRD of the used PdAu-SAA/SiO₂ catalysts shows diffraction peaks at 38, 44 and 65 degrees that correspond to metallic Au (Figure 6.13a). The peak intensity of Au(111) is same as that of fresh catalysts, no peak shift was observed compared to the fresh catalysts. Moreover, ICP-AES of the spent catalysts found no change in the elemental catalyst composition.

**Figure 6.11.** (a, c) TEM image and (b) particle size distribution of used PdAu-SAA NPs.

CO-IR for the spent PdAu-SAA NPs shows exclusively atop CO on Pd and Au sites but no bridging CO at lower wavenumbers, which confirms the stability of the single Pd atom sites (Figure 6.12c). XPS of spent PdAu-SAA exhibits an Au 4f7/2 peak at 83.4 eV.
(binding energy) corresponding to metallic Au (Figure 6.12b). This is in agreement with the reported values of Au 4f peaks in bimetallic PdAu NPs. There is indeed a slight negative shift of the binding energy compared to the 4f$_{7/2}$ of metallic Au (84 eV). This may be due to the charge transfer between Pd and Au. We were not able to obtain definitive XPS spectra for Pd as the Pd 3d peak overlaps with Au 4d$_{5/2}$ and Pd 3p intensity is lower than the detection limit of the instrument. Overall, the characterization of the spent catalysts demonstrates the structural and compositional stability of the PdAu-SAA NPs in hydrogenation reaction conditions, which explains the stability observed in the reaction tests.
Figure 6.12. Characterization of used PdAu-SAA/SiO$_2$ catalysts. (a) XRD, (b) XPS and (c) CO-DRIFTS. The tests were performed on spent catalysts without further treatment at ambient temperature.
To investigate the surface composition of PdAu catalysts, we performed XPS on a higher Pd loading sample, which is Pd$_{0.2}$Au. In this sample the bulk Pd to Au ratio is 0.2. As shown in Figure 6.14, both Au and Pd are in the metallic states. The surface Pd/Au ratio calculated with XPS data is 3/2, which indicates the Pd segregates to the surface of PdAu bimetallic NPs.

**FTIR studies**

To investigate the 1-hexyne hydrogenation on PdAu-SAA catalysts, FTIR spectroscopy was performed with DRIFTS and ATR reaction cells. As shown in Figure 6.14, DRIFTS spectra were collected following a dose of 1-hexyne (2 μL) in the pure hydrogen gas stream. The peaks at 2966, 2941 and 2877-2865 cm$^{-1}$ are assigned to v(C-H) in CH$_2$ and CH$_3$ groups. They are very
similar to the same features in the liquid 1-hexyne spectra (Figure 6.16). This shows there is no significant perturbation in the saturated part of the 1-hexyne molecules upon adsorption on the catalysts. The peaks between 3335 and 3296 cm\(^{-1}\) are assigned to \(\nu(\equiv\text{C-H})\). The peaks are broadening after adsorption compared to the liquid 1-hexyne, similar effects were observed on adsorbed phenylacetylene on Pt catalysts before.\(^{61}\) The peak at 2112 cm\(^{-1}\) is assigned to the \(\nu(\equiv\text{C})\), while the \(\nu(\equiv\text{C})\) of liquid 1-hexyne is at 2119 cm\(^{-1}\).

**Figure 6.14.** DRIFTS spectra recorded when 1-hexyne and \(\text{H}_2\) are carried over PdAu-SAA/SiO\(_2\) catalyst.
A peak at 3081 cm$^{-1}$ appears after a short period of time following 1-hexyne dose (Figure 6.15). This peak is assigned to $\nu(=\text{C-H})$. We find a similar $\nu(=\text{C-H})$ peak when 1-hexene and helium are carried over PdAu-SAA/SiO$_2$ catalyst. However, we do not see this peak when 1-hexyne and $\text{H}_2$ are carried over silica support. This is a direct evidence of the conversion of 1-hexyne to 1-hexene at room temperature. The 1-hexene formed on the surface of PdAu NPs has a certain residence time and can be desorbed and re-adsorbed on the PdAu NPs or on the silica support. The $\nu(=\text{C-H})$ of all these species of 1-hexene can be detected.

**Figure 6.15.** DRIFTS spectra between 3130 and 3120 cm$^{-1}$ recorded when 1-hexyne and $\text{H}_2$ are carried over PdAu-SAA/SiO$_2$ catalyst.
As shown in Figure 6.17, we observed a peak centered at around 3580 cm\(^{-1}\) and an adjacent negative peak at higher wavenumbers when carrying 1-hexyne and H\(_2\) over PdAu-SAA/SiO\(_2\) catalyst. The 3583 cm\(^{-1}\) peak is assigned to a 1-hexyne species π-adsorbed on the OH group of silica\(^{62}\). The negative peak has been reported previously, which is likely due to the perturbation of the silanol group on silica.
Figure 6.17. DRIFTS spectra between 3800 and 3450 cm$^{-1}$ recorded when 1-hexyne and H$_2$ are carried over PdAu-SAA/SiO$_2$ catalyst.

The DRIFTS spectra recorded when carry 1-hexyne and H$_2$ over silica support are shown in Figure 6.18 and 6.19. The peaks at 2966, 2941, 2879 and 2870 cm$^{-1}$ are assigned to $\nu$(C-H). They are at very similar positions as that of PdAu/SiO$_2$. A peak at 2115 cm$^{-1}$ is assigned to $\nu$(C≡C), which is slightly higher (3 cm$^{-1}$) than that of PdAu/SiO$_2$. We observed a broad absorption band centered at $\sim$ 3335 – 3316 cm$^{-1}$ ascribed to $\nu$(C≡C-H) as shown in Figure 6.19B. At the meantime, multiple peaks are shown for the PdAu/SiO$_2$ sample (Figure 6.14) at 3335, 3323, 3296 cm$^{-1}$ and a shoulder band at around 3260 cm$^{-1}$ in the $\nu$(C≡C-H) region. The differences between them is very likely due to the adsorption on PdAu NPs. We can also see a broad peak around 3700-3500 cm$^{-1}$ corresponding to 1-hexyne species π-adsorbed on the OH group of
This confirms the adsorption of 1-hexyne on silica support. We flow 1-hexyne and H\textsubscript{2} over silica support but did not observe any evidence to show the formation of 1-hexene.

**Figure 6.18.** DRIFTS spectra between 3450 and 2000 cm\textsuperscript{-1} recorded when 1-hexyne and H\textsubscript{2} are carried over silica.
The catalytic performance studies were performed in a batch reactor with liquid phase 1-hexyne and above ambient pressure of H₂. To better mimic the hydrogenation reaction conditions, we used an ATR reaction cell with liquid phase 1-hexyne and 120 psi H₂ applied to the PdAu-SAA catalysts without support.

As shown in Figure 6.20, 6.22 and 6.23, the IR spectra were recorded over the course of the hydrogenation reactions. In the spectra of H₂-hexyne over Pd₀.₀⁰₄Au-SAA system, the v(≡C-H) is at 3306 cm⁻¹, the v(C-H) are at 2960, 2934, 2902, 2873 and 2864 cm⁻¹, while the v(C≡C) is centered at 2117 cm⁻¹. The peak positions are very similar to that in the DRIFTS studies. The v(C≡C) band is broadened compared to the liquid 1-hexyne (Figure 6.16), which is in agreement with that has been reported in the literature. The peak intensity of v(C≡C) and v(≡C-H) decrease over the time while the peak intensity of v(C-H) essentially unchanged, this captures the
conversion of 1-hexyne in the hydrogenation conditions. The peak at 3080 cm\(^{-1}\) is ascribed to \(v(=C-H)\), which appears under the reaction condition (Figure 6.21), showing the formation of hexene molecules.

**Figure 6.20.** ATR-IR collected with 1-hexyne and 120 psi H\(_2\) applied on Pd\(_{0.004}\)Au-SAA catalysts in the batch mode at room temperature.

**Figure 6.21.** ATR-IR collected with 1-hexyne and 120 psi H\(_2\) applied on Pd\(_{0.004}\)Au-SAA catalysts in the batch mode at room temperature.
We also conducted ATR-IR with a Pd$_{0.01}$Au sample, which was prepared with the same method as Pd$_{0.004}$Au -SAA but with higher Pd loadings. CO-FTIR finds no bridge CO adsorbed on this sample. The data are shown in Figure 6.22. The broad band centered at 3306 cm$^{-1}$ is assigned to $\nu$(≡C-H), the peaks at 2960, 2933, 2902, 2873, 2862 cm$^{-1}$ are assigned to $\nu$(C-H) modes and the peak at 2115 cm$^{-1}$ is ascribed to $\nu$(C≡C).

Similarly, we observed the evolution of $\nu$(≡C-H), $\nu$(C-H) and $\nu$(C≡C) over Pd$_{0.2}$Au sample (Figure 6.23), which is an even higher Pd loading sample that CO-FTIR shows bridge CO.

![Figure 6.22](image-url)

**Figure 6.22.** ATR-IR collected with 1-hexyne and 120 psi H$_2$ applied on Pd$_{0.01}$Au catalysts in the batch mode at room temperature.
**Figure 6.23.** ATR-IR collected with 1-hexyne and 120 psi H$_2$ applied on Pd$_{0.2}$Au catalysts in the batch mode at room temperature.

The ATR-IR results are in good agreement with the DRIFTS data for 1-hexyne hydrogenation reactions over PdAu catalysts. ATR-IR mimics the batch reactor conditions while DRIFTS is in flow mode. The PdAu SAA catalysts were thoroughly characterized with both techniques. We demonstrated the conversion of 1-hexyne to hexene *in situ*. 1-hexyne adsorbed on the catalysts mainly through the non-dissociative fashion as we did not detect any dissociative adsorbed species through the conversion of 1-hexyne. 1-hexyne can be adsorbed on the silica support but cannot be hydrogenated by the silica.

To quantify the adsorption of 1-hexyne on the PdAu-SAA catalysts, we performed TGA of the catalyst pellets in nitrogen with different 1-hexyne concentrations, as shown in Figure 6.24. It corroborates the FTIR results that 1-hexyne adsorbed on both silica support and PdAu-SAA NPs. At 60 °C, the adsorption of 1-hexyne is much less than at 30 °C.
Figure 6.24. 1-hexyne adsorption isotherm at 30 and 60 °C.

Kinetics Studies

Comparing PdAu-SAA NP and Au NP catalysts, we observed a remarkable increase in hydrogenation activity of PdAu-SAA. Figure 6.25 compares the initial reaction rate normalized by the total Au loading between the two catalysts. It shows the reactivity is improved tenfold with the addition of a small amount of Pd (0.4 at.%) to the Au NPs. The rate-limiting step of hydrogenation on Au NPs at ambient temperature is hydrogen activation, and the single-atom Pd sites efficiently catalyse hydrogen dissociation as shown in Figure 6.25B. To confirm this in the PdAu NPs, we performed H$_2$-D$_2$ exchange experiments on PdAu-SAA/SiO$_2$ and Au/SiO$_2$, where HD production is a reporter of the H$_2$ activation capability of the catalysts. Figure 6.25B shows the temperature-programmed HD production for both catalysts. At the same Au loading, PdAu-SAA/SiO$_2$ shows much higher HD production rate than Au/SiO$_2$. This indicates that single
Pd atoms in the surface of Au NPs effectively catalyze the H₂ activation. The reaction rate per surface Pd atom on PdAu-SAA/SiO₂ is lower than that of the monometallic Pd catalysts.

![Graph showing reaction rates](image)

**Figure 6.25.** Hydrogenation of 1-hexyne. (A) Reaction rate of PdAu-SAA/SiO₂ and Au/SiO₂.

Reaction condition: 1% hexyne-ethanol solution, 5 bar H₂, 600 rpm, 25 °C. (B) H₂-D₂ exchange TPSR of PdAu-SAA/SiO₂ and Au/SiO₂ catalysts. H₂-D₂ exchange: 10% H₂, 10% D₂, balance Argon, 50 mL/min, 5 °C/min.

Comparing the initial rate over Pd-NP and PdAu-SAA catalysts in Figure 6.26, we found the rate per surface Pd atom was lower on PdAu-SAA. However, the selectivity PdAu-SAA to partial hydrogenation products is significantly higher than that of Pd/SiO₂. The decreased reactivity is likely due to the ligand effects through alloying with Au and the limited availability of adsorbed hydrogen on PdAu-SAA compared to that of Pd-NP.
Figure 6.26. 1-hexyne hydrogenation reaction rate and the selectivity to 1-hexene at 100% conversion of 1-hexyne with Pd-NP (5% Pd/SiO₂) and PdAu-SAA/SiO₂ catalysts. Reaction condition: 1% hexyne-ethanol solution, 5 bar H₂, 600 rpm, 25 °C.

In Figure 6.27, we compare the hydrogenation rate over Pd/SiO₂ and PdAu-SAA/SiO₂ in an Arrhenius-type plot. The PdAu-SAA shows lower activity per Pd atom than monometallic Pd. The apparent activation energy of monometallic Pd is around 26.7 kJ/mol, while the apparent activation energy of PdAu-SAA is around 41.2 kJ/mol. The difference in apparent activation energy indicates different reaction pathway or rate limiting steps. Considering monometallic Pd catalysts, hydrogenation activation is a facile process, while on the monometallic Au catalysts, the rate limiting step is hydrogenation activation. PdAu-SAA is a novel class of hydrogenation catalysts and its rate limiting step has not yet been determined.
Figure 6.27. Arrhenius type plot of 1-hexyne selective hydrogenation over 5% Pd/SiO$_2$ and PdAu-SAA/SiO$_2$ catalysts. Reaction conditions: 1% hexyne-ethanol solution, 5 bar H$_2$, 600 rpm, 6-70 °C.

To study the rate determining step of hydrogenation over PdAu-SAA catalysts, we performed kinetics isotope effect (KIE) measurements, as shown in Figure 6.28. In this study, either H$_2$ or D$_2$ was used in the selective hydrogenation of 1-hexyne at the same experimental conditions. We observed significant KIE with $R_{H2}/R_{D2} = 1.87$. This indicates the hydrogen is involved in the rate limiting step in hydrogenation reactions over these catalysts.
Figure 6.28. KIE measurements. The conversion vs. reaction time for 1-hexyne hydrogenation with PdAu-SAA/SiO$_2$ catalysts. The initial rate is calculated from the slope of the conversion-time curve. Reaction condition: 1% hexyne-ethanol solution, 5 bar H$_2$ or D$_2$, 600 rpm, 25 °C.

To further understand the kinetics of 1-hexyne hydrogenation over PdAu-SAA catalysts, we performed reaction order measurements, as shown in Figure 6.29. The reaction rate of 1-hexyne hydrogenation is linearly correlated with the H$_2$ pressure (Figure 6.29 A), which reveals first order kinetics to H$_2$. While the reaction order to 1-hexyne appears to be zero as the reaction rate is independent of the concentration of 1-hexyne between 0.5 and 1.5% (Figure 6.29 B). Combined with the results of KIE studies, we conclude the rate limiting step in 1-hexyne hydrogenation with PdAu-SAA catalyst is hydrogen activation. These results also corroborate
the surface science TPD results that 1-hexyne adsorbs on the PdAu surface much stronger than hydrogen atoms.

**Figure 6.29.** Reaction order to H$_2$ and 1-hexyne. (A) The reaction rate was measured with 1% 1-hexyne and 3-7.5 bar H$_2$, 600 rpm, 25 °C. (B) The reaction rate was measured with 0.5-1.5% 1-hexyne, 5 bar H$_2$, 600 rpm, 25 °C.

We investigated the H$_2$-D$_2$ exchange with PdAu SAA catalysts. H$_2$-D$_2$ exchange includes two steps, H$_2$ dissociative adsorption and desorption. Gellman et al. devised a mathematical model to describe the H$_2$-D$_2$ exchange on Pd-Cu alloys and fitted their experimental data to the model to calculate the energy barriers. When the H$_2$ partial pressure is low and conversion of H$_2$ and D$_2$ is low, the HD concentration in the outlet is approximately proportional to exp(-E$_{ads}$/RT). Thus, the apparent activation energy of H$_2$-D$_2$ exchange to produce HD can reflect the H$_2$ activation barrier. As shown in Figure 6.30, the results of steady state H$_2$-D$_2$ exchange are put in an Arrhenius-type plot. The apparent activation energy calculated from it is 41.9 kJ/mol,
which is essentially the same as that of apparent activation energy of 1-hexyne hydrogenation over PdAu-SAA catalysts. Thus, the rate limiting step in 1-hexyne hydrogenation is the hydrogenation activation.

Figure 6.30. Arrhenius-type plot of H₂-D₂ exchange over PdAu-SAA/SiO₂ catalysts. The RGA intensity of HD is proportional to the concentration of HD. Reaction condition: H₂-D₂ exchange: 10% H₂, 10% D₂, balance Argon, 50 mL/min, 170-110 °C.

In selective hydrogenation processes, alkyne and alkene hydrogenation are two competing processes. Pd is active for both reactions, but the favorable adsorption of alkyne over alkene leads to selective hydrogenation of alkyne to alkene especially at low
conversion. But with the increase of alkyne conversion, the alkyne surface coverage is reduced, and the selectivity to alkene is decreased. On the Cu catalysts, the alkene desorption barrier is lower than on Pd and the alkene hydrogenation barrier is higher than on Pd, which results in the desorption of reactively formed alkene instead of over-hydrogenation. Thus, Cu is intrinsically more selective than Pd in hydrogenation reactions. For Au NP catalysts, the low-coordination sites are believed to be the active sites, since the hydrogen activation barrier on Au terraces is too high. The favorable adsorption of alkynes compared to alkenes on the low-coordination Au sites determines the high selectivity of Au catalysts in selective hydrogenation. When single-atom Pd is added to the Au surface, the low-coordination Au sites are no-longer necessary for hydrogenation activation. The hydrogenation chemistry of single Pd atom and the surrounding Au atoms is of great interest in understanding the selective hydrogenation with PdAu-SAA catalysts.

Besides the kinetics studies reported in this thesis, surface science techniques and DFT calculations are employed by our collaborators to investigate the PdAu SAA system for selective hydrogenation of 1-hexyne. DFT calculations indicate the most stable adsorption site of 1-hexyne molecules is the bridged Pd-Au site, while the 1-hexene molecule favorably adsorbs in the atop configuration on the single-atom Pd sites. However, on the Pd surface, 1-hexyne adsorbs at the 3-fold hollow sites and 1-hexene adsorbs atop on one Pd atom. The adsorption sites for 1-hexyne on PdAu-SAA and Pd(111) are different. Moreover, the adsorption strength of 1-hexyne and 1-hexene was calculated with DFT and from TPD results. The values calculated with the two methods are in good agreement with each other. We find the addition of single-atom Pd stabilizes the adsorption of both 1-hexyne and 1-hexene, while the adsorption
strengths of 1-hexyne and 1-hexene are comparable. This indicates the favorable adsorption of 1-hexyne over 1-hexene no longer holds for PdAu-SAA catalysts. In other words, 1-hexyne hydrogenation is not thermodynamically favorable compared to 1-hexene hydrogenation on PdAu-SAA catalysts.

![Figure 6.31](image)

**Figure 6.31.** Reaction rate ratio of 1-hexyne hydrogenation over 1-hexene hydrogenation of PdAu-SAA/SiO₂ and Pd/SiO₂ catalysts. Reaction conditions: 1% hexane - or hexene-ethanol solution, 5 bar H₂, 600 rpm, 25 °C.
To better understand the hydrogenation selectivity of PdAu-SAA catalysts, we compared the reaction rate of 1-hexyne hydrogenation with that of 1-hexene hydrogenation. As shown in Figure 6.31, the 1-hexyne hydrogenation rate is one-order-of-magnitude higher than that of 1-hexene over PdAu-SAA catalysts, which reveals a highly selective hydrogenation kinetics. However, the difference between 1-hexyne and 1-hexene hydrogenation rate over Pd-NP catalysts is less than that. Indeed, the apparent activation energy of 1-hexene hydrogenation is higher than that of 1-hexyne over PdAu-SAA catalysts, as shown in Figure 6.32.

**Figure 6.32.** Arrhenius-type plot of 1-hexene hydrogenation over PdAu-SAA/SiO₂ catalysts.

Reaction condition: 1% hexene-ethanol solution, 5 bar H₂, 600 rpm.
6.4 Summary

In this work, we have demonstrated that PdAu SAA catalysts have excellent selectivity and stable activity for the hydrogenation of 1-hexyne to 1-hexene. PdAu SAA NPs were synthesized through a sequential reduction approach with a Pd/Au ratio of 1/250, and the exclusive presence of isolated single-atom Pd sites in the surface of Au NPs was confirmed by CO-IR. We found the hydrogenation activity of Au NPs was enhanced 10-fold through addition of this small amount of Pd atoms in the Au surface. This is attributed to the improved hydrogen activation on PdAu SAA NPs. Moreover, high selectivity to 1-hexene was achieved. PdAu SAA NPs retain stable structure and longer-term activity in the hydrogenation reaction in the liquid phase. The properties of the PdAu SAA NPs were rationalized by examining the model catalyst analogues in which a different amount of Pd was alloyed into the surface of Au(111). STM found that Pd atoms were isolated in Au(111) when the Pd coverage was low, while CO- and H$_2$-TPD showed weaker binding of CO and low-temperature uptake and release of H$_2$ on the single Pd atom sites as compared to Pd(111). Thus, PdAu SAAs can be used for the selective hydrogenation of alkynes to alkenes and serve as practical stable catalysts for these reactions under industrially relevant conditions. This study also provides a well-defined catalytic material that can be applied in a wide range of reactions including oxidation and dehydrogenation reactions.

The selective hydrogenation of 1-hexyne was investigated with IR in DRIFTS and ATR reaction cells. We observed the adsorption of 1-hexyne on both metal NPs and the silica supports. No IR peaks for dissociative adsorption species were detected. The
conversion of 1-hexyne and formation of 1-hexene were investigated under the reaction condition with liquid phase 1-hexyne and above-ambient pressure hydrogen.

Moreover, we demonstrated a different selective hydrogenation chemistry over PdAu SAA catalysts compared to that of monometallic Pd. DFT calculations reveal 1-hexyne adsorbed on the three-fold sites on extended Pd surface, but on the Pd-Au bridge sites on the surface of PdAu SAA. The two catalysts showed different reactivity, selectivity and apparent activation energy. The role of single-atom Pd in the surface of Au NPs is to activate the hydrogen molecules and stabilize the adsorption of reactant and product molecules, which activate the otherwise inert Au surface for selective hydrogenation of 1-hexyne. Moreover, PdAu SAA catalysts are more active for the 1-hexyne hydrogenation than the 1-hexene hydrogenation, such that PdAu SAAs are demonstrated as a class of highly selective catalysts for alkyne conversion selectively to alkenes.

Overall, this work demonstrates that the SAA approach can be applied in the design of various bimetallic catalysts to achieve high selectivity and reactivity.

6.5 References


Chapter 7. Conclusions and Recommendations

7.1 Conclusions

This thesis reports the preparation of single-atom alloy (SAA) nanoparticle (NP) catalysts. With well controlled galvanic replacement methods, PtCu SAA NPs were formulated on alumina and silica supports. Using a variety of characterization techniques including EDS, STEM, in situ FTIR, XRD, in situ XANES and in situ EXAFS, the presence of isolated Pt atoms in the surface of Cu NPs was demonstrated. Moreover, the PdAu SAA NP catalysts were prepared with sequential reduction method with silica support or unsupported.

This novel class of catalysts is of great fundamental and practical importance:

First, SAA NPs are the ideal platform materials in studying the chemical reactions in the single-atom sites under ambient pressure conditions. Unlike the conventional bimetallic catalysts, where single atoms, clusters and monometallic NPs co-exist, SAA NPs contain only single-atom sites in the surface of the host metal. SAAs indeed effectively bridge the materials gap between NP catalysts and model catalyst single crystals. We and our collaborators performed catalysis, surface science and DFT studies and found good convergence in the results from different sides. This capability enabled us to understand the specific chemical reactions from the molecular level to reactor level.

Second, SAA NP catalysts provide the solutions to many practical challenges in the Pt group metal (PGM) catalysts. PGMs are among the most widely used transition metal catalysts due to their superior catalytic performance for both dehydrogenation and hydrogenation reactions. Despite their favorable reactivity, there are three major drawbacks of PGM catalysts.
First, PGMs, especially Pt and Pd are very expensive, which imposes major limitations on their future role in catalysis. Second, the intrinsic selectivity of monometallic PGMs is low for hydrogenation and dehydrogenation at high conversions. Third, Pt and Pd are susceptible to carbon monoxide (CO) poisoning because CO binds strongly to metallic Pt and Pd. Indeed, SAAs are the hybrid systems containing only the minimum number of Pt (or Pd) sites necessary to perform or to assist a target reaction that retain most of the high activity of Pt (or Pd), at similar high selectivity as that of the host metal (Cu, Au), and increase the PGM tolerance to CO poisoning.

Following the invention of the SAA catalysts, we have applied them in the selective hydrogenation of 1,3-butadiene, CO tolerant H\textsubscript{2} activation, coke resistant selective dehydrogenation of butane, selective hydrogenation of 1-hexyne, selective dehydrogenation of methanol, selective dehydrogenation of ethanol and selective dehydrogenation of formic acid reactions. This thesis reports on the first four reactions among them. For selective hydrogenation and dehydrogenation reactions, the selectivity to partial hydrogenation/dehydrogenation products is very important. SAA catalysts with exclusive single-atom Pt or Pd sites largely avoid the side reactions and prove to be highly selective and stable for these reactions.

In PtCu SAA catalysts, Pt single atoms lower the hydrogen activation and desorption barrier on Cu, and enable the hydrogenation on Cu with spillover of hydrogen. With the addition of a small amount of Pt single atoms in the surface of Cu NPs, the 1,3-butadiene hydrogenation activity is improved more than one order of magnitude, while the high selectivity to butenes is maintained. In the presence of excess propylene, the PtCu SAA catalysts
are still highly selective in 1,3-butadiene hydrogenation to butenes with less than 1% conversion of propylene. Moreover, the PtCu SAA catalysts are highly stable in the hydrogenation reaction conditions.

PtCu SAAs are highly CO tolerant due to the electronic effects in SAAs. The alloying of Pt with Cu alters the electronic properties of Pt leading to weak adsorption of CO. Thus, PtCu SAA catalysts show higher CO tolerance compared to monometallic Pt catalysts from near ambient temperature to 200 °C with CO concentration from 200ppm to 3.3%. For instance, PtCu SAA shows one order of magnitude higher H₂ activation activity than monometallic Pt at 150 °C with 3.3% CO in the gas phase. In the practical low temperature and low CO partial pressure conditions (200 ppm CO, 70 °C), PtCu SAA shows much higher CO tolerance compared to monometallic Pt. Moreover, the interaction between SAA surface and CO molecules was studied with CO-FTIR and corroborative surface science studies. Therefore, the PtCu SAA is a promising candidate for improvement of many CO-poisoned systems, such as fuel cell electrocatalysts.

Pt is the most often used metal catalyst for alkane dehydrogenation reactions in industry, and its alloys with Sn are coke resistant to a certain extent but still need considerable coke removal treatments in the reaction process. In alkane dehydrogenation reactions, the first C-H activation is the rate limiting step. PtCu SAA can combine the C-H activation chemistry of single-atom Pt and the weak bonding nature of the Cu surface, which is ideal for coke resistance C-H activation. Our flow reactor studies demonstrate that with the addition of single-atom Pt in the surface of Cu, the “light off” temperature of C-H activation is lowered by almost 300 °C. Single-atom Pt, the smallest possible and most material efficient Pt ensemble, is enough
to facilitate the C-H activation in the surface of Cu. Moreover, PtCu SAA catalysts show excellent coke resistance in the C-H activation processes. This is indeed due to the highly selective nature of SAA catalysts, while the undesired by-product in this case is carbon coke. On extended Pt surfaces, the strong binding between Pt and reactively formed molecules lead to the over-dehydrogenation and other side reactions. But PtCu SAA binds the molecules weakly, the uphill energy barrier for over-dehydrogenation makes the desorption of the reactively formed molecules more favorable. The PtCu SAA catalysts were applied in the selective dehydrogenation of butane to butene under practical reaction conditions. They showed considerable reactivity with high stability without carbon deposition for at least 50 hours. The apparent activation energy of butane dehydrogenation over PtCu SAA is higher than that of monometallic Pt, indicating different energy landscape of the two catalysts.

In sum, this thesis has investigated the selective hydrogenation, CO-tolerant hydrogen activation and selective dehydrogenation reactions with PtCu SAA catalysts. For both selective hydrogenation and selective dehydrogenation reactions, commercial catalysts based on Pt or Pd are available, where Pt or Pd are in the bulk phase with additives to improve the selectivity or stability. In this thesis, we introduced a new SAA design idea for the selective hydrogenation and dehydrogenation catalysts. The Pt ensembles are designed as single-atom sites and loaded into the surface of Cu to activate the otherwise inert Cu for these reactions. The single-atom Pt can catalyze the rate limiting steps in these reactions but cannot catalyze the undesired side reaction. Although the rate limiting steps in selective hydrogenation and dehydrogenation are H₂ and C-H activation, respectively, which are the two reactions at the two ends of the spectrum of activation barrier, single-atom Pt is effective in both. On the other hand, Cu binds
the reactant and product molecules weakly, which makes it intrinsically selective. PtCu SAA is indeed a bifunctional catalytic system. We postulate that the SAA approach can be applied to a wide range of bimetallic systems for different reactions to design highly selective and stable catalysts. Indeed, the SAA strategy will become important for the “rational catalyst design” tool set.

Furthermore, to demonstrate the applicability of the SAA strategy in other catalytic systems, we designed and prepared PdAu SAA NP catalysts. A sequential reduction method was developed as galvanic replacement is not possible for Pd-Au. We find PdAu SAA catalysts are highly reactive and selective in partial hydrogenation of 1-hexyne to 1-hexene in the liquid phase. They have stable activity and selectivity without leaching. The kinetics studies find that PdAu SAAs have different reactivity, selectivity and apparent activation energy compared to monometallic Pd, revealing different hydrogenation chemistries. The role of single-atom Pd sites is to activate the hydrogen molecules, which is the rate limiting step, and stabilize the adsorption of reactant and product molecules. The surface science and DFT studies were performed to extend the understanding of PdAu SAA catalysts. Their results corroborate the findings in catalysis and in operando FTIR studies at ambient and above ambient pressures.

7.2 Recommendations

SAA catalysts based on Pt and Pd have shown promising performance in various selective hydrogenation and dehydrogenation reactions. To investigate Ni based SAA catalysts will be of great interest in decreasing the use of precious metals and providing the high-
performance catalysts for other reactions. Giannakakis et al. has demonstrated NiAu SAAs as the selective and active catalysts for the non-oxidative dehydrogenation of ethanol to acetaldehyde.\(^1\) Recent work by Shan et al. shows the alcohol dehydrogenation reactivity and selectivity of NiCu SAA catalysts. NiCu SAA appears to be more reactive in C-H bond breaking in alcohols compared to PtCu and PdCu SAAs.\(^2\) The NiCu SAA catalysts might be good candidates for a wide variety of selective dehydrogenation and oxidation reactions of hydrocarbons and alcohols. Moreover, DFT calculations performed by Stamatakis et al. show stronger adsorption of CO on Ni based SAAs compared to monometallic Ni, unlike the weaker adsorption on PtCu and PdCu SAAs. This suggests a different alloying effect in Ni-Cu, which might be useful for reactions such as water gas shift and CO oxidation reactions.

I recommend evaluating the reaction network and developing the reaction kinetics models for the selective hydrogenation and dehydrogenation reactions with PtCu SAA catalysts. The activation energy of each elementary steps can also be obtained. Through comparing the reaction mechanisms between PtCu SAA, Cu and Pt catalysts, we will be able to evaluate the structure-function relationship in the SAA catalysts. These results would indicate the nature of the active sites for the different reaction steps and which one is the rate-determining-step. This would extend our understanding of the SAA as a bifunctional catalyst. It would also be useful to identify reactions for which this is not a bifunctional catalyst. I also suggest performing the microkinetic modeling with the DFT-derived parameters to gain insight into the mechanistic aspects of the catalyst activity and guide further improvement of the SAA catalysts.
References
